

Corrigendum (Full version) to «**PYRITE IRON ISOTOPE COMPOSITIONS TRACK LOCAL SEDIMENTATION CONDITIONS THROUGH THE SMITHIAN-SPATHIAN TRANSITION (EARLY TRIASSIC, UTAH, USA)**»

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Balmat pyrite standard (*Whitehouse and Fedo, 2007; Marin-Carbonne et al., 2011*) is known to have variable $\delta^{56}\text{Fe}$ values (*Xu et al. 2022*). We have recently reanalyzed the batch of Balmat standard from UNIL (hereafter called Balmat-UNIL) at the University of Chicago (courtesy of Dr. Rego and Prof. Dauphas, unpublished data). The $\delta^{56}\text{Fe}$ bulk value of Balmat-UNIL pyrite used in this study is $-1.459 \pm 0.024 \text{‰}$, compared to the previous published value of -0.399‰ (*Whitehouse and Fedo, 2007*). As all the SIMS data are standardized to the Balmat-UNIL pyrite, the correction with this new value led to a shift of 1.06‰ of the entire published dataset. This corrigendum presents the corrected pyrite Fe isotope data, now displaying variations between -3.05‰ and $+4.33 \text{‰}$. Most of the figures (**Figs. 3, 4, 6, 7, 8, 9, 10 and 11**), the supplementary data (**Figs. S3, S5, S6 and Table S9**) and **sections 3.5** and **3.6** (results) need to be modified as presented below. As a result of this shift, we also propose hereafter an alternative model to explain the $\delta^{56}\text{Fe}$ signal of pyrite precipitated in the inner ramp system (**section 4.3.1** - discussion), combining petrographic observations and both pyrite S and newly corrected Fe isotope results. The authors would like to apologize for this mistake.

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Abbreviations: Smithian/Spathian Boundary (SSB), Secondary Ion Mass Spectrometer (SIMS), Scanning Electron Microscope (SEM), Microbial Sulfate Reduction (MSR), Dissimilatory Iron Reduction (DIR)

Abstract

The late Smithian and the Smithian-Spathian boundary (SSB) are associated with harsh environmental conditions, including abrupt temperature changes, oceanic acidification and oxygen deficiency causing an additional marked loss of biotic diversity in the aftermath of the end-Permian mass extinction. Such environmental disturbances are documented worldwide through large fluctuations of the C, O, S and N biogeochemical cycles. This study presents secondary ion mass spectrometry pyrite Fe isotope analyses from the Lower Weber Canyon (LWC) section (Utah, USA) combined with bulk rock $\delta^{34}\text{S}_{\text{py}}$ and $\delta^{34}\text{S}_{\text{CAS}}$ analyses in order to better understand the redox changes in different environmental settings along a ramp depositional system through the SSB. $\delta^{56}\text{Fe}$ analyses show a large variability along the studied ramp system of $\sim 7\text{‰}$ (from -3.05 to $+4.33\text{‰}$), over a set of 350 microscale analyses. Bulk sulfide sulfur isotope analyses, performed on 30 samples, show $\delta^{34}\text{S}_{\text{py}}$ varying from -20.5 to $+16.3\text{‰}$. The inner ramp domain is characterized by a mean negative $\delta^{34}\text{S}_{\text{py}}$ value of -11.4‰ . A progressive ^{34}S -enrichment (up to $+16.3\text{‰}$) is recorded in pyrite from mid and outer ramp settings. Carbonate-associated sulfate (CAS) sulfur isotope analyses, performed on 5 samples, show relatively steady $\delta^{34}\text{S}_{\text{CAS}}$ of $+30.2 \pm 2.2\text{‰}$. Variations in $\delta^{34}\text{S}_{\text{py}}$ are interpreted as reflecting the degree of connection between sediment porewaters and the overlying water column. Multiple lines of evidence point to a fully oxygenated water column and thus restricts pyrite formation to the sediments. Both the sedimentary environment and the nature of deposits seem to control $\delta^{56}\text{Fe}_{\text{py}}$. In the inner ramp, high $\delta^{56}\text{Fe}_{\text{py}}$ values averaging $+0.97\text{‰}$ are only observed in microbially induced sedimentary structures (MISS), which reflect the complete reduction of Fe-oxides. In the absence of MISS, $\delta^{56}\text{Fe}_{\text{py}}$ inner ramp values are lighter ($\delta^{56}\text{Fe}_{\text{mean}} = -0.16\text{‰}$) and reflect the partial reduction of Fe-oxides due to a limited production of microbial H_2S deeper in the sediments. In more distal and deeper mid and outer ramp settings, Fe isotope compositions are controlled by microbially-produced H_2S that scavenged iron into sulfides. This study unravels local redox state changes in the upper part of some marine sediments by coupling Fe and S isotope systematics. It demonstrates that pyrite grains, and their sulfur and iron isotopic compositions, formed throughout the SSB should be used with caution to infer the redox state of the ocean after the Permian-Triassic biotic crisis.

1. Introduction

Only ~2 Myr after the devastating end-Permian mass extinction (~252 Ma), the ocean-atmosphere system underwent successive and abrupt environmental and biotic changes at the Smithian-Spathian Boundary (SSB; e.g. *Galfetti et al., 2007*). Significant disturbances of the global carbon, sulfur, oxygen, and nitrogen biogeochemical cycles underline sustained environmental stresses. These include climate warming up to the beginning of the late Smithian, followed by a rapid cooling across the latest Smithian and the SSB (*Goudemand et al., 2019; Zhang et al., 2019*), the spread of anoxia through ocean stratification (*Song et al., 2019*) or development of oxygen minimum zones (*Algeo et al., 2011*), and ocean acidification potentially linked with greenhouse gases released by large igneous province eruption (*Grasby et al., 2013; Zhao et al., 2020*). All combined, these deleterious conditions led to a marked loss in ecosystem diversity and a severe extinction of several nekton-pelagic organisms during the late Smithian (*Brayard et al., 2006; Orchard, 2007; Jattiot et al., 2016*). Several proxies such as sulfate and sulfide $\delta^{34}\text{S}$ values, paired carbonate and organic matter $\delta^{13}\text{C}$ values, Ce anomalies, trace elements (e.g. Mn, V, U, Mo, Mo/Al ratio), iron speciation and size of framboidal pyrite suggest occurrences of transient anoxia in the water column that could have spread onto continental shelves (*Grasby et al., 2013; Song et al., 2013; Zhang et al., 2015; Elrick et al., 2017; Song et al., 2019; Zhang et al., 2019*). Although anoxic and/or euxinic conditions seem to be recurrent in the Early Triassic deep ocean, well-oxygenated shallow-marine environments, as well as diversified and complex ecosystems are also documented (*Beatty et al., 2008; Hautmann et al., 2011; Collin et al., 2015; Brayard et al., 2017; Olivier et al., 2018; Smith et al., 2021; Dai et al., 2023*). Reconstructing the redox state of the water column together with the sediment porewaters is therefore key to better understand the evolution of the environmental conditions in space and time that prevailed through the late Smithian and the SSB, and that may have markedly influenced the biotic recovery in the aftermath of the end-Permian mass extinction.

As a mineral made of redox-sensitive elements, pyrite (FeS_2) is used to infer the oxygenation state of the ocean through the study of its morphology, size, trace element content and isotope (S and/or Fe) compositions (*Wilkin et al., 1996; Wilkin and Barnes, 1997; Shen et al., 2003; Bond and Wignall, 2010; Gregory et al., 2015*). Oxidation of Fe(II) into Fe(III) is associated with a large equilibrium fractionation producing ^{56}Fe -depleted dissolved Fe(II) (*Welch et al., 2003*), which can be later recorded in Fe-bearing minerals such as pyrite. Large ^{56}Fe -depletion during redox processes is however only expressed for the partial reaction of the dissolved iron reservoir, i.e. during partial oxidation of aqueous Fe^{2+} (*Rouxel et al., 2005*). Consequently, assuming that redox reaction occurred through Earth history and that $\delta^{56}\text{Fe}$ values reflect primary processes, secular variations of Fe isotope compositions can be interpreted in terms of global oxygenation state changes (*Rouxel et al., 2005*). Other studies propose that low $\delta^{56}\text{Fe}$ values reflect transport of microbially reduced dissolved Fe(II) from the shelf to the basin along the chemocline (*Johnson and Beard, 2005; Severmann et al., 2008*). Alternatively, another scenario, which does not require Fe redox cycling, suggests that $\delta^{56}\text{Fe}$ values reflect primarily the rate of pyrite precipitation (kinetic or equilibrium) that is highly dependent on the sulfur availability (*Guilbaud et al., 2011; Mansor and Fantle, 2019*). Therefore,

the interpretation of pyrite iron isotope compositions in terms of water oxygenation state can be obscured by the involvement of redox and non-redox processes, inheritance of isotopic fractionation of the Fe source, size of the sulfide reservoir, and early diagenetic processes (*Severmann et al., 2008; Virtasalo et al., 2013; Busigny et al., 2014; Marin-Carbonne et al., 2020*). Further, on a more local scale, processes occurring within the upper part of the sedimentary column (from the upper centimeters to meter depth), can produce Fe isotope variations up to 5 ‰. Such variations are partially to entirely decoupled from global biogeochemical cycling in the water column, resulting in the isotopic evolution of Fe(II) or Fe-oxide reservoirs in the sediments (*Archer and Vance, 2006; Virtasalo et al., 2013; Lin et al., 2017*). Some ambiguities in the interpretation of the pyrite $\delta^{56}\text{Fe}$ can be lessened by pairing this measurement with the $\delta^{34}\text{S}$ signal. For instance, positive covariation between pyrite $\delta^{56}\text{Fe}$ and $\delta^{34}\text{S}$ signals through Archean-Paleoproterozoic transition suggests an expansion of the sulfidic Fe sink (*Heard and Dauphas, 2020*). Such positive covariations have also been evidenced in 2.7 Ga sedimentary pyrite from the Belingwe greenstone belt and interpreted as early diagenetic signatures of coexisting microbial Fe and S reduction (*Archer and Vance, 2006*).

The aim of this study is to distinguish in the context of the Smithian-Spathian transition whether pyrite paired Fe-S isotopic signatures (1) record redox condition variations of the seawater, (2) reflect variations of the Fe/S ratio or (3) are controlled by early diagenesis and/or microbial processes within the sediments or at the seawater/sediment interface. For this purpose, we present spatially resolved $\delta^{56}\text{Fe}$ associated with chemostratigraphic $\delta^{34}\text{S}$ records along the Lower Weber Canyon sedimentary succession (**Fig. 1a**; Utah, USA), which is characterized by various sedimentary rocks deposited between shallow inner to deeper outer ramp settings. The combination of petrological observations, in situ $\delta^{56}\text{Fe}_{\text{py}}$ analyses by SIMS, bulk rock $\delta^{34}\text{S}$ analyses and total organic carbon concentrations allowed us to unravel pyrite formation pathways within the sediments from different facies.

2. Materials and methods

2.1. Geological context, studied section and samples

During the Permian-Triassic interval, the near equatorial region of western Pangea was an active tectonic compression zone, leading to the formation of the Sonoma orogeny and the associated Sonoma Foreland Basin (SFB; **Fig. 1**; *Burchfiel and Davis, 1975; Ingersoll, 2008; Dickinson, 2013*). Lower Triassic sedimentary rocks of the SFB mainly cover present-day eastern Idaho, eastern Nevada and Utah (USA; *Caravaca et al., 2018*). The Early Triassic sea-level rise resulted in a sedimentary succession exhibiting continental terrigenous conglomerates and sandstones of the Moenkopi Group on the south and eastern sides of the basin, interfingering with marine carbonates of the Thaynes Group towards the north and western sides (*Paull and Paull, 1993; Lucas et al., 2007; Brayard et al., 2013*). In the SFB, a transgression that corresponds to the second 3rd order T-R (transgressive-regressive) sequence of the Smithian is recorded (*Haq et al., 1987; Embry, 1997*), which was

controlled by regional tectonics and climate (Olivier et al., 2014; Caravaca et al., 2018; Brayard et al., 2020). Calibrations of late Smithian to early Spathian sedimentary sections within the basin are based on ammonoid biostratigraphy (Guex et al., 2010; Brayard et al., 2013, 2021). The late Smithian Anasibirites beds represent the maximum flooding at the scale of the basin (Brayard et al., 2013, 2020). The SFB is also characterized by occurrence of morphologically various, mineralized or not, microbial deposits, mainly in the southern part of the SFB (Schubert and Bottjer, 1992; Pruss et al., 2004; Brayard et al., 2011; Olivier et al., 2014, 2016, 2018; Woods, 2014; Vennin et al., 2015; Grosjean et al., 2018; Kirton and Woods, 2021). The studied sedimentary section (described below) includes Smithian microbially deposits characterized by microbially induced sedimentary structures (MISS; see Grosjean et al., 2018).

This study focuses on the Lower Weber Canyon (LWC) section, for which petrographic descriptions and interpretation in terms of depositional environments are detailed in Grosjean et al. (2018). According to these authors, the LWC section can be divided in four lithologic units (A-D).

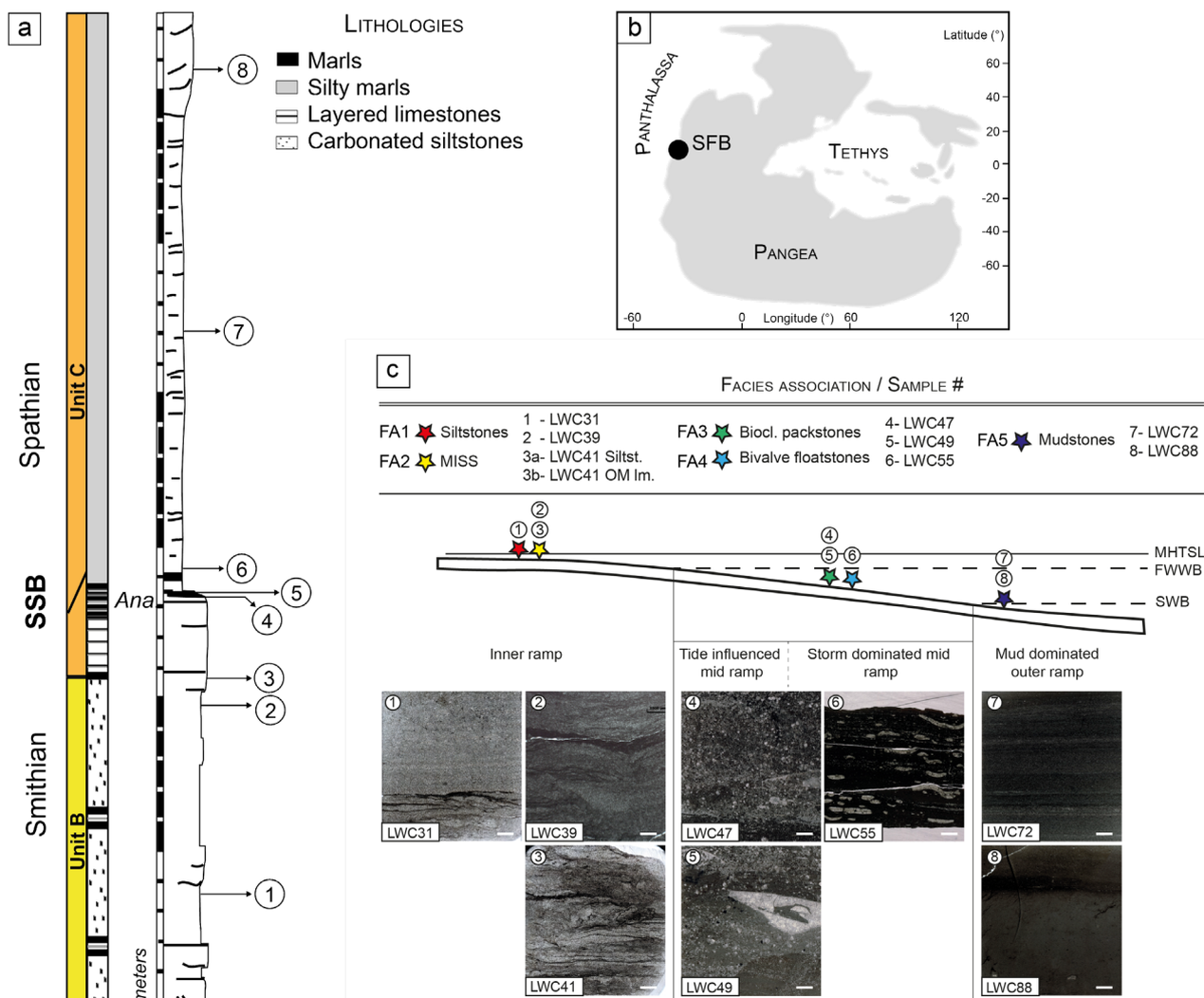


Figure 1: a) Log of the LWC section with focus on Units B and C (modified after Grosjean et al., 2018). b) Early Triassic paleogeographic map showing the location of the Sonoma Foreland Basin (SFB; modified after Brayard et al., 2013). c) Position along a ramp system and thin section pictures of studied samples (scale bar for thin sections: 2.5 mm). The different star colors refer to the corresponding facies association (FA1-FA5). MHTSL: Mean High Tide Sea Level; FWWB: Fair Weather Wave Base; SWB: Storm Wave Base.

This study concentrates on the two units (B and C) preceding and including the SSB (**Fig. 1**). Unit B is 111 m thick and mainly made of carbonated siltstones, including organic-rich laminae recognized as MISS, and few levels of bioclastic calcarenites. Siltstone-rich sediments of the Unit B reflect a peritidal depositional environment in an inner ramp system. Sediments of this unit are of middle Smithian age as they belong to the Meekoceras ammonoid assemblage (*Grosjean et al., 2018; Brayard et al., 2021*). The 36 m-thick Unit C consists of silty mudstones that are intercalated with bioclastic limestones. The lower part of this unit includes the late Smithian Anasibirites ammonoid assemblage (*Brayard et al., 2021*) and the SSB. The mudstone-rich deposits of the Unit C reflect a mid to outer ramp setting that occasionally recorded storm events. Eight rock samples were collected in the Units B and C, thus representing different depositional environments distributed along an inner to outer ramp system (**Fig. 1**). Their deposition during the SSB, the diversity of depositional environments (from peritidal siltstones to deeper outer ramp mudstones), and associated facies including occurrences of MISS in a shallow and proximal environment, are relevant parameters to decipher controlling parameters on Fe isotope signatures of pyrite. A description of facies associations and corresponding depositional environments is detailed in the **section 3.1**.

2.2. Methods

2.2.1. Petrographic observations

Petrographic descriptions were carried out on polished thick sections using a Scanning Electron Microscope (SEM) at ISTE Lausanne (Tescan Mira LMU). Backscattered electron images were performed at a working distance of 21 mm using a voltage of 20 kV and a current of 1.5 nA. Semi quantitative spot analyses were done by energy dispersive X-ray spectrometry to determine elemental composition of the sulfides. Bulk rock mineralogical analyses were conducted at the Institute of Earth Sciences of the University of Lausanne, Switzerland (Thermo Scientific ARL X-TRA Diffractometer) using the procedure described by *Klug and Alexander (1974)* and *Adatte et al. (1996)*. These analyses were performed on powdered samples pressed into powder holders. This method for semi-quantitative analysis of the bulk rock mineralogy used external standards with elemental compositions based on mineral stoichiometries. The error varies between 5 and 10 % for the phyllosilicates and 5 % for grain minerals. The non-quantified portion corresponds generally to poorly crystallized iron hydroxides, phyllosilicates, and nearly amorphous material. This portion represents here less than 2 % of the bulk composition.

