Environmental Insights From High-Resolution (SIMS) Sulfur Isotope Analyses of Sulfides in Proterozoic Microbialites With Diverse Mat Textures

The Harvard community has made this article openly available. Please share how this access benefits you. Your story matters

Citation

Citable link
http://nrs.harvard.edu/urn-3:HUL.InstRepos:41292726

Terms of Use
This article was downloaded from Harvard University’s DASH repository, WARNING: This file should NOT have been available for downloading from Harvard University’s DASH repository.
Environmental insights from high-resolution (SIMS) sulfur isotope analyses of sulfides in Proterozoic microbialites with diverse mat textures

M. L. Gomes¹,² | D. A. Fike¹ | K. D. Bergmann³ | C. Jones¹ | A. H. Knoll²

¹Washington University, Saint Louis, MO, USA
²Harvard University, Cambridge, MA, USA
³Massachusetts Institute of Technology, Cambridge, MA, USA

Correspondence
M. L. Gomes, Johns Hopkins University, Baltimore, MD, USA.
Email: mgomes@jhu.edu

Abstract
In modern microbial mats, hydrogen sulfide shows pronounced sulfur isotope ($\delta^{34}S$) variability over small spatial scales (~50‰ over <4 mm), providing information about microbial sulfur cycling within different ecological niches in the mat. In the geological record, the location of pyrite formation, overprinting from mat accretion, and post-depositional alteration also affect both fine-scale $\delta^{34}S$ patterns and bulk $\delta^{34}S_{pyrite}$ values. We report μm-scale $\delta^{34}S$ patterns in Proterozoic samples with well-preserved microbial mat textures. We show a well-defined relationship between $\delta^{34}S$ values and sulfide mineral grain size and type. Small pyrite grains (<25 μm) span a large range, tending toward high $\delta^{34}S$ values (~54.5‰ to 11.7‰, mean: −14.4‰). Larger pyrite grains (>25 μm) have low but equally variable $\delta^{34}S$ values (~61.0‰ to −10.5‰, mean: −44.4‰). In one sample, larger sphalerite grains (>35 μm) have intermediate and essentially invariant $\delta^{34}S$ values (~22.6‰ to −15.6‰, mean: −19.4‰). We suggest that different sulfide mineral populations reflect separate stages of formation. In the first stage, small pyrite grains form near the mat surface along a redox boundary where high rates of sulfate reduction, partial closed-system sulfate consumption in microenvironments, and/or sulfide oxidation lead to high $\delta^{34}S$ values. In another stage, large sphalerite grains with low $\delta^{34}S$ values grow along the edges of pore spaces formed from desiccation of the mat. Large pyrite grains form deeper in the mat at slower sulfate reduction rates, leading to low $\delta^{34}S_{sulfide}$ values. We do not see evidence for significant $^{34}S$-enrichment in bulk pore water sulfide at depth in the mat due to closed-system Rayleigh fractionation effects. On a local scale, Rayleigh fractionation influences the range of $\delta^{34}S$ values measured for individual pyrite grains. Fine-scale analyses of $\delta^{34}S_{pyrite}$ patterns can thus be used to extract environmental information from ancient microbial mats and aid in the interpretation of bulk $\delta^{34}S_{pyrite}$ records.

1 | INTRODUCTION

Microbial mats are multi-layered aggregations of phylogenetically and metabolically diverse microorganisms. Microbial mats can be preserved in the geological record as microbialites, which are ubiquitous through geological time (Grotzinger & Knoll, 1999) and thus have the potential to provide a rich suite of information about the evolution of microbial metabolisms and ancient environmental conditions. Precambrian microbialites have received special attention because they provide insight into our understanding of early life on Earth. Additionally, it is likely that metabolic activity of microbial mat communities played a dominant role in biogeochemical cycling on the early Earth and thus influenced both nutrient and oxygen availability (Hoehler, Bebout, & Des Marais, 2001; Anbar & Knoll, 2002; Hayes & Waldbauer, 2006).
Methodological and technological advances in geochemical imaging and fine-scale analyses have made it possible to evaluate geochemical heterogeneity in both modern microbial mats and microbialites over small spatial scales (i.e., µm-scale). By coupling these analyses with petrological information about mat textures, it is possible to evaluate potential signatures of biological process at play at the time of formation, ancient environmental conditions, evidence for post-depositional alteration, and the influence of fine-scale heterogeneity on traditional bulk-scale (i.e., >1 cm³) geochemistry. Here, we use this approach to evaluate sulfur isotope heterogeneity in sulfide minerals within microbialites from two time intervals in the Proterozoic Eon in order to explore patterns of sulfur cycling during this critical interval of Earth history.