The chemical composition of pyrite was investigated by Electron Probe Micro-Analyzer (EPMA) using a JEOL JXA-8530F at ISTE Lausanne in order to quantify the trace content of zoning pyrite. The acceleration voltage was 15 kV and the beam current was 15 nA focused in 3 μ m. Reference materials, including sulfides, oxides, and silicates, were tested before the analysis of S, Fe, Co, Mn, Pb, Ti, Cr, Zn, Cu and Ni. Detection limits were 133 ppm for S, 130 ppm for Fe, 141 ppm for Co, 129 ppm for Mn, 327 ppm for Pb, 79 ppm for Ti, 145 ppm for Cr, 238 ppm for Zn, 199 ppm for Cu and 150 ppm for Ni.

Carbonaceous matter has been characterized by Raman spectroscopy. Raman data were obtained using a Horiba Jobin Yvon LabRAM HR800 spectrometer (ISTE, UNIL) in a confocal configuration equipped with an Ar⁺ 532 nm laser and a CCD detector. Measurements were performed at room temperature on thin sections, with a laser power delivered at the sample surface <1 mW to prevent thermal damage. This technique allows to document the structural organization of the aromatic skeleton of carbonaceous matter, and to estimate the peak metamorphic temperature experienced by the carbonaceous matter (*Beysac et al., 2002*). Peak temperature estimations were done following the methodology proposed by *Kouketsu et al. (2014)*.

2.2.2. Hg concentrations and TOC contents

Mercury concentrations were measured using a Zeeman R-915F (Lumex, Saint-Petersburg, Russia) high-frequency atomic absorption spectrometer at the University of Lausanne. Hg was extracted by thermal evaporation at 700°C from solid samples that were previously powdered. Analyses were bracketed by the measurement of a certified external standard (GSD-11, Chinese alluvium, Hg concentration: 72 ±6 ppb) and two aliquots were systematically analyzed. To be exempt of biased Hg peaks due to affinity of Hg with organic carbon and sulfide, measured Hg concentrations were normalized to TOC and sulfur concentrations.

Total organic carbon (TOC in wt.%) contents were obtained through Rock-Eval 6 analyses at University of Lausanne following the procedure of *Espitalié et al. (1985)* and *Behar et al. (2001)*. The reference material IFP 160000 (French Institute of Petroleum) was used to calibrate the measurements and the precision was <0.1 %. Analyses consist of a combined pyrolysis of ~60 mg of powdered samples followed by oxidation of the residue.

The analyses of Trace element concentrations were performed by Inductively Coupled Plasma Mass Spectrometry (ICP-MS) at the Activation Laboratory (Actlabs) in Ancaster, Canada. The method used is Lithium Metaborate/Tetraborate Fusion ICP and ICP-MS. Reproducibility was checked by seven duplicate and replicate analyses of laboratory standards and is better than 2 % (1σ) for major and trace elements.

2.2.3. Bulk rock δ³⁴S

Sulfide sulfur was retrieved using a conventional wet chemical extraction at the Biogéosciences Laboratory, Dijon, France. Acid volatile sulfide (AVS), representing monosulfides, and chromium reducible sulfur (CRS) consisting primarily of pyrite were extracted sequentially. First, AVS was liberated using cold concentrated HCl for 2 hours. If any, resulting hydrogen sulfide was precipitated as Ag₂S with a 0.3 M AgNO₃ solution. Subsequently, CRS was released using a hot and acidic 1.0 M CrCl₂ solution following the method described in *Gröger et al. (2009)*. The resulting H₂S was precipitated as Ag₂S. After centrifugation, the silver sulfide precipitate was washed several times with deionized distilled water and dried at 50°C for 48 hours in an oven and weighted. Mass balance was used to calculate the amount of AVS and CRS. Concentrations in ppm are reported individually in **Table S1** (note that AVS were below the detection limit in all samples).

CAS sulfur was retrieved using a wet chemical extraction. Fifty grams of carbonate samples

(with carbonate contents typically >70 wt.% of the total rock) were powdered, soluble sulfates being leached by a 1.7 M NaCl solution for 4 hours, followed by four rinses in deionized water. This step was repeated five times and the powder was then dissolved in 4 N HCl (12 h). The acidified samples were then filtered, on a 0.45 μm nitrocellulose paper and an excess of 250 g.l^{-1} of BaCl_2 solution was added to the filtrate to precipitate BaSO_4 . After centrifugation, the barium sulfate precipitate was washed several times with deionized distilled water and dried at 60°C for 24 hours in an oven.

The $\delta^{34}\text{S}$ measurements were performed at the Biogéosciences Laboratory, Dijon, France, on SO_2 molecules via combustion of $\sim 500 \mu\text{g}$ of samples (silver sulfide and barium sulfate) homogeneously mixed with an equal amount of tungsten trioxide using a Vario PYRO cube (Elementar GmbH) connected online via an open split device to an IsoPrime isotope ratio mass spectrometry (IRMS) system (Isoprime, Manchester, UK). International standards IAEA-S-1, IAEA-S-2, IAEA-S-3 and NBS-127 were used for calibration assuming $\delta^{34}\text{S}$ values of -0.3 ‰, +22.7 ‰, -32.3 ‰ and 20.3 ‰ (VCDT), respectively. Analytical reproducibility was better than $\pm 0.4 \text{‰}$ (1σ) based on replicates for standard materials and samples.

2.2.4. Fe isotope analyses by SIMS

Iron isotope compositions were measured over four SIMS sessions (March 2020, June 2020, November 2020, September 2021) using the Cameca ims 1280 HR2 at the SwissSIMS (University of Lausanne). Iron isotope compositions are expressed in delta notation, reporting permil variations of the $^{56}\text{Fe}/^{54}\text{Fe}$ ratios normalized to IRMM-014 as the following:

$$\delta^{56}\text{Fe} = \left[\frac{\left(\frac{^{56}\text{Fe}}{^{54}\text{Fe}} \right)_{\text{sample}}}{\left(\frac{^{56}\text{Fe}}{^{54}\text{Fe}} \right)_{\text{IRMM014}}} - 1 \right] * 1000$$

In order to preserve high vacuum conditions of $\sim 10^{-9}$ mbar in the analytical chamber and avoid hydride formations, thick sections were polished and pressed into one-inch indium mounts. Balmat-UNIL pyrite standard was used over the sessions to assess the stability of the instrument ($\delta^{56}\text{Fe} = -1.46 \pm 0.02 \text{‰}$, courtesy of Dr. Rego and Prof. Dauphas, unpublished data) and was included in each sample mount. The conductivity of the sample surface was ensured by a 35nm-thick gold coating. A total of 350 analyses of pyrite were performed in 8 different samples following the method described in Decraene et al. (2021). Briefly, a 10 kV, 3 nA Gaussian $^{16}\text{O}^+$ primary beam was focused into a 2.5 to 3 μm spot. $^{52}\text{Cr}^+$, $^{54}\text{Fe}^+$, $^{56}\text{Fe}^+$ and $^{57}\text{Fe}^+$ secondary ions were collected simultaneously by electron multipliers (L2 for $^{52}\text{Cr}^+$) and off-set faraday cups (C for $^{54}\text{Fe}^+$, H1 for $^{56}\text{Fe}^+$ and H'2 for $^{57}\text{Fe}^+$). We used an entrance slit width of 61 μm and the slit 3 of the multicollection to obtain a Mass Resolving Power at ~ 7000 (interference of ^{53}CrH on ^{54}Fe). The field aperture was set at 2000 μm and the energy slit at 50 eV. A 90 seconds presputtering time was applied to remove the surface contamination, followed by secondary ion beam centering and background detector measurements. Then, data acquisition is conducted for 300 seconds. Typical count rate for $^{56}\text{Fe}^+$ on Balmat-UNIL standard was $4.50\text{E}+7$ counts per second (cps) and the mean reproducibility was $\pm 0.21 \text{‰}$ (2SD; **Table S2**). Instrumental mass fractionation (IMF) was corrected by standard bracketing. Iron isotope

compositions are represented in the form of histogram or box plot. Histograms have been set with a bin size of 0.4 ‰, to both limit the production of artificial secondary peaks and bin boundary effects, and in a range of -2 to +6 ‰. The Shapiro-Wilk test was used to test whether the data are normally distributed (**Table S3**). In the case of Gaussian distributions, the parametric Student's T test was employed to test the hypothesis of equal means for two independent samples. In the case of non-Gaussian distributions, the non-parametric Mann-Whitney test (two sided) was used to test the hypothesis of equal medians for two independent samples. Both tests gave results as p-values with a confidence interval of 95 % (**Tables S4-S7**).

3. Results

3.1. Facies description

According to *Grosjean et al. (2018)*, the eight studied samples belong to five different facies associations. These facies are distributed along a ramp depositional profile from proximal and shallow siliciclastic dominated deposits to more distal and deeper mud-dominated sediments (**Fig. 1**). The bulk mineralogical composition of all samples is predominantly composed of calcite, quartz, phyllosilicates, and plagioclase.

The inner peritidal ramp setting is dominated by two facies associations, termed carbonated-siltstones (FA1) and organic-rich laminae (FA2). It should be noted that LWC41 encloses these two facies FA1 and FA2, named respectively LWC41 Siltst. and LWC41 OM, in which pyrite is observed. FA1 consists of calcite, quartz, phyllosilicates, and dolomite (see supplementary materials; **Fig. S1**). Small fragmented bioclasts are observed as well as accessory minerals such as apatite, zircon and anatase. This facies association corresponds to samples LWC41 Siltst. and LWC31. FA2 is composed of undulated organic-matter laminae that are laterally discontinuous and locally reworked in a carbonated and silty matrix. These organic laminae and chips are interpreted by *Grosjean et al. (2018)* as MISS. Samples LWC39 and LWC41 OM are attributed to FA2. For mineralogical analyze purposes, LWC41 was not divided into two parts to distinguish FA1 and FA2. The bulk composition of LWC41 being similar to that of LWC39, LWC41 is processed here as only a part of FA2. The bulk mineralogical composition of LWC31 (**Fig. 2**) is dominated by calcite (37 %) and quartz (32 %). The phyllosilicate content is up to 14 % and is similar to the plagioclase content (13 %). K-feldspars are present in small proportion (2 %), as well as dolomite and ankerite, which contents are below 1 %. The average bulk composition of FA2 is dominated by quartz (33 %), plagioclase (20 %), phyllosilicates (15 %) and calcite (15 %; **Fig. 2**). The proportion of carbonates is much lower than that of FA1. The dolomite content is higher than in FA1 (5 %) whereas the ankerite content is still low (1 %; **Fig. 2**).

Samples from the mid ramp setting reflect two facies associations, termed bioclastic packstones (FA3; LWC47 and LWC49) and bioclastic wackestones (FA4; LWC55). FA3 consists of calcite, phyllosilicates and quartz (**Fig. S1**). Few apatites and oxides are also observed. Bivalve

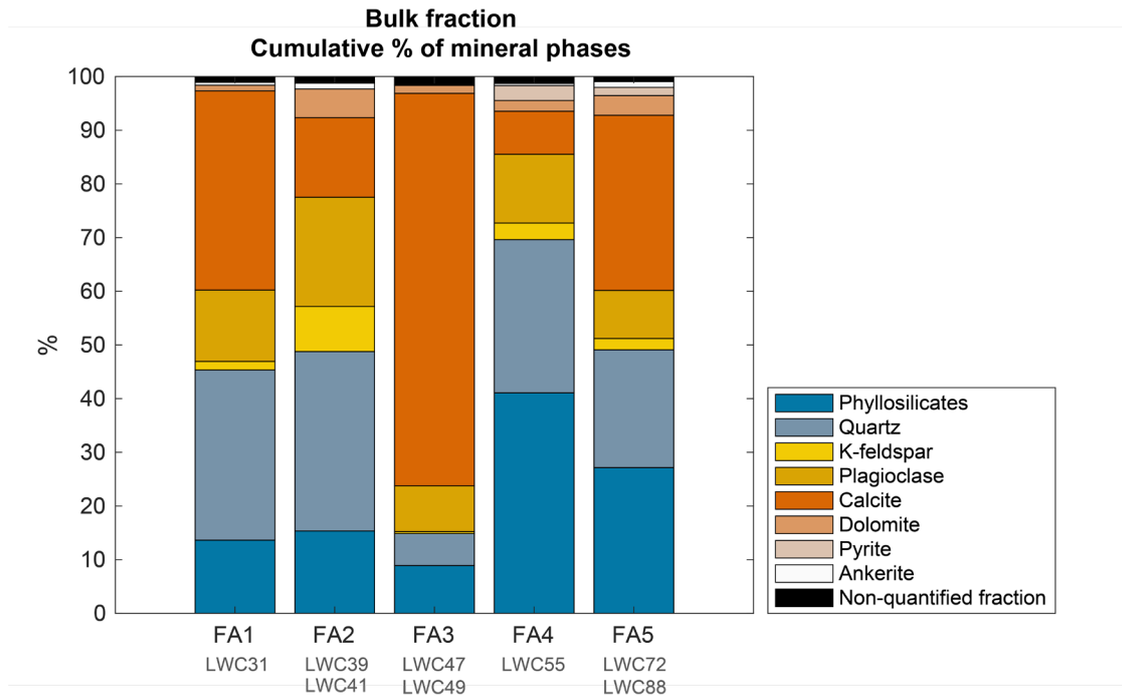


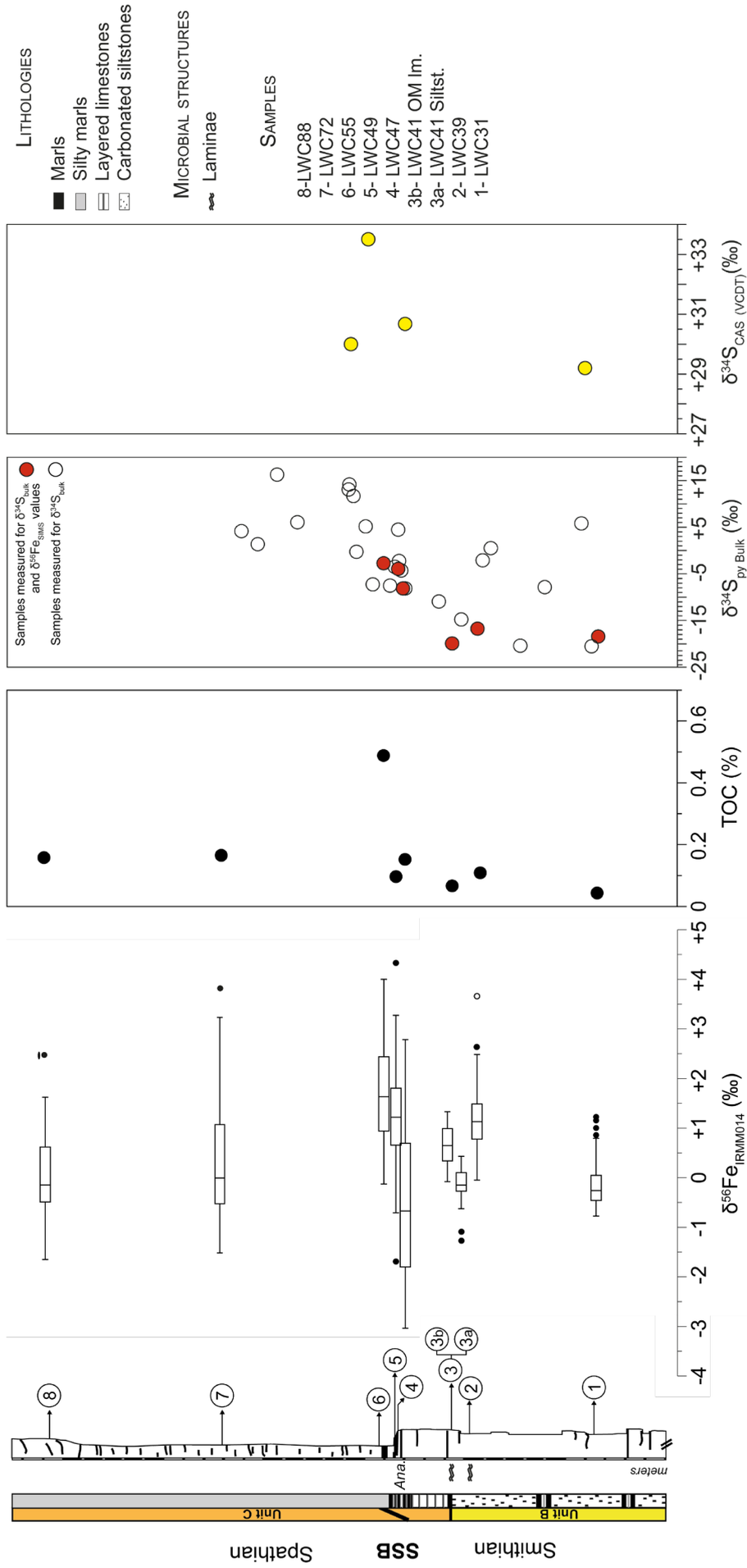
Figure 2: Bulk mineralogical composition of the 8 LWC samples classified as a function of their facies.

and ammonoid shells are recrystallized by sparite. The average bulk mineralogical composition of samples from FA3 shows a dominance of calcite (73 %) relative to the other phases (Fig. 2). Phyllosilicates and quartz represent 9 and 6 % of the bulk composition. The amount of K-feldspar is low (below 1 %) whereas the amount plagioclase is up to 9 %. The sum of the dolomite and ankerite content is below 2 % (Fig. 2). FA4 is made of sparite-filled bivalves following a planar distribution in a calcite dominated matrix with common small quartz grains. Apatite and zircon grains are present as accessory minerals. The mineralogy of LWC55 (Fig. 2) is dominated by phyllosilicates (41 %) and quartz (29 %). This sample contains 13 % of plagioclase and 8 % of calcite. The amount of K-feldspar and dolomite are of 3 % and 2 % respectively. The ankerite content is below 1 %. Pyrite was detected in a small proportion of 3 %.

Samples from the mud-dominated outer ramp correspond to a mudstone facies association (FA5; LWC72 and LWC88). These sediments deposited closed to the mid to outer ramp transition as they consist of mudstone sometimes interbedded with rare thin layers made of small quartz grains. Apatite, zircon, anatase and chalcopyrite are present as accessory mineral phases. The bulk mineralogical composition of samples from FA5 (Fig. 2) contains up to 33 % of calcite, 27 % of phyllosilicates and 22 % of quartz. The plagioclase and K-feldspar contents are respectively of 9 and 2 %. The respective amount of dolomite and ankerite are of 4 % and 1 %. Finally, pyrite is present in a smaller proportion than in FA4, representing 2 % of the bulk composition.

3.2. Carbon, sulfur and mercury contents

Total organic contents vary from 0.04 to 0.49 wt.% and show an enrichment across the SSB, with a maximum TOC content recorded in LWC55 (corrected Fig. 3). TOC concentrations for samples LWC39 and LWC41 are biased due to the heterogeneous distribution of organic-rich



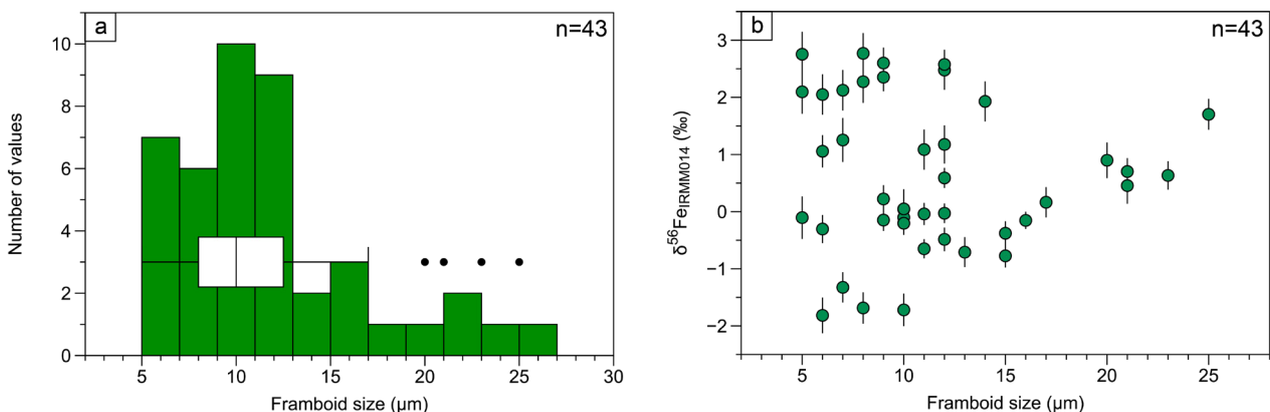
Corrected Figure 3: Evolution of Fe isotope compositions measured by SIMS on 8 LWC samples with corresponding total organic carbon concentrations (in %), pyrite $\delta^{34}\text{S}$ values on 30 samples (LWC31 to LWC70) and carbonate associated sulfate (CAS) $\delta^{34}\text{S}$ values on 5 samples of the section. The simplified log reports lithologies and positions of samples measured by SIMS.

laminae in the analyzed samples. Sulfur content of 30 LWC samples (from LWC31 to LWC70) vary between 21 and 12357 ppm with a mean value of 2585 ppm (**corrected Fig. 3 and Table S1**). Sulfur content is highly variable with highest concentrations recorded in samples deposited during and immediately after the SSB. Inner ramp samples (LWC31 to LWC42) show S concentrations between 21 and 3780 ppm, with an average of 1222 ppm (**Table S1**). There is no difference in sulfide content between samples from siltstone facies (FA1) and MISS facies (FA2). Sulfur concentrations in samples from mid ramp setting range from 296 to 11472 ppm and show the highest average value (4232 ppm). Samples from the outer ramp system exhibit the larger variation, from 61 to 12358 ppm. The Sr and Mn contents have also been measured in these 30 samples (LWC31 to LWC70; **Table S1**). The Sr content varies from 17 to 284 ppm, with an average value of 105 ppm. The Mn content varies between 264 ppm and 6900 ppm, with an average value of 2308 ppm. These 30 samples show an average Mn/Sr ratio of 24, with a range of variation between 11 and 40. Mercury concentrations vary between 2.70 and 25.55 ppb (**Table S8, Fig. S2**). Normalization with TOC and S contents does not show any Hg enrichment in the studied samples.

3.3. Bulk pyrite and CAS $\delta^{34}\text{S}$

The $\delta^{34}\text{S}_{\text{py}}$ values show a large range of variations from -20.5 to +16.3 ‰ (**corrected Fig. 3, Table S1**). Except for three samples, only negative $\delta^{34}\text{S}$ values are recorded in the inner and the mid ramp systems, while the majority of $\delta^{34}\text{S}$ values in the outer ramp are positive. The inner ramp setting shows highly variable $\delta^{34}\text{S}$ values, ranging from -20.5 to +5.8 ‰, with an average value of -11.4 ± 9.3 ‰ (1SD). The mid ramp setting documents small variations with a $\delta^{34}\text{S}$ average of -4.0 ± 4.0 ‰ (1SD). The outer ramp system displays the highest S isotope compositions, ranging from -7.3 to +16.3 ‰, and an average value of $+6.4 \pm 7.4$ ‰ (1SD). The $\delta^{34}\text{S}$ signal of sedimentary sulfide shows a ^{34}S enrichment tendency from the base to the top of the LWC section. The $\delta^{34}\text{S}_{\text{CAS}}$ signal measured on five samples across the section does not exhibit noticeable secular variations (**corrected Fig. 3; Table S1**). Indeed, $\delta^{34}\text{S}_{\text{CAS}}$ values range from 29.2 to 33.5 ‰, with a mean value of 30.2 ± 2.2 ‰ (1SD).

3.4. Pyrite description



Corrected Figure 4: a) Framboid size distribution (histogram and box plot) of the 43 framboids measured in five LWC samples (LWC47 to LWC88) and b) relationship between the size of framboid pyrite grains and their Fe isotope compositions.

In the eight studied samples, pyrite grains display various morphologies, categorized as euhedral, framboid, aggregated and framboid with secondary overgrowth. Their size ranges between 5 and $\sim 100 \mu\text{m}$ (**corrected Fig. S3**). Only LWC31 displays a large pyrite nodule of $\sim 3 \text{ mm}$ wide. Pyrite with a size of 5 to $40 \mu\text{m}$ tends to record a wider range of Fe isotopic compositions than pyrite larger than $40 \mu\text{m}$ (**corrected Fig. S3**). However, it should be noted that a sampling bias may have been introduced by measuring a larger amount of pyrite $10\text{-}20 \mu\text{m}$ in size relative to pyrite larger than $30 \mu\text{m}$.

Framboids are of particular interest as they are extensively studied to reconstruct paleoredox conditions and to constrain the locus of pyrite precipitation (*Wilkin et al., 1996*). In this study, the size of framboids ranges from 5 to $25 \mu\text{m}$ with an average size of $11 \mu\text{m}$ ($n=43$; **corrected Fig. 4a**). Moreover, the size distribution of framboids is monomodal, with a mode at around $10 \mu\text{m}$, and highlights that most of the pyrite grains have a size between 5 and $12 \mu\text{m}$ (**corrected Fig. 4a**).

The absence of correlation between the grain size and their Fe isotope compositions (**corrected Fig. 4b**) does not allow to define several generations of framboidal grains. The number of analyzed pyrite per morphology and facies and their corresponding Fe isotope compositions is detailed in **section 3.6**. Several remarkable pyrite features have been observed in the samples of the LWC section. In sample LWC41 (Siltst. and OM parts), pyrite is affected by secondary rims but still preserves framboidal cores (**Fig. 5a**). The nodule observed in LWC31 is made of an aggregate of euhedral pyrite grains whose former boundaries are still discernible at the nodule edge (**Fig. 5b**). Sample LWC49 encloses aggregated pyrite with chemical zoning enriched in Ni (**Fig. 5c**). These

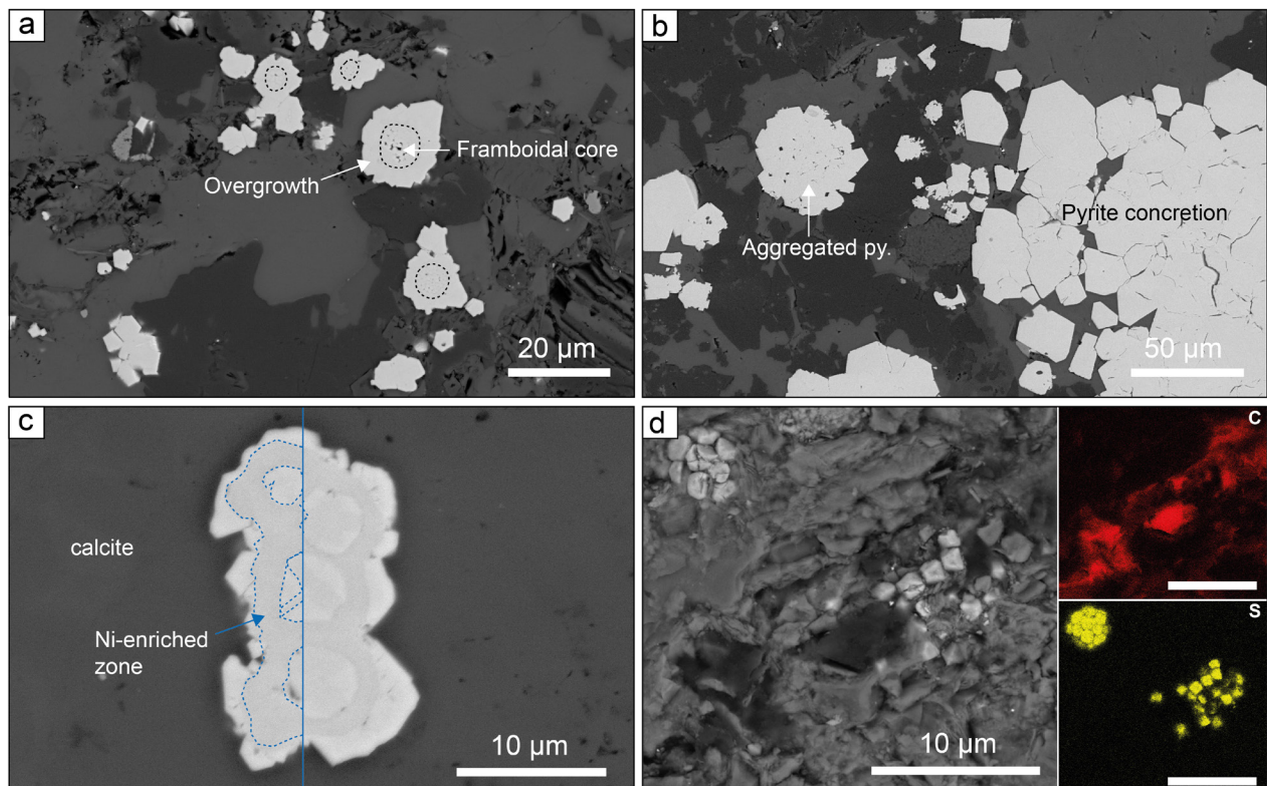
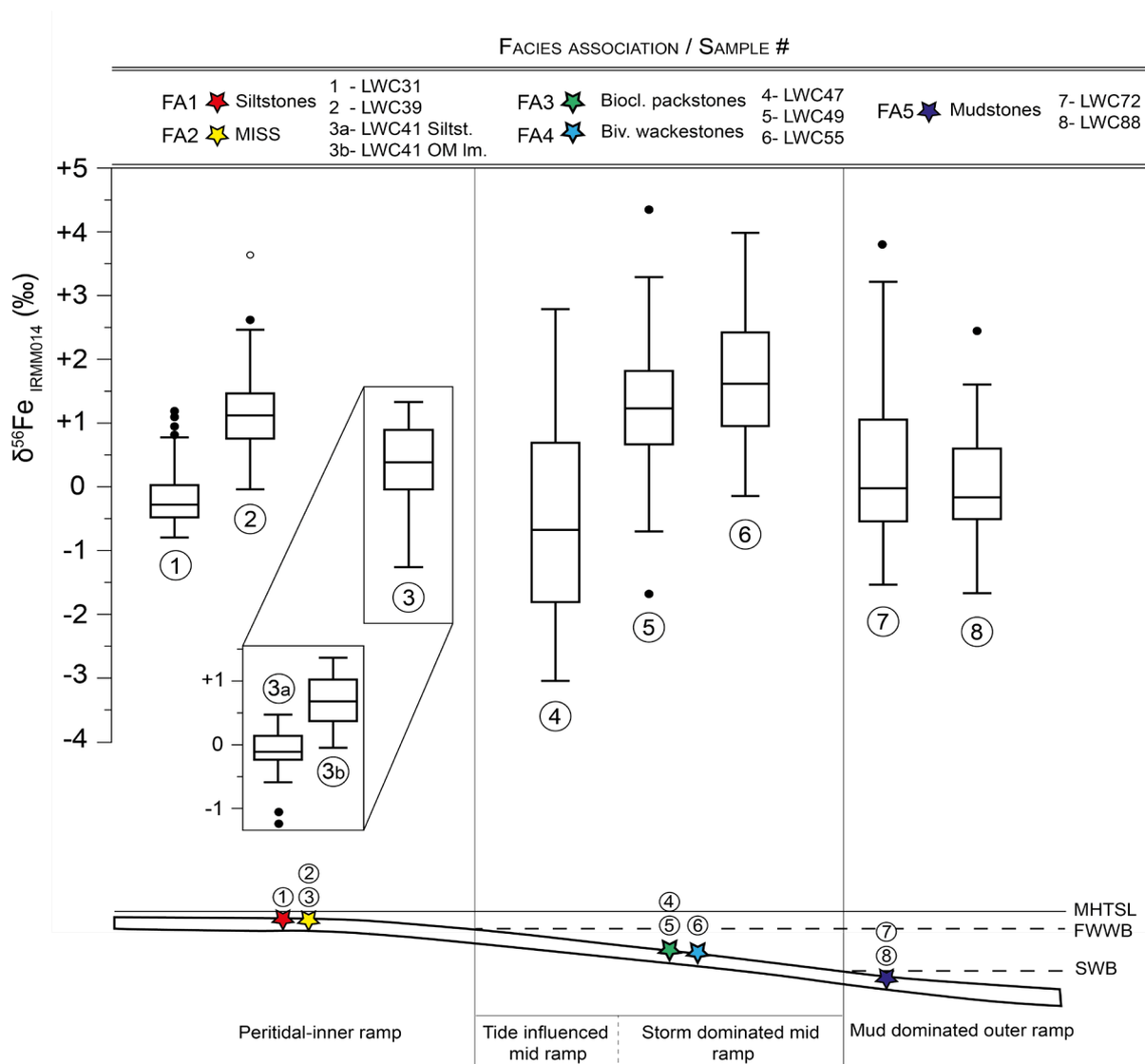


Figure 5: Backscattered images of early diagenetic pyrite observed in LWC samples. Examples of a) framboids with overgrowths from LWC41 Siltst., b) aggregated grains and edge of the pyrite nodule observed in LWC31, c) a zoned (Ni-enriched) aggregate from LWC49. The Ni-enrichment was evidenced by electronic microprobe analyses. d) Framboids associated with carbonaceous material in LWC88 and maps of carbon (in red) and sulfur (in yellow). Scale bars are $10 \mu\text{m}$.

Ni enrichments have been evidenced through electronic microprobe analyses but, due to the size of the grains and the spatial resolution of electron beam, Ni concentrations were not quantified. A close association between pyrite and organic matter is shown in samples LWC31, LWC39, LWC41 and LWC88 (Fig. 5d). All Raman spectra obtained on these laminae show characteristics of poorly ordered carbonaceous material (Fig. S4), corresponding to maximum temperature of 150°C consistent with burial diagenesis (Heydari and Moore, 1989).

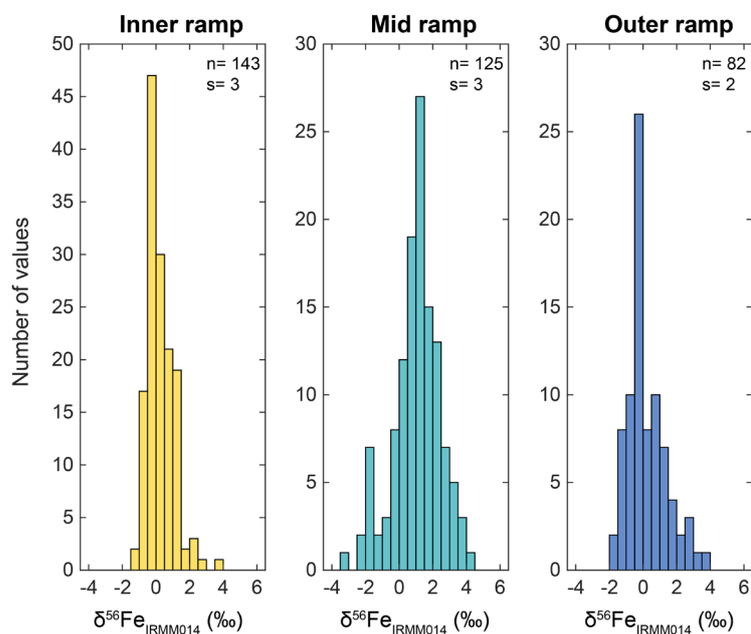
3.5. In situ iron isotope compositions



Corrected Figure 6: Fe isotope compositions of pyrite from the Lower Weber Canyon section as a function of the depositional environment along a ramp system. The different star colors refer to the corresponding facies associations. Sample LWC31 is part of FA1 (red star). Samples LWC39 and LWC41 are MISS (FA2, yellow star). Please note that sample LWC41 encloses two facies in which pyrite is observed. The first is the silty part (reported as LWC41 Siltst.) and the second is the OM-rich laminae (noted LWC41 OM). F3 (green star) includes samples LWC47 and LWC49. Sample LWC55 is part of FA4 (blue star). Samples LWC72 and LWC88 are included in FA5 (purple star). Fe isotope compositions are reported using outlier box plot representation. Line through the box is the median and the edges of the box represent the quartiles (lower line is the 1st quartile Q1, i.e. the 25th percentile, and upper line is the 3rd quartile Q3, i.e. the 75th percentile). Lower and upper whiskers represent 1.5*IQR (interquartile range, i.e. the range between Q1 and Q3). The minimum and maximum are marked by small lines at the end of whiskers and represent respectively the lowest and highest values of the distribution, excluding outlier values of the dataset (filled and unfilled dots below and above whiskers). See Fig. 1 for abbreviations.