Biogeochemical sulfur transformations are an important component of elemental cycling in microbial mats (Canfield & Des Marais, 1993; Habicht & Canfield, 1996; Fike, Gammon, Ziebis, & Orphan, 2008; Fike et al., 2009). Many modern microbial mats contain an upper pigmented layer with oxygenic phototrophs (e.g., Cyanobacteria) overlying and admixed with anoxygenic phototrophs (e.g., certain taxa of the Chloroflexi and Proteobacteria such as the Chloroflexales and Chromatiales, respectively). These phototrophic communities fix carbon to produce energy and biomass, fueling mat growth and thereby providing a substrate for heterotrophic communities. Sulfide is generated in the mats via microbial sulfate reduction (MSR), a process by which microorganisms (e.g., certain orders of the Proteobacteria such as the Desulfobacterales) use sulfate to respire organic carbon (CH₂O) or hydrogen (H₂, not shown in the reaction below), generating hydrogen sulfide (H₂S) and bicarbonate (HCO₃⁻):

\[
SO⁴^{2−} + 2CH₂O → H₂S + 2HCO₃⁻.
\]

The depth in the mat at which sulfide accumulates is called the sulfide chemocline. Diurnal cycles stimulate vertical migration of the sulfide chemocline (Canfield & Des Marais, 1993; Fike et al., 2009). During the day, photosynthetic oxygen production and sulfide oxidation drive the sulfide chemocline to deeper within the mat. At night, the chemocline moves upwards as the result of continued sulfide chemocline (Canfield & Des Marais, 1993; Fike et al., 2009). The magnitude of δ¹³S fractionation during MSR (εSR) depends on cell-specific sulfate reduction rates (csSRR; Habicht & Canfield, 1996; Sim, Bosak et al., 2011; Sim, Ono, Donovan, Templer, & Bosak, 2011; Leavitt, Haley, Bradley, & Johnston, 2013) and ambient sulfate concentrations (Habicht, Gade, Thamdrup, Berg, & Canfield, 2002; Wing & Haley, 2014; Bradley et al., 2015). First, there is a negative correlation between net sulfate reduction rate and εSR values that has been well documented in microbial cultures (Harrison & Thode, 1958; Kaplan & Rittenberg, 1964; Chambers, Trudinger, Smith, & Burns, 1975) and marine sediments (Goldhaber & Kaplan, 1975). Low csSRR, which contribute to a low net sulfate reduction rates in a system, can result from limitation by organic carbon availability, nutrient availability, low sulfate levels, high sulfide concentrations, or a combination of these factors (Wing & Haley, 2014; Bradley et al., 2015). Second, εSR values tend to scale with ambient sulfate concentrations, which can reflect strain-specific relationships between sulfate concentration and fractionation (Habicht et al., 2002; Wing & Haley, 2014; Bradley et al., 2015) or Rayleigh isotope fractionation as the sulfate reservoir is more readily depleted when sulfate levels are low (Gomes & Hurtgen, 2013). However, the relationship between sulfate concentration, csSRR, and εSR values differs among microbial species (Bradley et al., 2015) and may have varied over Earth history in association with secular changes in ocean chemistry and/or biological evolution.

Sulfide oxidation is also a globally significant process (Jorgensen, 1982; Canfield, 1991). However, the magnitude of δ³⁴S fractionations reported for various sulfide oxidation pathways is small (<10‰; Kaplan & Rafter, 1958; Kaplan & Rittenberg, 1964; Ivanov, Gogotova, Matrosov, & Zyakun, 1977; Fry, Gest, & Hayes, 1988; Zerkle, Farquhar, Johnston, Cox, & Canfield, 2009; Poser et al., 2014) relative to εSR values. Although sulfide oxidation can be important for driving overall δ³⁴S patterns, particularly in shallow marine sediments (Aller, Madrid, Chistoserdov, Aller, & Heilbrun, 2010; Fike, Bradley, & Rose, 2015), the consensus is that environmental controls on MSR play the dominant role in determining the evolution of δ³⁴Ssulfide values with depth in a sedimentary environment.