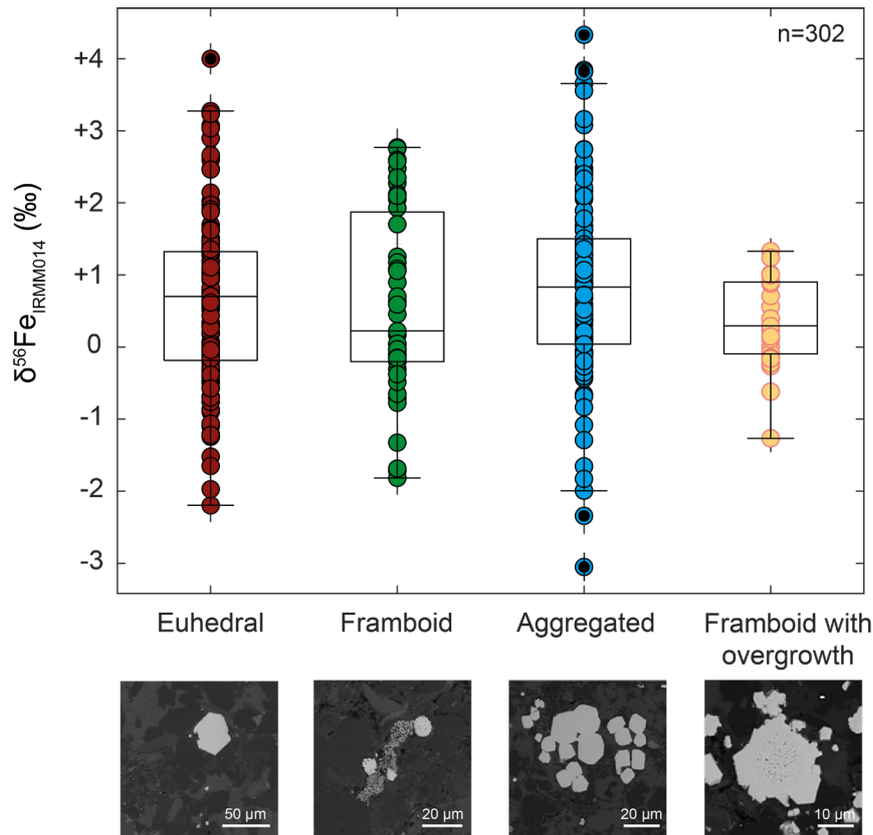
Large variations of pyrite $\delta^{56}\text{Fe}$ values, from -3.05 to +4.33 ‰, are evidenced over 350 analyses through the studied section (**corrected Fig. 3; corrected Table S9**). In the inner ramp deposits (FA1 and FA2), 143 pyrite analyses show $\delta^{56}\text{Fe}$ values ranging from -1.27 to +3.66 ‰, and an average $\delta^{56}\text{Fe}_{\text{py}}$ value of +0.26 ‰ (**corrected Fig. 6**). In the mid ramp sediments (FA3 and FA4), the 125 $\delta^{56}\text{Fe}_{\text{py}}$ values display the largest range from -3.05 to +4.33 ‰, and an average $\delta^{56}\text{Fe}_{\text{py}}$ value of +1.04 ‰ (**corrected Fig. 6**). Samples from inner and mid ramp systems display monomodal distributions (**corrected Fig. 7**) for which median values are statistically different (Mann-Whitney $p=1.69\text{E}-10$; **Table S4**). In the outer ramp deposits (FA5), $\delta^{56}\text{Fe}_{\text{py}}$ values ($n=82$) range from -1.65 to +3.82 ‰, with an average $\delta^{56}\text{Fe}_{\text{py}}$ value of +0.22 ‰ (**corrected Fig. 6**).

In details, the inner ramp FA1 includes samples LWC31 ($n=77$) and LWC41 Siltst. ($n=13$). The Fe isotope compositions range from -0.78 to +1.21 ‰ in LWC31, and from -1.27 to +0.44 ‰, in LWC41 Siltst. (**corrected Fig. 6**) and show similar distribution with the respective mean $\delta^{56}\text{Fe}_{\text{py}}$ values of -0.15 ‰ and -0.22 ‰ (Mann-Whitney $p=0.662$; **Table S5**). The inner ramp FA2 includes samples LWC39 ($n=30$) and LWC41 OM ($n=23$) with higher $\delta^{56}\text{Fe}$ values, varying respectively from -0.05 to +3.66 ‰ and from -0.08 to +1.33 ‰. These two samples display statistically different mean $\delta^{56}\text{Fe}$ values of +1.22 ‰ and +0.65 ‰, respectively (T test $p=2.7\text{E}-3$; **Table S5**). Fe isotope compositions of mid ramp FA3, including samples LWC47 ($n=25$) and LWC49 ($n=52$), range from -3.05 to +2.77 ‰ and from -1.69 to +4.33 ‰. They display statistically different $\delta^{56}\text{Fe}_{\text{mean}}$ values of -0.60 ‰ and +1.24 ‰, respectively (T test $p=8.01\text{E}-9$; **Table S5**). Mid ramp FA4 corresponds to sample LWC55 ($n=48$), which shows $\delta^{56}\text{Fe}$ values ranging from -0.13 to +4.00 ‰, with the highest average $\delta^{56}\text{Fe}$ value of +1.67 ‰. Outer ramp FA5 is represented by samples LWC72 ($n=35$) and LWC88 ($n=47$). Pyrite grains in LWC72 have Fe isotope compositions ranging from -1.52 to +3.82 ‰. In LWC88, pyrite $\delta^{56}\text{Fe}$ values vary between -1.65 and +2.46 ‰. These two samples show similar distribution with the respective mean $\delta^{56}\text{Fe}$ values of +0.49 ‰ and +0.02 ‰ (Mann-Whitney $p=0.249$; **Table S5**). No correlation between $\delta^{56}\text{Fe}$ values and S content or between $\delta^{56}\text{Fe}$ and $\delta^{34}\text{S}$ values has been observed (**corrected Fig. S5**).



Corrected Figure 7: Pyrite $\delta^{56}\text{Fe}$ distributions as a function of the domain of the ramp, i.e. the inner ramp, the mid ramp and the outer ramp system. The number of values (n), and the number of samples from which pyrite grains were measured (s) are indicated for each histogram.

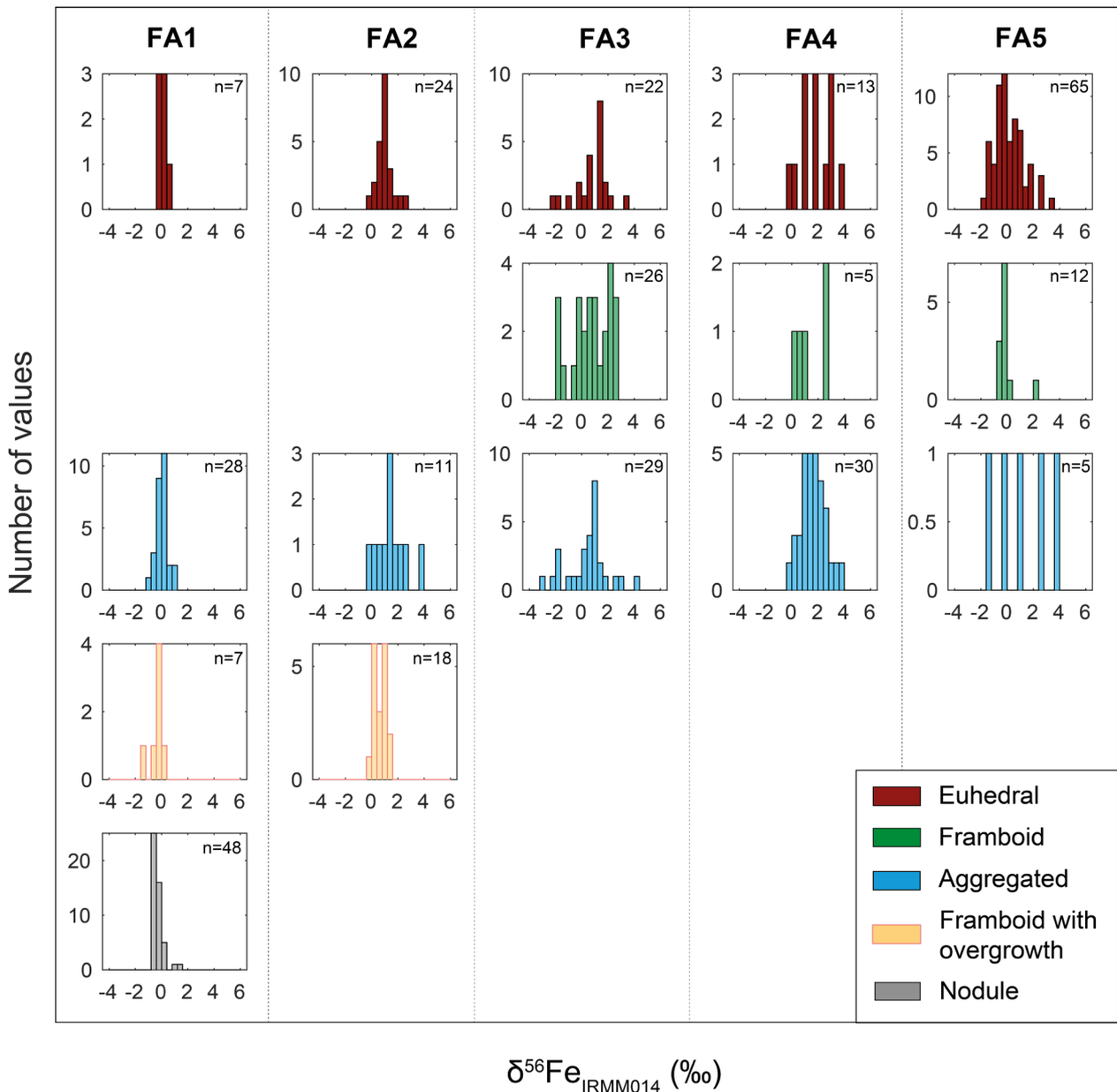
3.6. Iron isotope signatures, facies and pyrite morphologies



Corrected Figure 8: $\delta^{56}\text{Fe}_{\text{py}}$ as a function of the morphology of pyrite grains. The different pyrite types are categorized as euhedral, framboids, aggregated and framboids with secondary overgrowth. No statistical differences are observed between each morphology.

The Fe isotope variations can be represented as a function of the four pyrite morphologies (excepting the nodule from LWC31; **corrected Fig. 8**). The $\delta^{56}\text{Fe}$ values of euhedral grains range between -2.20 to +4.00 ‰ (n=131), with an average value of +0.60 ‰. Framboidal pyrite have a lower range of variations, with $\delta^{56}\text{Fe}$ values from -1.82 to +2.77 ‰ (n=43) and a mean $\delta^{56}\text{Fe}$ value of +0.59 ‰. Aggregated pyrite grains show an Fe isotope range between -3.05 and +4.33 ‰ (n=103) and an average $\delta^{56}\text{Fe}$ value of +0.83 ‰. The Fe isotope compositions of framboidal grains with secondary overgrowth range from -1.27 to +1.33 ‰ (n=25). These grains display an average $\delta^{56}\text{Fe}$ value of +0.35 ‰. These four pyrite morphologies describe monomodal Gaussian-type distributions (**corrected Fig. S6**) and the differences of mean $\delta^{56}\text{Fe}$ values are not statistically relevant (T test p value= [0.082-0.929]; **Table S6**). Moreover, there is no significant differences in the mean $\delta^{56}\text{Fe}$ values between the core $\delta^{56}\text{Fe}_{\text{mean}} = +0.32 \pm 0.57$ ‰ and the rim ($\delta^{56}\text{Fe}_{\text{mean}} = +0.42 \pm 0.76$ ‰) among the framboid with secondary overgrowth pyrite grains (Mann-Whitney p= 0.432). The observation of these 4 different pyrite types depends on the associated facies (**corrected Fig. 9**):

- In FA1 (siltstones), pyrite grains are euhedral, aggregated, framboidal (with diagenetic overgrowth) or forming a nodule (**corrected Fig. 9**). Euhedral pyrite grains have a mean $\delta^{56}\text{Fe}$ value of $+0.14 \pm 0.31$ ‰ (n=7). The aggregated grains display an average $\delta^{56}\text{Fe}$ value of $+0.03 \pm 0.44$ ‰ (n=28). The distributions observed for euhedral and aggregated pyrite in FA1 are monomodal and similar (Mann-Whitney p=0.695; **Table S7**). Framboids with secondary overgrowth and the pyrite nodule show the respective mean $\delta^{56}\text{Fe}$ values of -0.37 ± 0.45 ‰ (n=7) and -0.29 ± 0.40 ‰ (n=48). These two morphologies display monomodal and similar distributions (Mann-Whitney p=0.640;



Corrected Figure 9: Pyrite $\delta^{56}\text{Fe}$ distributions as a function of pyrite morphologies in each facies (FA1 to FA5). Data from euhedral pyrite are reported in red, from framboids in green, from aggregates in blue, from framboids with secondary rims in yellow and from the nodule in grey. The number of values (n) is indicated for each histogram.

Table S7). Moreover, the mean values observed in framboids with overgrowth and the nodule are lower than those observed in euhedral and aggregated grains. However, the difference in means between these two groups is covered by the uncertainties of the analyses (**corrected Fig. 9**).

- Excluding the nodule, which has only been observed in FA1, the same pyrite morphologies are reported in FA2 (MISS; **corrected Fig. 9**). Euhedral grains in FA2 show a monomodal distribution and a mean $\delta^{56}\text{Fe}$ value of $+1.02 \pm 0.60 \text{‰}$ ($n=24$). In the aggregated grains, the $\delta^{56}\text{Fe}$ values are distributed along a monomodal distribution on a larger range than euhedral grains. The average value is $+1.44 \pm 1.05 \text{‰}$ ($n=11$). However, the difference between euhedral and aggregated grains is not statistically significant (T test $p=0.137$; **Table S7**). Framboidal pyrite with secondary overgrowth also displays a monomodal distribution, which is characterized by the lowest mean $\delta^{56}\text{Fe}$ value of $+0.63 \pm 0.43 \text{‰}$ ($n=18$) among the other pyrite morphologies observed in FA2. However, once again

the difference in means between the recrystallized framboids and the other pyrite types falls into the uncertainties of the analyses.

- The overgrowth affecting framboidal pyrite are no longer observed in FA3 (bioclastic packstones) and the other following facies (**corrected Fig. 9**). Euhedral grains in FA3 have an average $\delta^{56}\text{Fe}$ value of $+0.79 \pm 1.30 \text{ ‰}$ ($n=22$), which is similar to the Fe isotope signature of framboidal grains ($+0.72 \pm 1.42 \text{ ‰}$, $n=26$). Aggregated grains have a slightly lower mean $\delta^{56}\text{Fe}$ value of $+0.45 \pm 1.59 \text{ ‰}$ ($n=29$). The distributions of these 3 pyrite types spread over a larger range than those of FA1 and FA2 and are similar (Mann-Whitney $p= [0.168-0.828]$; **Table S7**).

- The three pyrite types described in FA3 are also observed in the following facies FA4 and FA5 (**corrected Fig. 9**). Only one sample (LWC55) is part of FA4 (bivalve wackestones), in which pyrite grains record the highest Fe isotope compositions. Euhedral pyrite shows a sparse distribution with a mean $\delta^{56}\text{Fe}$ value of $+1.86 \pm 1.22 \text{ ‰}$ ($n=13$). Five framboids were measured and display a mean $\delta^{56}\text{Fe}$ value of $+1.41 \pm 1.12 \text{ ‰}$. Aggregated grains show a monomodal distribution with an average Fe isotope composition of $+1.64 \pm 0.97 \text{ ‰}$ ($n=30$). The isotopic differences between these three pyrite types are statistically insignificant (T test $p= [0.528-0.633]$; **Table S7**).

- Samples from FA5 (mudstones) enclose euhedral pyrite grains, which describe an asymmetric distribution with a mean $\delta^{56}\text{Fe}$ value of $+0.19 \pm 1.08 \text{ ‰}$ ($n=65$), framboids, characterized by a lower mean $\delta^{56}\text{Fe}$ value of $-0.05 \pm 0.80 \text{ ‰}$ ($n=12$), and a few aggregated grains displaying a $\delta^{56}\text{Fe}$ value of $+1.23 \pm 2.09 \text{ ‰}$ ($n=5$; **corrected Fig. 9**). Yet, there is no significant difference between these distributions (Mann-Whitney $p= [0.269-0.523]$; **Table S7**).

Although some pyrite exhibits secondary features, i.e. recrystallization (**Fig. 5a and b**), their iron isotope compositions remain similar (**corrected Fig. 9**). For example, there is no profile of variation of the $\delta^{56}\text{Fe}$ values in the framboidal pyrite with overgrowths. Overall, there is no clear difference of Fe isotope compositions between the pyrite types observed among each facies.

4. Discussion

4.1. Diagenetic and syngenetic pyrite grains

Pyrite in three different facies (FA3, FA4, FA5) exhibits preserved primary morphological features such as framboids, characterized by an average size of $11 \pm 5.0 \mu\text{m}$ (**corrected Fig. 4**). The study of *Rickard (2019)*, in line with the pioneer work of *Wilkin and Barnes (1997)*, documented that diagenetic framboids yield a mean diameter of $6.7 \mu\text{m}$, meaning that framboids from our samples most likely precipitated in the sediment porewaters. As framboids are closely associated with well-preserved carbonaceous materials, we suggest that the Fe isotope compositions of those grains are inherited from early diagenetic processes. The Fe isotope compositions (ranges and mean values) of these framboids is similar to the Fe isotope compositions of euhedral and aggregated grains within the same facies. Therefore, we can reasonably hypothesize that framboids, euhedral and aggregated

pyrite formed in the sediments through the same precipitation event.

The presence of overgrowth around framboidal cores in FA1 and FA2 samples (LWC31, LWC39 and LWC41) marks a subsequent event of precipitation. Supposedly, pyrite resulting from secondary fluid circulation should display contrasting $\delta^{56}\text{Fe}$ values or different mineralogical features. However, the Fe isotope compositions of overgrowth are similar to those of framboidal cores. Based on the fact that (1) there is also no significant isotopic difference between rimmed framboidal pyrite and the other pyrite types (i.e. euhedral or aggregated pyrite) observed within the same facies (FA1 or FA2), and (2) in FA2 sample LWC41, an isotopic heterogeneity is preserved between pyrites from the silty sediments (LWC41 Siltst.) and pyrite from the organic laminae (LWC41 OM), we suggest that this secondary fluid circulation did not affect the Fe isotope compositions of the initial pyrite grains (framboid dissolution and recrystallization without additional Fe supply, i.e. low fluid/mineral ratio).

The influence of late diagenetic processes is illustrated by the isotope signatures of CAS. The comparison between the Fe and S isotope compositions of pyrite and the S isotope composition of CAS reveals that the variation of $\delta^{56}\text{Fe}$ and $\delta^{34}\text{S}_{\text{py}}$ values does not match with the S isotopic value of sulfate reservoir ($\delta^{34}\text{S}_{\text{CAS}}$; **corrected Fig. 3**). The decoupling between pyrite and CAS means that the isotopic signatures of pyrite are not related to the evolution of the sulfate reservoir from which carbonate precipitate. This decoupling raises concerns about secondary overprint that could have affected CAS (Swart, 2015). In this study, $\delta^{34}\text{S}_{\text{CAS}}$ values were measured in samples characterized by Mn/Sr ratios between 11 and 40. Due to the decrease of Sr concentrations in carbonates during late/burial diagenetic processes, the Mn/Sr ratio is widely used to assess the degree of carbonate recrystallization. Consequently, samples showing high Mn/Sr ratio (> 2) reflect secondary alteration (Kaufman and Knoll, 1995; Derry, 2010). Here, the analyzed samples have minimum Mn/Sr ratio of 11 (**Table S1**), ratio showing an extensive carbonate recrystallization. Consequently, the $\delta^{34}\text{S}_{\text{CAS}}$ signal is likely inherited from late diagenetic processes.