In the case of cyanobacterial mats from Guerrero Negro, Baja California Sur, Mexico, δ³⁴Ssulfide patterns with depth in the mat are interpreted in terms of changes in average csSRR because sulfide abundances were <1% of ambient sulfate concentrations, and therefore, neither sulfate limitation nor Rayleigh isotope effects influence δ³⁴Ssulfide trends (Fike et al., 2009). In the top 1–4 mm of the mat, δ³⁴Ssulfide values are ~15‰–40‰ higher than δ³⁴Ssulfide values deeper in the mat (Fike et al., 2008, 2009). Additional δ³⁴Ssulfide oscillations (~20%) occur well below the chemocline (~1.5 cm), and these are interpreted to reflect variable εSR at depth due to differential microbial metabolic activity that arises from different microbial populations, variations in availability of carbon substrates or other metabolites, or additional sulfur species (e.g., polysulfides) at depth. However, the consistent trend that emerges is that δ³⁴Ssulfide values are high near the surface redox boundary and decrease with distance from the redox boundary (Figure 1). Similarly, microbial consortia in marsh environments show a trend of high δ³⁴Ssulfide values near the redox boundary due to differential microbial metabolic activity rather than closed-system Rayleigh isotope effects (Wilbanks et al., 2014). Experimental evidence indicates that...
Sulfate concentration also influences $\delta^{34}$S$_{\text{sulfide}}$ patterns; mat incubations at low sulfate concentration (<1 mM) show smaller (~5‰) $\delta^{34}$S enrichments in sulfide toward the top of the mat and $\delta^{34}$S$_{\text{sulfide}}$ values tend to be closer to $\delta^{34}$S$_{\text{sulfate}}$ values (Fike et al., 2009). Sulfide yields were low in these incubations, so it is unlikely that the $\delta^{34}$S$_{\text{sulfide}}$ trends are due to closed-system Rayleigh isotope effects. Rather, the $\delta^{34}$S$_{\text{sulfide}}$ patterns can be attributed to low $\varepsilon_{\text{SR}}$ values associated with MSR at low sulfate concentrations, particularly when carried out by organisms that are otherwise adapted to higher sulfate concentrations (Bradley et al., 2015). Given how these trends vary with differential metabolic activity associated with microbial communities present at different depths in the mat and ambient sulfate concentration, $\delta^{34}$S$_{\text{sulfide}}$ Patterns preserved in ancient mats can provide additional paleoenvironmental context to complement bulk $\delta^{34}$S$_{\text{pyrite}}$ records.

Pyrite (FeS$_2$) is an important geological archive of $\delta^{34}$S$_{\text{sulfide}}$ trends in sedimentary systems where it is a ubiquitous component of marine sediments. However, there is still uncertainty about pathways of pyrite formation in sedimentary environments (see Rickard, 2012 for a review). Although studies of modern systems have shown that $\delta^{34}$S$_{\text{pyrite}}$ values can be offset from pore water $\delta^{34}$S$_{\text{sulfide}}$ values by up to 40‰, the working hypothesis is that $\delta^{34}$S$_{\text{pyrite}}$ values reflect $\delta^{34}$S$_{\text{sulfide}}$ values at the location of pyrite formation (Lyons, 1997) and the offset is due to transport of pyrite grains, unsteady diagenetic regimes, or pyrite formation in microenvironments (Bruchert & Pratt, 1996; Canfield, Raiswell, & Bottrell, 1992; Chanton, Martens, & Goldhaber, 1987; Raven, Sessions, Fischer, & Adkins, 2016). Petrographic information can be used to assess these effects.