Although late diagenetic processes occurred in the LWC samples, the association of pyrite with preserved organic materials and Fe isotope measurements indicate that most of the facies (FA1, FA3, FA4 and FA5) contain pyrite of early diagenetic origin. In FA2 (MISS), the presence of framboid remains within organic-rich laminae is suggestive of the formation of pyrite when biofilms were the site of metabolic activities, i.e. framboids of syndimentary origin. Those pyrite grains have been partly recrystallized (i.e. overgrowth around framboids and euhedral grains) but still likely preserve their original Fe isotope compositions. Therefore, we propose that samples of the LWC section contain two generations of pyrite. The first generation represents syndimentary pyrite associated with MISS (FA2), which formed during the activity of the biofilm close to the water column/sediment interface. The second generation is represented by early diagenetic pyrite (FA1, FA3, FA4 and FA5), whose formation is still controlled by a microbial activity spreading over the upper tens of centimeters of the sediments. Thus, syndimentary to early diagenetic processes likely largely controlled Fe isotopic signatures of pyrites in the studied samples during the late Smithian and across the SSB.

4.2. Iron isotope kinetic and equilibrium isotope effects

Pyrite precipitation associated with various degrees of kinetic and equilibrium fractionation is a possible hypothesis to explain large Fe isotope range of isotopic composition (*Mansor and Fantle, 2019*). This hypothesis links Fe/S ratio changes with highly variable Fe pyrite isotope signatures. Indeed, an elevated Fe/S ratio enhances the expression of kinetic fractionation, i.e. formation of pyrite with negative $\delta^{56}\text{Fe}$ values, in a sulfate (and thus sulfide) poor environment. In contrast, pyrite expressing the equilibrium isotope fractionation, i.e. positive $\delta^{56}\text{Fe}$ values, reflects low Fe/S ratio and thus an environment with higher sulfate concentrations. Therefore, the increase of $\delta^{56}\text{Fe}$ values through the SSB can reflect variations in the supply of sulfate to the ocean, for example by volcanic degassing as already suggested during the SSB interval (*Bond and Grasby, 2017*). In this latter case, the absence of high Hg concentrations through the SSB does not support sulfate concentration variations driven by volcanism. Finally, although not entirely excluded, the absence of covariation between the sulfur content and pyrite $\delta^{56}\text{Fe}$ values or between pyrite $\delta^{56}\text{Fe}$ and $\delta^{34}\text{S}$ values rather suggests that sulfur availability during pyritization did not primarily control the $\delta^{56}\text{Fe}$ signal observed in the studied samples through the SSB.

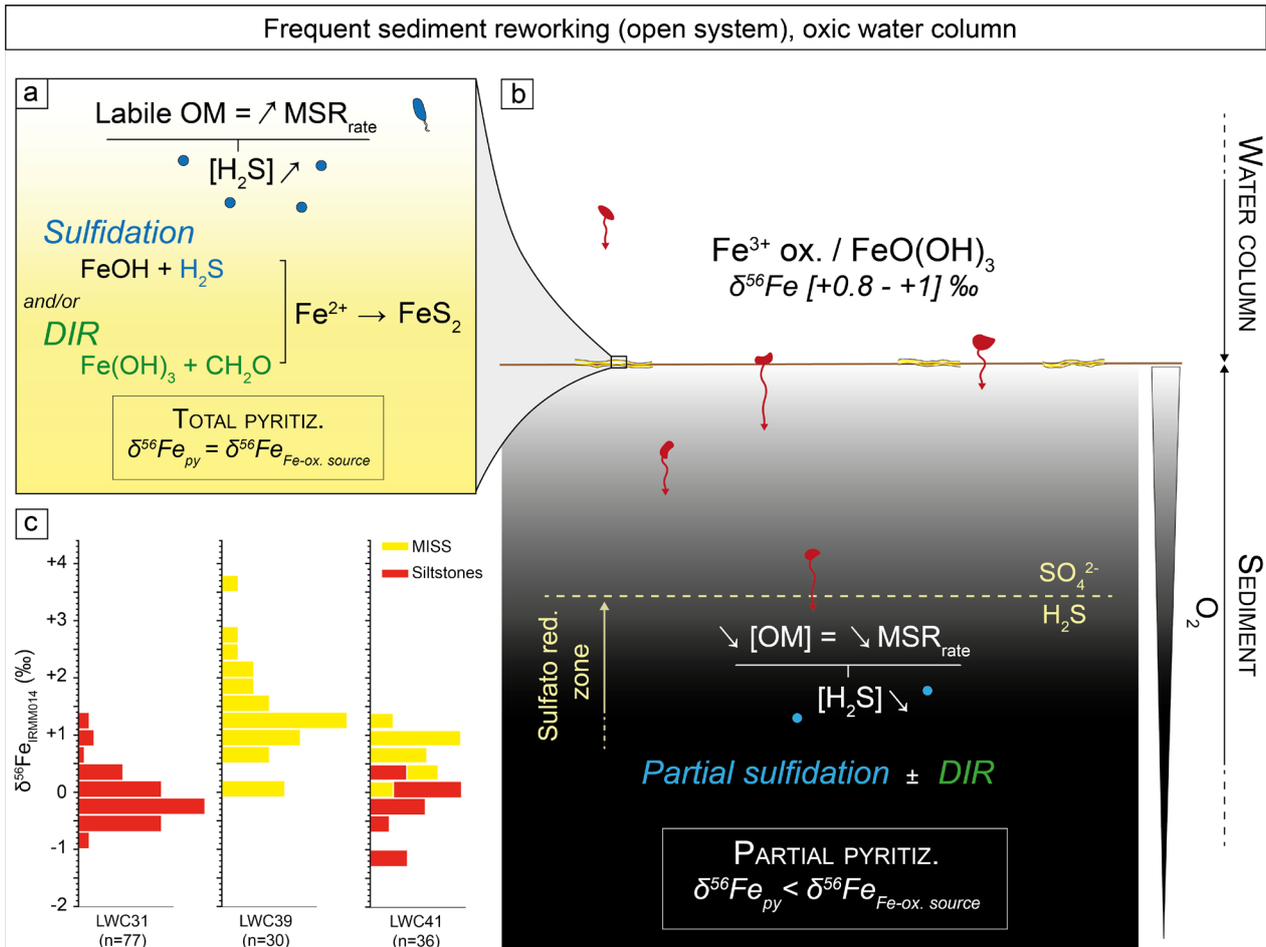
4.3. Pyrite $\delta^{56}\text{Fe}$ values controlled by sedimentary depositional settings and microbial communities

We suggest that $\delta^{56}\text{Fe}$ variability is driven by redox cycling proceeding within the sediment, including biotic Fe-oxide reduction by microorganisms, a process named dissimilatory iron reduction (DIR), and reductive dissolution of Fe-oxides by sulfides. Pyrite Fe isotope signatures may also partly reflect kinetic and equilibrium isotope effect as the reductive dissolution of Fe-oxides likely change, at a local scale, the Fe/S ratio. The petrographic observations (**Fig. 1**), $\delta^{34}\text{S}_{\text{py}}$ values (**corrected Fig. 3**) and $\delta^{56}\text{Fe}$ values (**corrected Fig. 7**) allow to identify two endmembers. Both endmembers are largely dependent of the degree of connectivity of the sediment porewaters with the overlying water column, this latter likely being fully oxygenated as suggested by the relatively low S content (mean S content= 2585 ppm) and a low TOC content (mean TOC= 1477 ppm), which are not expected under euxinic or anoxic conditions (*Lyons et al., 2009* for a review). Endmember 1 is depicted in samples from the inner ramp system. In these samples, the negative $\delta^{34}\text{S}_{\text{py}}$ (**corrected Fig. 3**) and the disturbed OM-rich laminae (**Fig. S7**) suggest a good ventilation of the sediments, leading to a constant replenishment of the S and Fe reservoirs. However, this parameter alone does not explain the variability and the significant difference in mean $\delta^{56}\text{Fe}$ values between MISS and siltstones. These differences can be explained by the variation in the amount of microbially-produced H_2S , resulting from different microbial sulfate reduction (MSR) rates in MISS and siltstones. Endmember 2 is illustrated by samples from the mid and outer ramp system. In these samples, $\delta^{34}\text{S}_{\text{py}}$ are positive (**corrected Fig. 3**) and the position along the ramp suggests that the sediments are dominated by a relatively low energy regime. The increasing trend described by both $\delta^{34}\text{S}_{\text{py}}$ and $\delta^{56}\text{Fe}$ suggest that pyrite records diagenetic processes occurring under closed system in the sediment porewaters.

Therefore, two models are proposed below to account for the $\delta^{56}\text{Fe}$ variations observed within the LWC ramp system deposits: (1) an “inner ramp” model, functioning in an open system (i.e. frequent sediment reworking leading to a good connection between the sediment porewaters and the overlying water column) and (2) a “mid and outer ramp” model, operating in a closed system (i.e. rare sediment reworking leading to a disconnection between the sediment porewaters and the overlying water column) and allowing the buildup of Fe-oxide pools within the sediments.

4.3.1. Inner ramp model: $\delta^{56}\text{Fe}$ values controlled by the nature of deposit

The LWC inner ramp is characterized by frequent sediment remobilization by physical reworking (tidal currents or waves) and bioturbation, which facilitated the connection between sediment porewaters and the overlying water column. Therefore, sulfates, Fe-oxides and organic matter that fuel metabolic reactions related to sedimentary sulfides precipitation are assumed to be non-limiting. An open system is consistent with the average negative $\delta^{34}\text{S}$ value recorded in the inner ramp samples ($\delta^{34}\text{S} = -11.4 \pm 9.3 \text{‰}$, 1SD) and specifically in samples LWC31, LWC39 and LWC41, with values lower than -16.8‰ . The model described in the original version of this article may still explain the variability of Fe isotope compositions in FA1 and FA2 samples of the inner ramp system, assuming that the $\delta^{56}\text{Fe}$ value of the Fe source is close to 0‰ . Such iron isotopic signal can be derived, for example, from quantitative iron oxidation into Fe^{3+} -minerals after weathering of igneous rocks, aerosols, or river loads (*Dauphas and Rouxel, 2006* for a review). However, the Fe isotope compositions in FA1 and FA2 samples may also be alternatively interpreted, assuming that the $\delta^{56}\text{Fe}$ value of the initial Fe oxides is close to the canonical Fe isotope composition of $+0.9 \text{‰}$ known for Fe oxides throughout Earth history (*Rouxel et al., 2005; Wu et al., 2011; corrected Fig. 10*). In that case, the Fe isotope variability measured in pyrite grains from the inner ramp would be primarily controlled by the availability of microbially-produced H_2S (**corrected Fig. 10**). Through this alternative model, the average $\delta^{56}\text{Fe}$ value of $+0.97 \text{‰}$ (1SD) associated with FA2 (MISS) samples would reflect the complete sulfidation of Fe-oxides, allowing the preservation of the initial $\delta^{56}\text{Fe}$ value of Fe-oxides within pyrite (**corrected Fig. 10a and c**). This process can be supported by H_2S production in biofilms after microbial sulfate reduction (MSR). Consequently, in sediments devoid of biofilms (FA1, siltstones), the sulfidation of Fe-oxides may be limited by lower production rates of H_2S by sulfate reducers, related to a lower reactivity of organic compounds (**corrected Fig. 10b**). Indeed, under well-ventilated conditions, MSR occurs deeper in the sediments where the quantity and reactivity of organic matter decrease as a function of the sediment depth and redox potential (*Middelburg, 1989; Dauwe et al., 1999; Meister et al., 2013*). Although iron reducers may have been involved in the reduction of Fe oxides, the newly corrected data are still not sufficiently negative to support dissimilatory iron reduction (DIR) as the dominant reduction pathway (*Crosby et al., 2005; corrected Fig. 10c*). Instead of a model requiring several oxidation-reduction cycles to explain pyrite $\delta^{56}\text{Fe}$ values as originally exposed, this new model proposes that the key driver of pyrite formation is the sulfidation of Fe oxides by microbial H_2S , the production of which depends on the sedimentary facies.



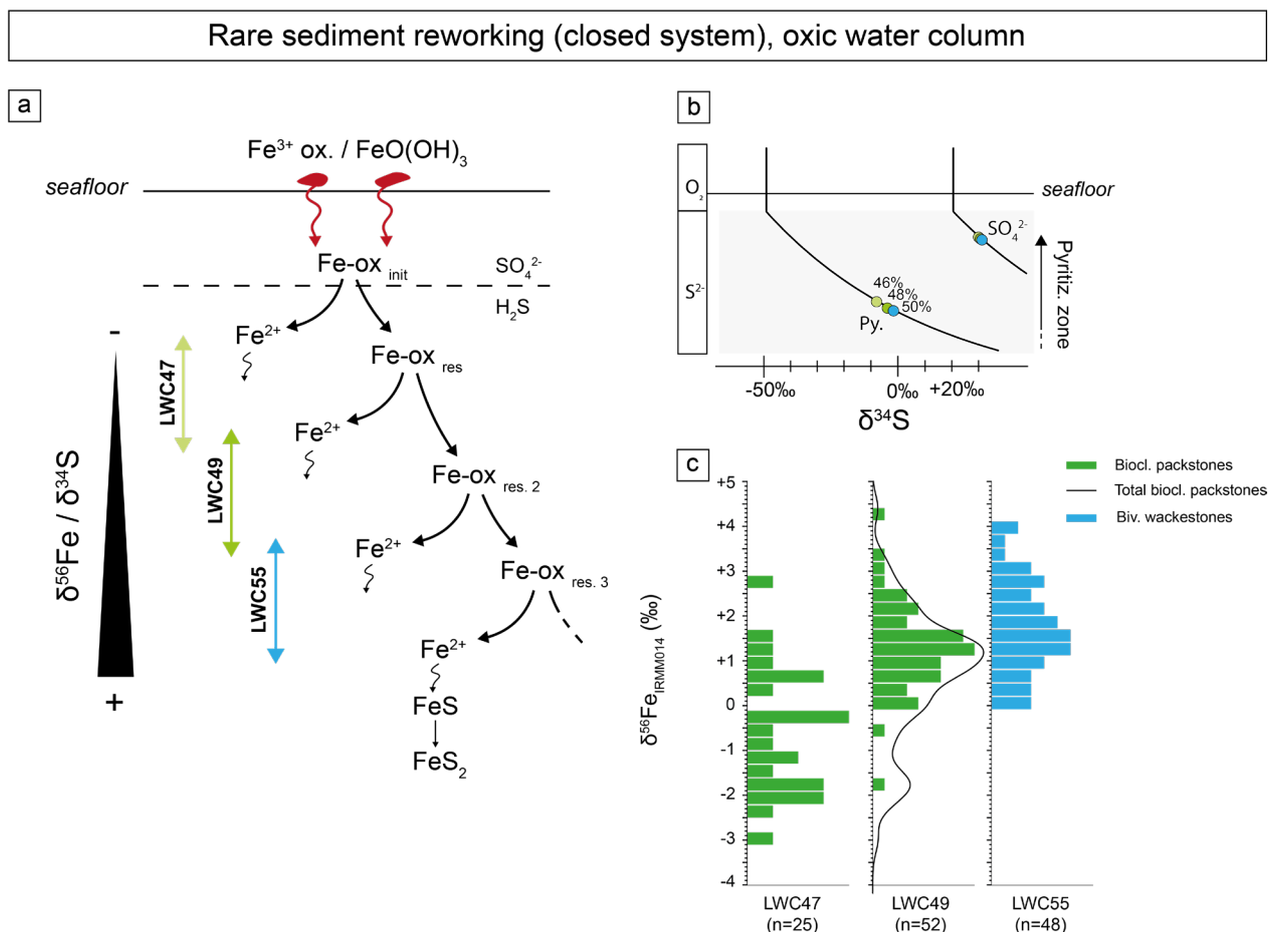
Corrected Figure 10: Iron cycling model in the sediments from inner ramp samples. a) Model of Fe reduction for MISS samples (FA2), where high rate of sulfate reduction promotes the complete conversion of Fe-oxides into Fe²⁺ or FeS and subsequently pyrite by reaction with H₂S (blue pathway: Sulfidation) combined with the possible effect of iron reducing bacteria (green pathway: Dissimilatory Iron Reduction). b) Model of Fe reduction for siltstone samples (FA1). In this model, less abundant and less labile organic matter is available for sulfate reducers, leading to a lower production of H₂S. In this case, the quantity of H₂S limits the reduction of Fe-oxides. c) Pyrite $\delta^{56}Fe_{py}$ distributions for the facies associations FA1 (siltstone) and FA2 (MISS).

4.3.2. Mid and outer ramp model: $\delta^{56}Fe_{py}$ controlled by H₂S production in a closed system

Samples from the mid ramp system record a progressive increase in average $\delta^{56}Fe$ values, from -0.60 to +1.67 ‰, that is also recorded in S isotope compositions with an increase from -8.2 to -2.8 ‰ (corrected Fig. 3). A concurrent increase of $\delta^{56}Fe$ and $\delta^{34}S$ values for mid ramp samples can reflect a consumption of the Fe-oxide and sulfate reservoirs within the sediments under closed system conditions relative to the overlying water column (corrected Fig. 11). Samples LWC47 and LWC49 show disarticulated bioclasts of bivalves and ammonoids, reflecting rare storm events and limited transport (Seilacher et al., 1985; Kidwell et al., 1986; Grosjean et al., 2018). LWC55 also shows fragmented bivalves and thin layers of silty beds (Fig. 1), but the higher proportion of micritic matrix compared to LWC47 and LWC49 suggests a slightly deeper depositional environment (Fig. 1). In such environments, episodic storms resulted in rapid sediment accumulations that promoted disconnections between porewaters and the water column. In this environmental context replenishment of solute oxidized species is restricted (corrected Fig. 11a). Consequently, a closed

system sulfate reduction developed and may have driven the partial reduction of Fe-oxides, leading to the buildup of a residual Fe-oxide pools in the sediments that are progressively ^{56}Fe -enriched (**corrected Fig. 11a and b**). The large range and non-Rayleigh-type distribution of $\delta^{56}\text{Fe}$ values in LWC47 (**corrected Fig. 11c**) support that (1) pyrite recorded multiple steps of partial reduction of a residual Fe-oxide pool, and (2) a possible recycling (reoxidation-reduction) of the produced Fe^{2+} .

The average $\delta^{56}\text{Fe}$ value of outer ramp samples LWC72 and LWC88 is of $+0.22\text{‰}$, i.e. 0.82‰ lower than that of the mid ramp samples (mean $\delta^{56}\text{Fe}_{\text{Mid}} = +1.04\text{‰}$). The large variability of $\pm 1.14\text{‰}$ observed in these outer ramp samples is not distributed along a Rayleigh-type process. In contrast, S isotope signal increases, from -4.0‰ in the mid ramp to $+6.4\text{‰}$ in the outer ramp setting. LWC72 and LWC88 pertained to FA5 (**corrected Fig. 6**), which is dominated by sediment decantation and very low hydrodynamic regime. Deposition of FA5 occurred in a more distal position along the ramp where sediment reworking is very rare, meaning that the sediments likely evolved under closed system conditions. In such setting, sulfides generated in sediments by sulfate reduction may promote the partial reduction of reactive Fe-oxides (**corrected Fig. 11**) in outer ramp sediments.



Corrected Figure 11: Iron cycling model in the sediments from mid and outer ramp samples. a) Model for Fe-oxide reduction in closed-system, where Fe-oxides are partially reduced by reaction with a restricted amount of H_2S (as sulfate reservoir also evolves in a closed-system), leading to the buildup of different residual Fe-oxide pools. b) Evolution of newly formed pyrite and sulfate in a closed sediment. Percentage of newly formed pyrite during the Rayleigh process is reported for LWC 47 (light green), LWC49 (green) and LWC55 (blue). c) Pyrite $\delta^{56}\text{Fe}$ distributions for samples from FA3 and FA4. Green and blue colors correspond to facies associations FA3 (bioclastic packstones) and FA4 (bivalve wackestones) respectively. Black line represents probability density for both LWC47 and LWC49.

This partial reduction produces Fe^{2+} that can be then converted into pyrite. Due to the activity of sulfate reducing metabolisms and the progressive buildup of residual Fe-oxide pools from an initial reservoir, reactive Fe-oxides tend to be rare. Therefore, sulfides produced deeper in the sediments are not consumed in situ by reaction with Fe^{2+} and accumulate, leading to a diffusion upward its zone of production (Riedinger *et al.*, 2017; Liu *et al.*, 2020). Hypothetically, this diffusing sulfide flux may be generated at the sulfate-methane transition zone (SMTZ), as recent studies documented the precipitation of ^{34}S -enriched pyrite in this deeper area in sediments (Liu *et al.*, 2020; Pasquier *et al.*, 2021). However, the data presented here are not conclusive evidence for the existence of a paleo-SMTZ in this ramp system. This diffusion process results in the successive sulfidation of the (more) reactive residual and initial Fe-oxides pools (**corrected Fig. 11**), converting them to pyrite characterized by ^{34}S -enriched S isotope signatures and large $\delta^{56}\text{Fe}$ range of values. The mean $\delta^{56}\text{Fe}$ values of FA5 samples ($\delta^{56}\text{Fe}_{\text{mean}} = +0.22 \pm 1.14 \text{‰}$) further suggests a partial sulfidation of Fe-oxides, which is consistent with sporadic H_2S diffusion flux.

4.4. Pyrite formation during an interval of biotic diversity loss: a depositional environment control rather the deleterious anoxic conditions

The oceanic redox conditions through the SSB were often documented by studying pyrite. Indeed, many studies focused on the occurrence and size distribution of pyrite framboids, Fe speciation and multiple sulfur isotope signals to assess euxinia, anoxia or ferruginous conditions of the water column (e.g. Shen *et al.*, 2011; Sun *et al.*, 2015; Clarkson *et al.*, 2016; Song *et al.*, 2019; Dai *et al.*, 2021). However, there have been no studies using iron isotopes to document this biotic crisis, although iron is also a redox-sensitive element. Here we evidence that these depositional environments and more specifically the degree of sediment connection with the water column, which is modulated by sediment accumulation and reworking, controlled the S and Fe isotopic compositions through variations of Fe-oxide and sulfate supply in the sediment microbial active zones. When sediments are well-connected to the water column, pyrite displays negative $\delta^{34}\text{S}$ values and their Fe isotope compositions reflect the availability of H_2S supplied by the activity of sulfate reducers. On the other hand, the disconnection between the water column and the sediments leads to the isotopic consumption of sulfate and Fe-oxide reservoirs.

The control of the sediment-seawater connectivity between each depositional environment is well evidenced by the S isotope signatures, while Fe isotopes show the additional influence of the nature of deposit (presence of MISS) and the reactivity of Fe-oxides with H_2S fluxes. This suggests that both Fe and S isotopic variations through the SSB at LWC are driven by local processes within the sediments rather than the water column redox structure. In this study, no direct causality can be established between Fe isotopic signal and interval of biotic crisis, as already shown through the SSB using multiple sulfur isotope signals (Thomazo *et al.*, 2019). Finally, the combination of Fe isotope compositions along with S isotopes and petrographic observations is consistent with local microbial and sedimentological processes as primary drivers of sedimentary geochemical signals within the studied biotic diversity loss interval.

5. Conclusions

Pyrite Fe isotope compositions obtained from LWC through the Smithian-Spathian transition indicate a sedimentary depositional system control, locally coupled with an influence of the nature of deposits (i.e. the presence or absence of MISS) on pyrite formation. LWC sediments deposited along a tide and wave influenced ramp system, ranging from shallow inner ramp to deeper outer ramp settings, was overlaid by an oxygenated water column. In these environments, the intensity of biological and physical induced sediment reworking produces two regimes of water column-sediment exchanges, an open-system characterized by well-ventilated sediments and a closed-system govern by diffusion within the porewater sediment. We propose two iron cycling models both influenced by the position along the ramp and nature of the deposits. The first model is relevant to reworked sediments from inner ramp settings, where the $\delta^{56}\text{Fe}_{\text{py}}$ values ($\delta^{56}\text{Fe}_{\text{mean}} \sim +1\text{‰}$) in MISS reflect the complete reduction of Fe-oxides and the lighter $\delta^{56}\text{Fe}_{\text{py}}$ ($\delta^{56}\text{Fe}_{\text{mean}} \sim 0\text{‰}$) in siltstones are attributed to the partial sulfidation of Fe-oxides. This partial reduction results from the restricted production of microbial H_2S that occurred deeper in the silty sediments compared to MISS. The second model is relevant for sediments deposited in the mid and outer ramp zones that were rarely affected by reworking. The concurrent increase of Fe and S isotope compositions suggests that the disconnection between the sediment porewaters and water column drives the consumption of the Fe-oxide and sulfate sedimentary reservoirs.

This study highlights the importance of using different isotope systematics the sedimentary depositional/nature of sediments to unravel local versus global control on geochemical signatures and call for caution in the “redox” interpretation of Fe and S isotopic signals, especially digging into geological intervals associated with extensive oceanic anoxia.

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Corrected supplementary materials

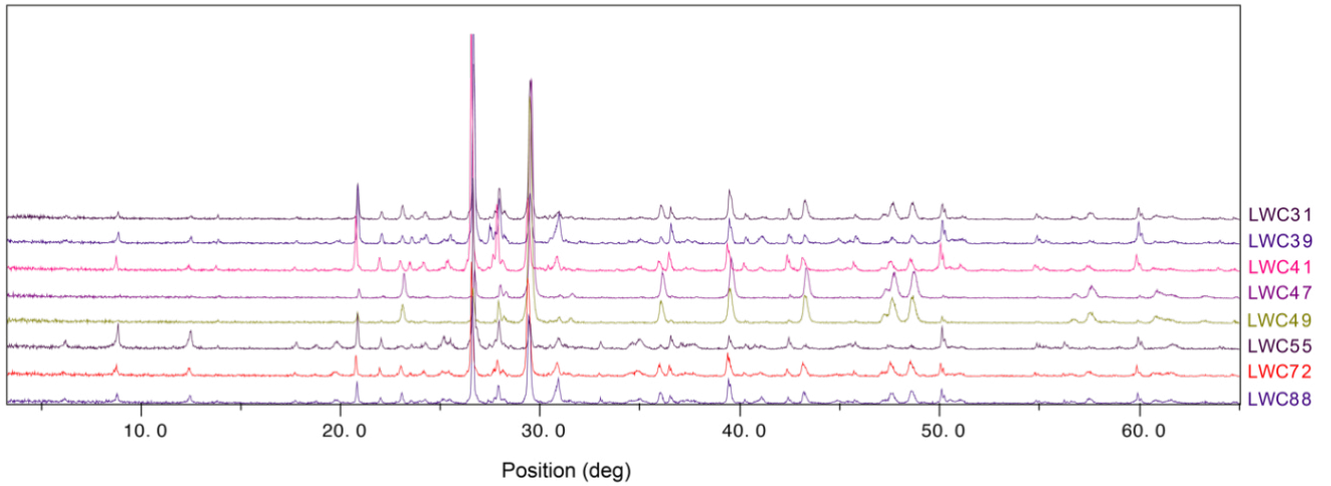


Figure S1: XRD analyses on samples from the Lower Weber Canyon section.

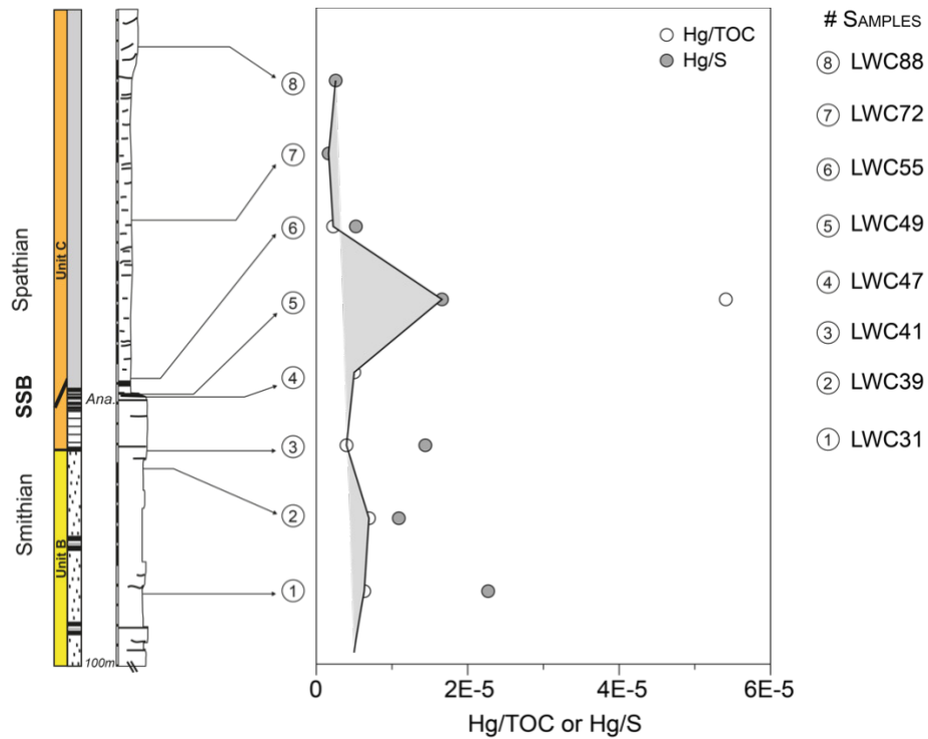


Figure S2: Hg concentrations normalized to TOC or S concentrations along the LWC stratigraphy.

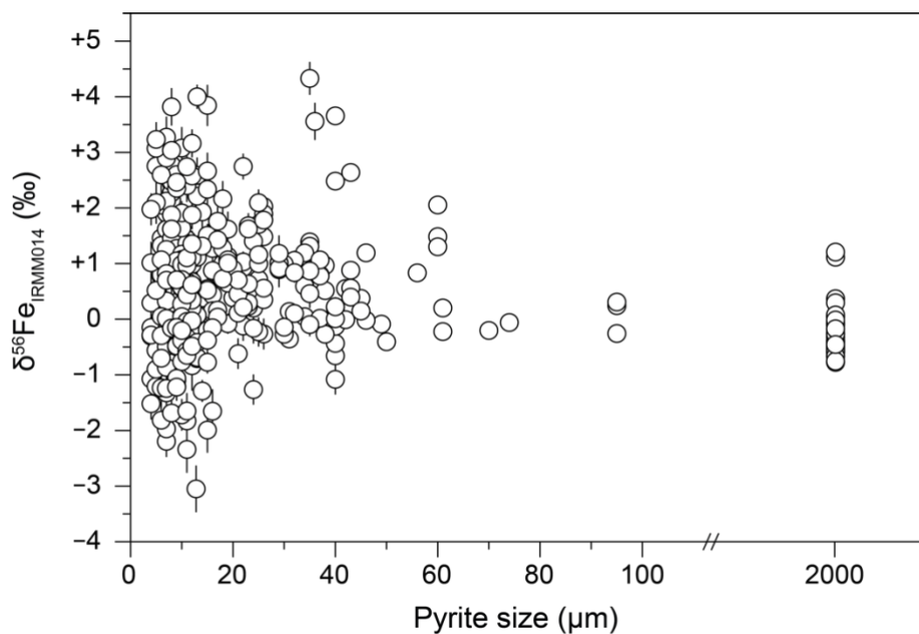


Figure S3: $\delta^{56}\text{Fe}_{\text{py}}$ values as a function of pyrite size.

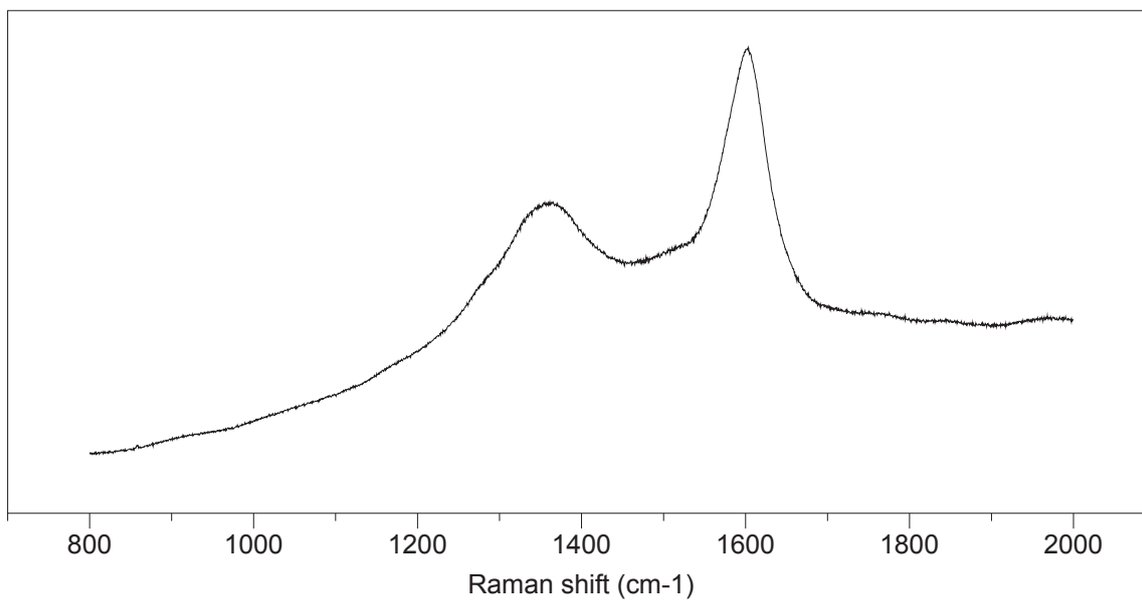


Figure S4: Raman spectra of organic matter in LWC54 (equivalent to LWC55). Graphite (G) and defect (D) bands provide information on the structural organization of the aromatic skeleton (Beysac et al., 2002), which underwent here temperature of $\sim 150^\circ\text{C}$.

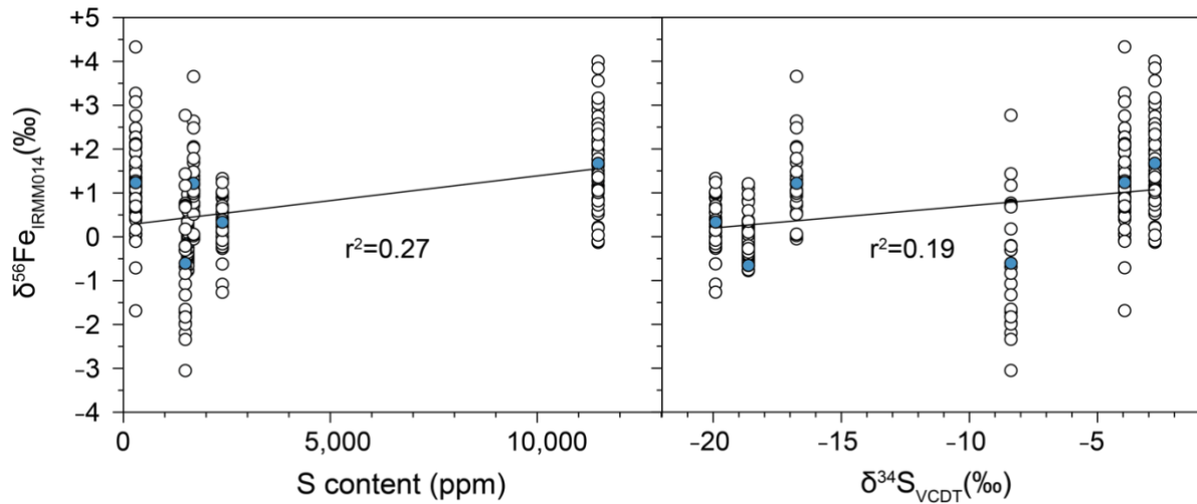


Figure S5: $\delta^{56}\text{Fe}_{\text{py}}$ measured by SIMS as a function of the S content or the bulk S isotope composition in LWC31 to LWC55. Blue dots correspond to the mean $\delta^{56}\text{Fe}$ values.

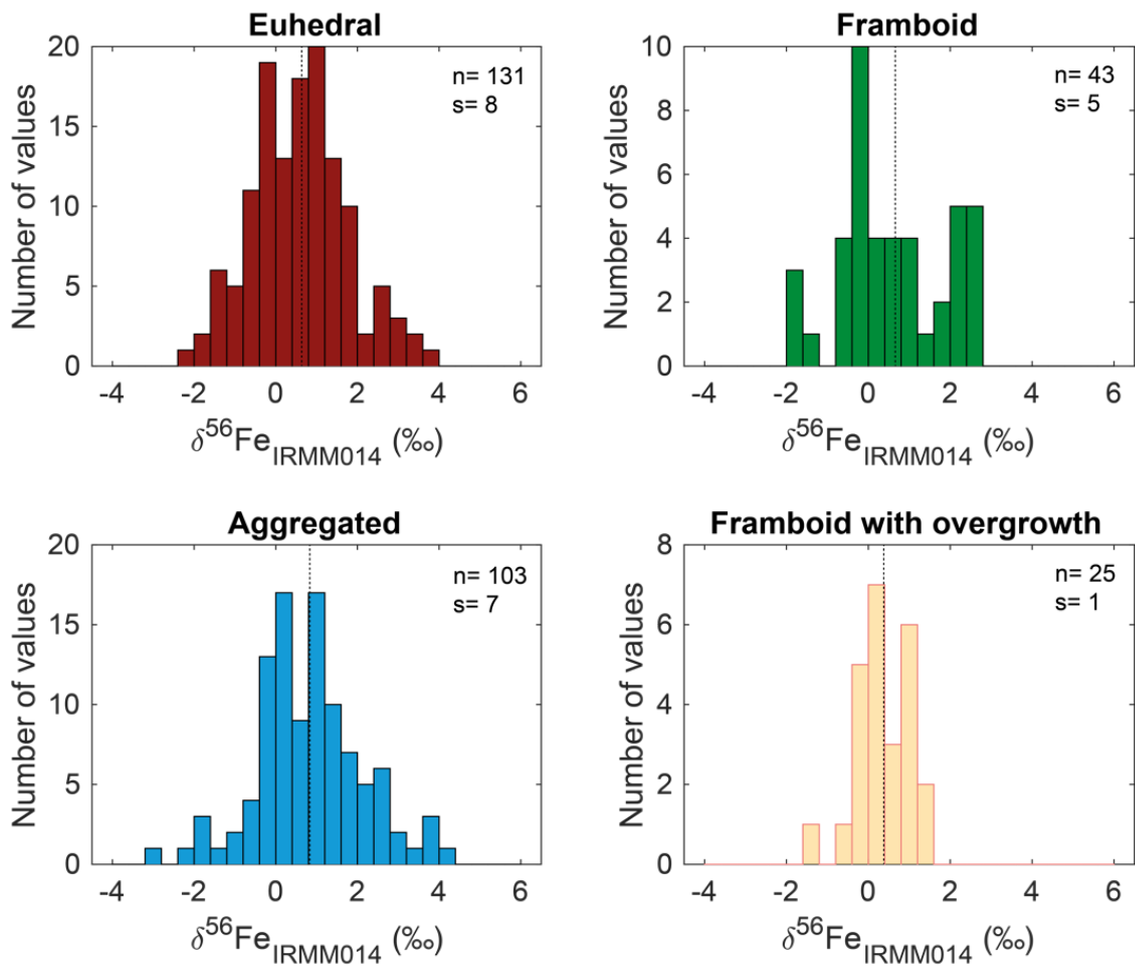


Figure S6: Pyrite $\delta^{56}\text{Fe}$ distributions as a function of pyrite morphologies. The dashed line in each histogram represents the mean value. The number of values (n), and the number of samples from which pyrite grains were measured (s) are indicated for each histogram.

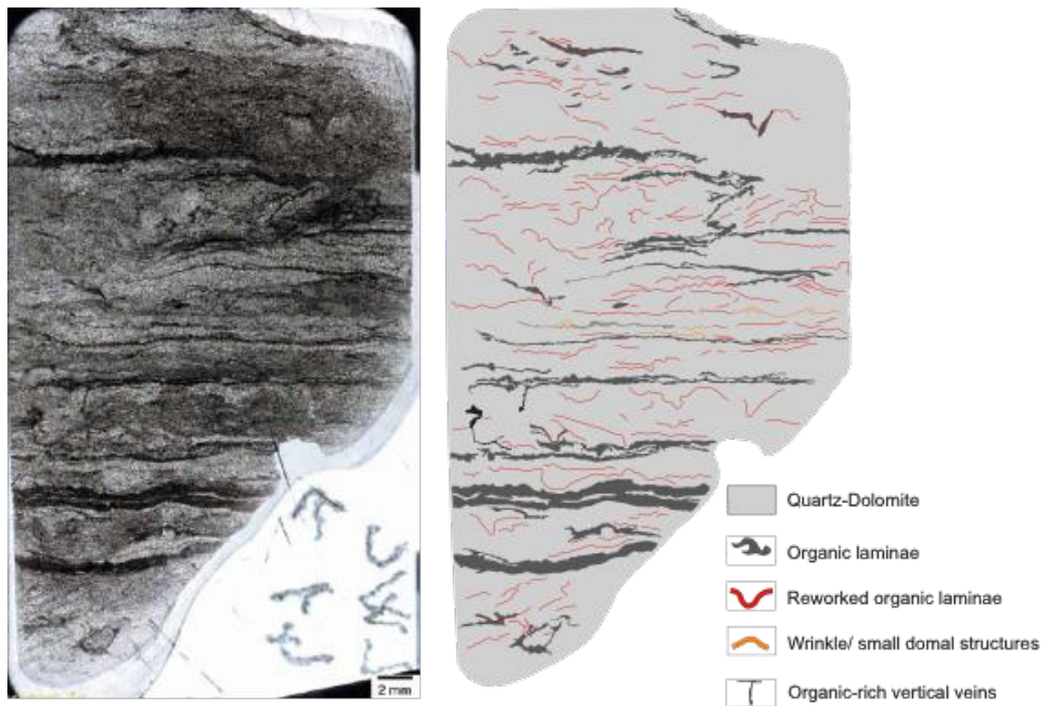


Figure S7: Optical microscope large view of sample LWC41 showing reworked organic laminae (bioturbation)

Table S1: Mn/Sr ratios, S content (ppm), $\delta^{34}\text{S}$ values for pyrite and CAS in LWC samples

Sample #	Mn/Sr	S content (ppm)	$\delta^{34}\text{S}$ S1 norm (‰)	SD	Setting	$\delta^{34}\text{S}_{\text{CAS}}$ NBS-127 norm (‰)
LWC70	17.4	<i>n.d.</i>	4.2	0.4	Outer ramp	
LWC69	26.8	126	1.4	0.0		
LWC68	<i>n.d.</i>	75	16.3	0.2		
LWC67	10.9	61	6.1	0.2	Average	5.6 ‰
LWC64	<i>n.d.</i>	661	13.1	0.1	Stdv	7.5 ‰
LWC63	21.3	71	14.1	0.1	Min	-7.3 ‰
LWC61	21.9	82	11.7	0.1	Max	16.3 ‰
LWC60	<i>n.d.</i>	781	-0.3	0.9		
LWC59	21.9	12358	5.2	0.0		33.5
LWC57	<i>n.d.</i>	9226	-7.3	0.9		
LWC55	21.5	11472	-2.8	0.9	Mid ramp	
LWC54	<i>n.d.</i>	11267	-7.5	0.9		
LWC53	40.0	1621	-3.5	0.0	Average	-4.2 ‰
LWC52	22.3	5298	4.5	0.3	Stdv	4.2 ‰
LWC50	29.1	1409	-2.2	0.4	Min	-8.4 ‰
LWC49	<i>n.d.</i>	296	-3.9	0.0	Max	4.5 ‰
LWC48	13.1	3811	-4.3	0.5		
LWC47	<i>n.d.</i>	1496	-8.4	0.2		30.7
LWC46	14.0	1422	-8.2	0.1		
LWC42	<i>n.d.</i>	59	-10.9	0.1	Inner ramp	
LWC41	31.6	2391	-19.9	0.0		
LWC40	36.6	1548	-14.8	0.1		
LWC39	<i>n.d.</i>	1695	-16.8	0.1	Average	-11.4 ‰
LWC38	<i>n.d.</i>	62	-2.1	0.0	Stdv	9.3 ‰
LWC37	<i>n.d.</i>	21	0.5	ND	Min	-20.5 ‰
LWC35	32.4	1318	-20.3	0.1	Max	5.8 ‰
LWC34	22.2	3780	-7.8	0.04		
LWC32	<i>n.d.</i>	68	5.8	0.0		29.2
LWC30	28.7	956	-20.5	0.1		
LWC31	25.2	1555	-18.6	0.3		

Table S2: $^{56}\text{Fe}^+$ intensities, $\delta^{56}\text{Fe}$ values and yields of pyrite standards (Balmat and Ruttan; all SIMS sessions)

Analysis #	$^{56}\text{Fe}^+$ intensity	$\delta^{56}\text{Fe}_{\text{corr}}$	2SE	Yield $^{56}\text{Fe}^+$
Balmat_LWC88@01	4.48E+07	-33.37	0.15	1.50E+07
Balmat_LWC88@2	4.51E+07	-33.16	0.15	1.51E+07
Balmat_LWC88@3	4.44E+07	-33.29	0.17	1.49E+07
Balmat_LWC88@4	4.51E+07	-33.22	0.16	1.50E+07
Balmat_LWC88@5	4.41E+07	-33.29	0.16	1.47E+07
Balmat_LWC88@6	4.48E+07	-33.33	0.19	1.50E+07
Balmat_LWC88@7	4.90E+07	-33.84	0.16	1.55E+07
Balmat_LWC88@8	4.94E+07	-33.78	0.17	1.56E+07
Balmat_LWC88@9	4.93E+07	-33.83	0.16	1.56E+07
Balmat_LWC88@10	5.01E+07	-33.42	0.14	1.59E+07
Balmat_LWC88@12	4.86E+07	-33.61	0.18	1.54E+07
Balmat_LWC88@11	4.98E+07	-33.51	0.15	1.58E+07
BalmatLWC72@12	4.49E+07	-33.58	0.19	1.63E+07
BalmatLWC72@13	4.49E+07	-33.54	0.21	1.63E+07
BalmatLWC72@14	4.54E+07	-33.45	0.20	1.64E+07
BalmatLWC72@15	4.49E+07	-33.49	0.19	1.62E+07
BalmatLWC72@16	4.49E+07	-33.38	0.20	1.63E+07
BalmatLWC72@17	4.57E+07	-33.44	0.23	1.64E+07
BalmatLWC72@18	4.53E+07	-33.46	0.24	1.63E+07
BalmatLWC72@19	4.51E+07	-33.62	0.23	1.62E+07
BalmatLWC72@20	4.65E+07	-33.32	0.22	1.67E+07
BalmatLWC72@21	4.59E+07	-33.78	0.24	1.65E+07
BalmatLWC72@22	4.60E+07	-33.56	0.23	1.65E+07
BalmatLWC72@23	4.68E+07	-33.41	0.25	1.67E+07
BalmatLWC72@24	4.60E+07	-33.55	0.26	1.64E+07
BalmatLWC72@25	4.59E+07	-33.61	0.26	1.65E+07
BalmatLWC72@26	4.62E+07	-33.36	0.26	1.66E+07
BalmatLWC72@27	4.77E+07	-33.16	0.15	1.70E+07
BalmatLWC72@28	4.77E+07	-33.14	0.15	1.70E+07
BalmatLWC72@29	4.78E+07	-33.07	0.14	1.70E+07
BalmatLWC72@30	4.76E+07	-33.74	0.15	1.70E+07
BalmatLWC72@31	4.74E+07	-33.76	0.15	1.68E+07
BalmatLWC72@32	4.71E+07	-33.93	0.16	1.68E+07
BalmatLWC47@1	4.69E+07	-33.88	0.16	1.68E+07
BalmatLWC47@2	4.62E+07	-34.16	0.16	1.66E+07
BalmatLWC47@3	4.69E+07	-33.81	0.16	1.69E+07
BalmatLWC47@8	4.53E+07	-33.98	0.15	1.61E+07
BalmatLWC47@9	4.55E+07	-33.95	0.10	1.62E+07
BalmatLWC47@10	4.74E+07	-33.96	0.14	1.69E+07
BalmatLWC47@11	4.72E+07	-34.04	0.16	1.69E+07

(Continued)

Analysis #	$^{56}\text{Fe}^+$ intensity	$\delta^{56}\text{Fe}_{\text{corr}}$	2SE	Yield $^{56}\text{Fe}^+$
BalmatLWC47@12	4.70E+07	-33.83	0.15	1.68E+07
BalmatLWC47@13	4.65E+07	-33.90	0.15	1.66E+07
BalmatLWC47@14	4.67E+07	-33.96	0.14	1.67E+07
BalmatLWC47@15	4.65E+07	-33.40	0.14	1.67E+07
BalmatLWC47@16	4.61E+07	-33.59	0.15	1.66E+07
BalmatLWC47@17	4.62E+07	-33.50	0.15	1.67E+07
BalmatLWC47@18	4.60E+07	-33.59	0.16	1.66E+07
BalmatLWC47@20	4.67E+07	-33.54	0.16	1.69E+07
BalmatLWC47@21	4.61E+07	-33.55	0.15	1.66E+07
BalmatLWC47@22	4.62E+07	-33.72	0.17	1.66E+07
BalmatLWC47@23	4.61E+07	-33.60	0.14	1.67E+07
BalmatLWC47@24	4.57E+07	-33.76	0.15	1.65E+07
BalmatLWC47@25	4.57E+07	-33.75	0.15	1.65E+07
BalmatLWC47@26	4.56E+07	-33.85	0.16	1.65E+07
BalmatLWC47@27	4.56E+07	-33.66	0.15	1.65E+07
BalmatLWC88@9	4.51E+07	-33.86	0.17	1.64E+07
BalmatLWC88@10	4.53E+07	-34.05	0.15	1.65E+07
BalmatLWC88@11	4.53E+07	-34.02	0.15	1.64E+07
BalmatLWC88@12	4.54E+07	-33.99	0.15	1.64E+07
BalmatLWC88@15	4.58E+07	-33.96	0.14	1.66E+07
BalmatLWC88@16	4.57E+07	-34.11	0.15	1.66E+07
BalmatLWC88@17	4.56E+07	-33.87	0.15	1.66E+07
BalmatLWC88@18	4.63E+07	-33.98	0.17	1.68E+07
BalmatLWC55@01	4.19E+07	-34.05	0.22	1.52E+07
BalmatLWC55@2	4.16E+07	-34.00	0.22	1.51E+07
BalmatLWC55@3	4.22E+07	-33.83	0.22	1.53E+07
BalmatLWC55@4	4.17E+07	-33.94	0.23	1.51E+07
BalmatLWC55@5	4.33E+07	-33.62	0.16	1.57E+07
BalmatLWC55@6	4.39E+07	-33.56	0.17	1.59E+07
BalmatLWC55@7	4.35E+07	-33.60	0.17	1.58E+07
BalmatLWC55@8	4.36E+07	-33.52	0.15	1.58E+07
BalmatLWC55@12	4.42E+07	-33.77	0.16	1.60E+07
BalmatLWC55@13	4.45E+07	-33.76	0.15	1.61E+07
BalmatLWC55@14	4.47E+07	-33.94	0.16	1.63E+07
BalmatLWC55@15	4.46E+07	-33.80	0.17	1.62E+07
BalmatLWC55@16	4.43E+07	-33.87	0.15	1.61E+07
BalmatLWC55@17	4.44E+07	-33.59	0.14	1.62E+07
BalmatLWC55@18	4.41E+07	-33.54	0.16	1.61E+07
BalmatLWC55@19	4.36E+07	-33.63	0.16	1.59E+07
BalmatLWC55@20	4.37E+07	-33.47	0.17	1.59E+07
BalmatLWC55@22	4.45E+07	-33.43	0.15	1.62E+07

(Continued)

Analysis #	$^{56}\text{Fe}^+$ intensity	$\delta^{56}\text{Fe}_{\text{corr}}$	2SE	Yield $^{56}\text{Fe}^+$
BalmatLWC55@23	4.46E+07	-33.44	0.15	1.63E+07
BalmatLWC55@24	4.50E+07	-33.54	0.16	1.64E+07
BalmatLWC55@25	4.44E+07	-33.37	0.17	1.62E+07
BalmatLWC49@1	4.61E+07	-33.73	0.31	1.66E+07
BalmatLWC49@2	4.54E+07	-34.04	0.32	1.64E+07
BalmatLWC49@3	4.55E+07	-33.95	0.32	1.65E+07
BalmatLWC49@4	4.56E+07	-34.04	0.32	1.65E+07
BalmatLWC49@5	4.55E+07	-34.01	0.31	1.65E+07
BalmatLWC49@6	4.60E+07	-33.51	0.32	1.66E+07
BalmatLWC49@7	4.63E+07	-33.73	0.31	1.67E+07
BalmatLWC49@8	4.63E+07	-33.50	0.31	1.68E+07
BalmatLWC49@9	4.76E+07	-33.56	0.31	1.72E+07
BalmatLWC49@10	4.62E+07	-33.87	0.31	1.66E+07
BalmatLWC49@11	4.69E+07	-33.51	0.32	1.68E+07
BalmatLWC49@12	4.70E+07	-33.62	0.32	1.69E+07
Balmat@01	4.19E+07	-32.84	0.12	1.40E+07
Balmat@2	4.35E+07	-32.97	0.12	1.44E+07
Balmat@3	4.29E+07	-32.74	0.13	1.40E+07
Balmat@4	4.35E+07	-32.51	0.12	1.42E+07
Balmat@5	4.26E+07	-32.53	0.13	1.38E+07
Balmat@6	4.30E+07	-32.81	0.11	1.40E+07
Ruttan@01	4.13E+07	-30.97	0.11	1.34E+07
Ruttan@2	4.22E+07	-31.03	0.13	1.38E+07
Ruttan@3	4.15E+07	-30.77	0.11	1.34E+07
Balmat@7	4.30E+07	-32.50	0.13	1.39E+07
Balmat@8	4.28E+07	-32.54	0.13	1.38E+07
Balmat@9	4.37E+07	-32.42	0.11	1.42E+07
Balmat@10	4.36E+07	-32.54	0.13	1.41E+07
Balmat@11	4.19E+07	-32.45	0.12	1.35E+07
Balmat@12	4.31E+07	-32.40	0.10	1.40E+07
Balmat@13	4.28E+07	-32.47	0.10	1.35E+07
Balmat@14	4.44E+07	-32.52	0.12	1.41E+07
Balmat@15	4.29E+07	-32.31	0.11	1.37E+07
Balmat@16	4.36E+07	-32.38	0.13	1.39E+07
Balmat@17	4.36E+07	-32.62	0.12	1.39E+07
Balmat@18	4.29E+07	-32.41	0.12	1.39E+07
Balmat@19	4.25E+07	-32.63	0.12	1.37E+07
Balmat@20	4.33E+07	-32.66	0.10	1.40E+07
Balmat@21	4.36E+07	-32.63	0.11	1.40E+07
Balmat@22	4.32E+07	-32.43	0.12	1.39E+07
Balmat@23	4.63E+07	-32.41	0.12	1.48E+07

(Continued)

Analysis #	$^{56}\text{Fe}^+$ intensity	$\delta^{56}\text{Fe}_{\text{corr}}$	2SE	Yield $^{56}\text{Fe}^+$
Balmat@24	4.58E+07	-32.45	0.10	1.47E+07
Balmat@25	4.44E+07	-32.43	0.13	1.43E+07
Balmat@26	4.40E+07	-32.46	0.11	1.42E+07
Balmat@27	4.45E+07	-32.48	0.10	1.43E+07
Balmat@28	4.52E+07	-32.36	0.12	1.50E+07
Balmat@29	4.51E+07	-32.02	0.10	1.50E+07
Balmat@30	4.51E+07	-32.25	0.11	1.51E+07
Balmat@31	4.55E+07	-32.17	0.09	1.52E+07
Balmat@32	4.50E+07	-32.16	0.10	1.50E+07
Balmat@33	4.38E+07	-32.41	0.11	1.42E+07
Balmat@34	4.32E+07	-32.31	0.11	1.42E+07
Balmat@35	4.37E+07	-32.31	0.12	1.43E+07
Balmat@36	4.32E+07	-32.42	0.11	1.41E+07
Balmat@37	4.32E+07	-32.35	0.11	1.42E+07
Balmat@39	4.47E+07	-32.44	0.10	1.45E+07
Balmat@40	4.46E+07	-32.38	0.10	1.44E+07
Balmat@41	4.42E+07	-32.35	0.12	1.43E+07
Balmat@42	4.51E+07	-32.22	0.13	1.45E+07
Balmat@43	4.42E+07	-32.27	0.11	1.42E+07
Balmat@44	4.42E+07	-32.43	0.11	1.43E+07
Balmat@45	4.32E+07	-32.15	0.10	1.41E+07
Balmat@46	4.47E+07	-32.20	0.09	1.47E+07
Balmat@48	4.41E+07	-32.12	0.12	1.45E+07
Balmat@49	4.48E+07	-32.13	0.10	1.47E+07
Balmat@50	4.36E+07	-32.51	0.12	1.43E+07
Balmat@52	4.40E+07	-32.35	0.12	1.43E+07
Balmat@53	4.32E+07	-32.50	0.13	1.41E+07
Balmat@54	4.32E+07	-32.38	0.11	1.41E+07
Balmat@55	4.47E+07	-32.14	0.10	1.47E+07
Balmat@56	4.47E+07	-32.14	0.11	1.47E+07
Balmat@57	4.51E+07	-31.98	0.09	1.48E+07
Balmat@58	4.50E+07	-32.07	0.11	1.49E+07
Balmat@59	4.46E+07	-31.95	0.11	1.47E+07
Balmat@60	4.45E+07	-32.24	0.11	1.46E+07

Table S3: Results of the Shapiro-Wilk test reported as p-values.

SHAPIRO-WILK TEST	SAMPLE #	P-VALUE	MAIN PYR. TYPE	P-VALUE	PYR. TYPE	P-VALUE	RAMP DOMAIN	P-VALUE	
	LWC88	0.197	Euhedral	0.591	FA1	Nodule	1.5E-6	Inner ramp	9.2E-7
	LWC72	0.038	Framboid	0.053		Framb.+ overg.	0.117	Mid ramp	0.023
	LWC55	0.633	Aggregated	0.180		Aggregated	0.941	Outer ramp	3.2E-4
	LWC49	0.408	Framboid + overgrowth	0.423		Euhedral	0.011		
	LWC47	0.545	Framb. core	0.484	FA2	Euhedral	0.229		
	LWC41-3a	0.151	Overgrowth	0.016		Framb.+ overg.	4.3E-3		
	LWC41-3b	0.453			Aggregated	0.779			
	LWC39	0.069			FA3	Euhedral	0.046		
	LWC31	5.7E-5				Aggregated	0.176		
				Framboid	0.179				
				FA4	Euhedral	0.964			
					Aggregated	0.891			
				Framboid	0.198				
				FA5	Euhedral	0.036			
					Aggregated	0.865			
				Framboid	2.2E-4				

Table S4: Results of the Mann-Whitney test reported as p-values for the comparison of $\delta^{56}\text{Fe}$ values the different domains of the ramp.

MANN-WHITNEY TEST		
P-VALUE		
Ramp domain	Inner ramp	Mid ramp
Mid ramp	1.7E-10	0.011
Outer ramp	0.235	2.8E-7

Table S5: Results of the Mann-Whitney test and Student test reported as p-values for the comparison of Fe isotope compositions of the different LWC samples. LWC41-3a corresponds to LWC41 Siltst. and LWC41-3b to LWC41 OM

STUDENT TEST OR MANN-WHITNEY TEST
P-VALUE

Sample #	LWC72	LWC55	LWC49	LWC47	LWC41-3a	LWC41-3b	LWC39	LWC31
LWC88	0.249	2.9E-13	4.9E-9	0.023	0.342	0.001	2.5E-8	0.413
LWC55	5.1E-5	-	0.036	3.6E-11	3.5E-8	2.1E-5	0.005	9.4E-18
LWC49	0.002	-	-	8.0E-9	5.5E-6	0.009	0.964	1.7E-15
LWC47	0.007	-	-	-	0.354	1.7E-4	1.8E-7	0.079
LWC41-3a	0.219	-	-	-	-	2.1E-6	5.0E-7	0.663
LWC41-3b	0.167	-	-	-	-	-	0.003	2.7E-9
LWC39	0.003	-	-	-	-	-	-	0.003
LWC31	0.049	-	-	-	-	-	-	-

Table S6: Results of the Student test reported as p-values for the comparison of Fe isotope compositions of the different pyrite types in all samples.

STUDENT TEST
P-VALUE

Pyr. type #	Euhedral	Framboid	Aggregated	Framb+overg.
Euhedral	-	0.929	0.175	0.287
Framboid	-	-	0.318	0.397
Aggregated	-	-	-	0.083

Table S7: Results of the Mann-Whitney test and Student test reported as p-values for the comparison of Fe isotope compositions of the different pyrite types per facies.

STUDENT TEST OR MANN-WHITNEY TEST
P-VALUE

	Pyr. type per facies	Aggregated	Framb+ overg.	Nodule
FA1	Euhedral	0.695	0.011	0.002
	Aggregated	-	0.041	1.6E-4
	Framb+ overg.	-	-	0.640

	Pyr. type per facies	Aggregated	Framb+ overg.
FA2	Euhedral	0.137	0.007
	Aggregated	-	0.011

	Pyr. type per facies	Aggregated	Framboid
FA3	Euhedral	0.168	0.828
	Aggregated	-	0.510

	Pyr. type per facies	Aggregated	Framboid
FA4	Euhedral	0.528	0.483
	Aggregated	-	0.633

	Pyr. type per facies	Aggregated	Framboid
FA5	Euhedral	0.269	0.523
	Aggregated	-	0.328

Table S8: Hg content (in ppb), TOC and S contents in LWC samples

# Sample	Hg content (ppb)	TOC (%)	TOC (ppb)	Hg/TOC	S content (%)	S content (ppb)	Hg/S
LWC31	9.85	0.04	4.34E+05	2.27E-05	0.16	1.55E+06	6.33E-06
LWC39	11.80	0.11	1.09E+06	1.09E-05	0.17	1.70E+06	6.96E-06
LWC41	9.55	0.07	6.65E+05	1.44E-05	0.24	2.39E+06	3.99E-06
LWC47	7.50	0.15	1.52E+06	4.93E-06	0.15	1.50E+06	5.01E-06
LWC49	16.00	0.10	9.64E+05	1.66E-05	0.03	2.96E+05	5.41E-05
LWC55	25.55	0.49	4.88E+06	5.23E-06	1.15	1.15E+07	2.23E-06
LWC72	2.70	0.17	1.65E+06	1.63E-06	<i>Not meas.</i>	-	-
LWC88	4.05	0.16	1.58E+06	2.56E-06	<i>Not meas.</i>	-	-

Table S9: $\delta^{56}\text{Fe}$ values measured in 8 samples from the LWC section

Analyses #	$\delta^{56}\text{Fe}$ (‰)	2σ	Analyses #	$\delta^{56}\text{Fe}$ (‰)	2σ
P1@01	-0.65	0.13	P2@10	-0.47	0.13
P1@02	-0.38	0.14	P2@11	-0.58	0.15
P1@03	-0.09	0.11	P2@12	-0.57	0.13
P1@04	-0.55	0.11	P2@13	-0.27	0.11
P1@05	-0.40	0.10	P2@14	-0.17	0.13
P1@06	-0.48	0.13	P2@15	-0.67	0.14
P1@07	-0.43	0.11	P2@16	1.21	0.13
P1@08	0.37	0.11	P2@17	-0.75	0.10
P1@09	-0.39	0.11	P2@18	-0.45	0.12
P1@10	-0.12	0.11	LWC31@01	-0.66	0.11
P1@11	-0.51	0.12	LWC31@02	0.37	0.11
P1@12	-0.39	0.13	LWC31@03	0.24	0.11
P1@13	-0.23	0.13	LWC31@04	0.31	0.12
P1@14	-0.77	0.13	LWC31@05	-0.26	0.13
P1@15	-0.05	0.12	LWC31@06	-0.06	0.13
P1@16	-0.50	0.11	LWC31@07	0.83	0.11
P1@17	-0.41	0.11	LWC31@08	0.08	0.15
P1@18	-0.46	0.11	LWC31@09	-0.22	0.13
P1@19	-0.27	0.20	LWC31@10	0.20	0.11
P1@20	0.04	0.12	LWC31@11	0.79	0.13
P1@21	0.04	0.12	LWC31@12	-0.43	0.11
P1@22	-0.61	0.13	LWC31@13	-0.13	0.11
P1@23	0.30	0.13	LWC31@14	-0.01	0.14
P1@24	-0.40	0.11	LWC31@15	0.02	0.14
P1@25	0.08	0.10	LWC31@16	-0.09	0.14
P1@26	-0.58	0.13	LWC31@17	0.21	0.10
P1@27	-0.43	0.14	LWC31@18	-0.40	0.13
P1@28	-0.52	0.11	LWC31@19	0.97	0.15
P1@29	-0.54	0.14	LWC31@20	0.60	0.14
P1@30	-0.08	0.13	LWC31@21	0.05	0.13
P2@01	-0.01	0.14	LWC31@22	-0.20	0.13
P2@02	-0.18	0.11	LWC31@23	0.20	0.16
P2@03	-0.59	0.10	LWC31@24	-0.35	0.14
P2@04	-0.26	0.14	LWC31@25	0.14	0.13
P2@05	-0.32	0.12	LWC31@26	0.10	0.13
P2@06	-0.32	0.11	LWC31@27	-0.27	0.12
P2@07	-0.50	0.12	LWC31@28	0.35	0.13
P2@08	1.11	0.12	LWC31@29	-0.27	0.12
P2@09	-0.78	0.12	LWC39@01	0.55	0.14

(Continued)

Analyses #	$\delta^{56}\text{Fe}$ (‰)	2σ	Analyses #	$\delta^{56}\text{Fe}$ (‰)	2σ
LWC39@3	0.72	0.14	LWC41@13	0.01	0.17
LWC39@7	1.22	0.14	LWC41@14	0.22	0.17
LWC39@8	1.70	0.14	LWC41@15	0.98	0.25
LWC39@9	1.39	0.17	LWC41@16	1.00	0.16
LWC39@10	1.31	0.17	LWC41@17	1.01	0.17
LWC39@12	1.19	0.15	LWC41@18	0.71	0.18
LWC39@13	-0.02	0.13	LWC41@19	1.33	0.18
LWC39@14	1.19	0.14	LWC41@21	1.02	0.20
LWC39@15	2.05	0.15	LWC41@22	0.65	0.20
LWC39@16	1.48	0.13	LWC41@23	-0.08	0.18
LWC39@17	1.30	0.14	LWC41@25	1.24	0.18
LWC39@18	0.51	0.15	LWC41@26	0.38	0.18
LWC39@19	0.78	0.14	LWC41@27	0.44	0.18
LWC39@20	1.07	0.15	LWC41@28	0.11	0.17
LWC39@21	2.64	0.14	LWC41@29	0.38	0.17
LWC39@23	1.48	0.16	LWC41@30	-1.08	0.27
LWC39@24	0.93	0.17	LWC41@36	-0.27	0.28
LWC39@25	1.03	0.19	LWC41@37	-0.14	0.27
LWC39@26	2.02	0.16	LWC41@38	-0.23	0.26
LWC39@27	1.06	0.17	LWC41@39	-0.12	0.27
LWC39@28	1.79	0.17	LWC41@41	-0.16	0.27
LWC39@29	2.48	0.17	LWC41@42	-1.27	0.27
LWC39@30	3.66	0.16	LWC41@43	0.11	0.27
LWC39@31	1.00	0.17	LWC41@44	-0.01	0.28
LWC39@35	0.99	0.18	LWC41@45	-0.62	0.28
LWC39@41	1.04	0.25	LWC41@46	0.15	0.27
LWC39@42	-0.05	0.25	LWC47@01	-2.20	0.28
LWC43@46	0.02	0.25	LWC47@2	-0.30	0.24
LWC39@49	0.06	0.24	LWC47@5	-1.07	0.26
LWC41@01	0.56	0.18	LWC47@22	-1.33	0.27
LWC41@2	0.40	0.17	LWC47@23	0.77	0.25
LWC41@3	0.56	0.16	LWC47@24	1.44	0.32
LWC41@4	0.88	0.17	LWC47@25	0.18	0.35
LWC41@5	0.29	0.18	LWC47@27	-0.20	0.21
LWC41@6	0.23	0.17	LWC47@28	2.77	0.36
LWC41@9	0.90	0.17	LWC47@29	-1.72	0.29
LWC41@10	0.90	0.16	LWC47@30	-0.30	0.25
LWC41@11	1.00	0.24	LWC47@31	-1.82	0.31
LWC41@12	0.55	0.17	LWC47@34	-1.97	0.31

(Continued)

Analyses #	$\delta^{56}\text{Fe}$ (‰)	2σ	Analyses #	$\delta^{56}\text{Fe}$ (‰)	2σ
LWC47@36	-0.21	0.28	LWC49@44	2.44	0.35
LWC47@01	-3.05	0.42	LWC49@45	1.93	0.35
LWC47@2	-2.34	0.42	LWC49@46	2.48	0.34
LWC47@4	-1.65	0.39	LWC49@47	2.14	0.35
LWC47@5	-0.68	0.41	LWC49@48	1.45	0.36
LWC47@7	0.72	0.42	LWC49@49	2.27	0.37
LWC47@8	0.71	0.44	LWC49@50	2.12	0.36
LWC47@9	-0.84	0.45	LWC49@51	2.10	0.38
LWC47@10	0.68	0.44	LWC49@53	2.75	0.39
LWC47@11	-1.99	0.41	LWC49@54	1.18	0.33
LWC47@12	-1.83	0.41	LWC49@55	-0.10	0.37
LWC47@13	1.17	0.48	LWC49@56	1.09	0.35
LWC49@01	0.89	0.32	LWC49@57	1.27	0.35
LWC49@2	0.91	0.32	LWC49@59	0.80	0.36
LWC49@3	1.18	0.32	LWC49@1	0.16	0.27
LWC49@4	1.13	0.34	LWC49@2	0.63	0.25
LWC49@5	1.62	0.32	LWC49@3	-1.69	0.27
LWC49@6	1.46	0.34	LWC49@4	0.46	0.26
LWC49@7	1.27	0.34	LWC49@5	4.33	0.30
LWC49@8	0.15	0.32	LWC49@6	0.87	0.30
LWC49@9	1.33	0.34	LWC49@7	0.68	0.27
LWC49@11	1.36	0.33	LWC49@8	1.70	0.27
LWC49@12	3.27	0.37	LWC49@9	0.70	0.24
LWC49@14	0.90	0.31	LWC49@11	3.08	0.38
LWC49@15	0.46	0.32	LWC49@12	-0.71	0.26
LWC49@16	-0.08	0.37	LWC55@2	2.16	0.32
LWC49@18	1.02	0.36	LWC55@3	1.17	0.29
LWC49@21	0.39	0.36	LWC55@4	2.58	0.32
LWC49@22	0.04	0.35	LWC55@5	1.98	0.29
LWC49@26	0.07	0.37	LWC55@7	1.76	0.26
LWC49@30	1.25	0.39	LWC55@8	0.22	0.24
LWC49@31	1.91	0.36	LWC55@10	2.60	0.27
LWC49@33	1.52	0.36	LWC55@11	2.90	0.28
LWC49@36	1.36	0.34	LWC55@12	2.21	0.32
LWC49@37	1.34	0.36	LWC55@13	2.75	0.23
LWC49@38	0.56	0.36	LWC55@16	3.07	0.37
LWC49@39	1.11	0.35	LWC55@17	2.40	0.29
LWC49@40	2.05	0.35	LWC55@18	0.52	0.25
LWC49@43	1.62	0.37	LWC55@19	1.68	0.23

(Continued)

Analyses #	$\delta^{56}\text{Fe}$ (‰)	2σ	Analyses #	$\delta^{56}\text{Fe}$ (‰)	2σ
LWC55@20	1.64	0.35	LWC72@8	0.29	0.27
LWC55@21	-0.13	0.26	LWC72@9	-0.56	0.25
LWC55@22	4.00	0.22	LWC72@10	-1.21	0.29
LWC55@23	2.58	0.26	LWC72@11	-0.56	0.27
LWC55@24	1.27	0.20	LWC72@12	-0.09	0.20
LWC55@25	1.40	0.21	LWC72@13	-0.49	0.23
LWC55@26	0.21	0.30	LWC72@14	-1.21	0.22
LWC55@27	3.56	0.33	LWC72@16	-0.20	0.20
LWC55@28	3.03	0.26	LWC72@17	0.33	0.23
LWC55@29	-0.10	0.21	LWC72@18	-1.29	0.20
LWC55@30	3.84	0.37	LWC72@22	1.88	0.26
LWC55@31	1.62	0.20	LWC72@23	-0.34	0.25
LWC55@32	1.16	0.21	LWC72@24	2.74	0.30
LWC55@33	1.91	0.21	LWC72@27	2.66	0.33
LWC55@34	1.51	0.20	LWC72@29	1.07	0.25
LWC55@35	0.87	0.19	LWC72@30	-0.90	0.29
LWC55@36	0.20	0.19	LWC72@31	-0.07	0.27
LWC55@38	0.59	0.18	LWC72@32	3.23	0.31
LWC55@39	1.89	0.26	LWC72@33	-0.15	0.32
LWC55@40	0.84	0.18	LWC72@36	0.52	0.28
LWC55@41	1.43	0.19	LWC72@41	-0.01	0.30
LWC55@42	0.73	0.18	LWC72@42	-1.23	0.30
LWC55@43	1.78	0.21	LWC72@44	0.58	0.31
LWC55@44	0.82	0.23	LWC72@45	1.05	0.34
LWC55@46	3.16	0.25	LWC72@47	0.98	0.39
LWC55@48	2.09	0.24	LWC72@48	-0.20	0.27
LWC55@50	2.47	0.26	LWC72@49	1.07	0.30
LWC55@51	0.04	0.18	LWC72@50	3.82	0.34
LWC55@52	1.11	0.26	LWC72@51	1.02	0.38
LWC55@53	1.36	0.23	LWC72@52	-0.87	0.30
LWC55@54	1.07	0.22	LWC88@02	-0.50	0.27
LWC55@55	1.01	0.18	LWC88@06	0.82	0.26
LWC55@56	1.06	0.28	LWC88@20	0.70	0.22
LWC55@58	2.34	0.29	LWC88@22	-0.10	0.18
LWC72@01	-0.20	0.22	LWC88@23	0.05	0.14
LWC72@2	1.87	0.32	LWC88@24	-0.04	0.19
LWC72@3	2.35	0.25	LWC88@25	-0.03	0.17
LWC72@5	2.59	0.30	LWC88@26	-0.77	0.20
LWC72@7	-1.52	0.28	LWC88@27	-0.65	0.17

(Continued)

Analyses #	$\delta^{56}\text{Fe}$ (‰)	2σ
LWC88@28	-0.15	0.19
LWC88@29	-0.20	0.18
LWC88@30	-0.49	0.21
LWC88@38	-0.47	0.15
LWC88@37	-0.48	0.19
LWC88@54	-0.15	0.15
LWC88@2	0.05	0.28
LWC88@3	0.96	0.35
LWC88@4	1.31	0.26
LWC88@5	-0.69	0.22
LWC88@7	-0.42	0.22
LWC88@8	1.62	0.27
LWC88@9	-0.28	0.24
LWC88@10	0.71	0.30
LWC88@11	1.10	0.22
LWC88@14	-0.29	0.29
LWC88@15	-0.38	0.21
LWC88@16	-0.76	0.23
LWC88@17	0.01	0.23
LWC88@18	-1.10	0.24
LWC88@20	-0.16	0.23
LWC88@22	-1.06	0.24
LWC88@23	0.34	0.22
LWC88@24	0.62	0.21
LWC88@25	0.27	0.21
LWC88@26	1.35	0.24
LWC88@27	2.46	0.23
LWC88@31	-0.04	0.21
LWC88@32	-0.38	0.24
LWC88@33	0.44	0.22
LWC88@34	-1.24	0.31
LWC88@35	0.71	0.26
LWC88@36	-1.22	0.25
LWC88@37	-1.65	0.32
LWC88@38	-0.70	0.29
LWC88@39	0.62	0.21
LWC88@40	-0.57	0.22
LWC88@41	1.62	0.22