It should be noted that above $\delta^{34}$S$_{\text{sulfide}}$ trends with depth in microbial mats differ from pore water $\delta^{34}$S$_{\text{sulfide}}$ trends in some modern marine sediments (e.g., Hartmann & Nielsen, 1968: Figure 1). Specifically, in some organic-rich modern marine sediments, pore water $\delta^{34}$S$_{\text{sulfide}}$ values are low near the top of the zone of sulfide accumulation and increase with depth. This pattern is a result of closed-system Rayleigh isotope fractionation effects where $\delta^{34}$S$_{\text{sulfide}}$ values evolve toward $\delta^{34}$S values of the overlying sulfate reservoir as the sulfate reservoir is progressively consumed. Such isotopic profiles are common in systems with high organic carbon availability, low rates of diffusion, and/or low sulfate concentrations (Gomes & Hurtgen, 2015). It is possible that the trend of $\delta^{34}$S-enrichment due to high rates of sulfate reduction near the top of the zone of sulfate reduction could be superimposed upon the closed-system trend (e.g., Raven et al., 2016; Figure 1). The closed-system trend has not been documented in modern microbial mats because the fraction of sulfate consumed in the studied systems is too low (on the order of 1%) to produce measurable Rayleigh isotope fractionation effects (Fike et al., 2009). However, taking a holistic view that considers both modern microbial mat and marine environments, high $\delta^{34}$S$_{\text{pyrite}}$ values could reflect the capture of the $S$-isotope signature of sulfide near the top of the zone of sulfate reduction (associated with high $c$SRR, low $\varepsilon_{\text{SR}}$ values, and oxidative sulfur cycling; Lyons, 1997; Habicht & Canfield, 2001; Dale, Bruchert, Alperin, & Regnier, 2009) or at greater depths where a large portion of the sulfate reservoir has been consumed. Another potential mechanism to generate high $\delta^{34}$S$_{\text{sulfide}}$ values, which could be captured in pyrite, is via Rayleigh isotope fractionation associated with MSR in microniches that are not well connected with the surrounding pore water. Intragrain $\delta^{34}$S variability could add insight into the influence of this process on $\delta^{34}$S$_{\text{pyrite}}$ values. Petrography is a key tool that can be used to assess the environmental context of pyrite formation and to distinguish between these possibilities. Thus, by coupling petrographic observations with fine-scale geochemical analyses, it is possible to make inferences about the location of pyrite formation and extract additional paleoenvironmental information from fine-scale $\delta^{34}$S$_{\text{pyrite}}$ patterns.

To evaluate how fine-scale $\delta^{34}$S$_{\text{sulfide}}$ patterns in modern microbial mats are preserved in sulfides in microbialites, we present fine-scale $\delta^{34}$S values of sulfide minerals from two microbialites from the Sukhaya Tunguska Formation, Siberia (Mesoproterozoic–Neoproterozoic transition; Sergeev, Knoll, & Petrov, 1997) and one microbialite from the Draken Formation, Spitsbergen (Neoproterozoic; Knoll, Swett, & Mark, 1991). We evaluate fine-scale $\delta^{34}$S$_{\text{pyrite}}$ patterns in the context of petrographic information and elemental mapping and compare the results to $\delta^{34}$S$_{\text{pyrite}}$ patterns from modern microbial mats (Fike et al., 2008, 2009) and microbial consortia (Wilbanks et al., 2014). This approach enables us to propose a framework for understanding the pyrite formation dynamics that influence bulk $\delta^{34}$S$_{\text{pyrite}}$ values in microbialites, an approach that can be extended to other types of sedimentary materials in order to gain additional insights into the spectrum of biogeochemical sulfur cycling associated with major Earth-life transitions in the past.
2 | MATERIALS AND METHODS

2.1 | Geological setting

Our samples come from silicified microbialites preserved in two Proterozoic carbonate platforms. The Sukhaya Tunguska Formation is exposed in the Turukhansk Uplift, along the northwestern margin of the Siberian Platform. This 530- to 670-m-thick succession records a shallowing-upward sequence from subtidal, basinal limestones to peritidal dolomites (Petrov, 1993; Petrov, Semikhatov, & Sergeev, 1995). Samples K95-101 and K95-103 come from a tidal flat/lagoonal unit midway through the formation (Petrov et al., 1995), which consists of intercalated dolarenite with low angle cross-stratification, dolomite, intraclastic conglomerate, and microbial laminites. Early diagenetic chert is common, particularly in the microbial mats, with excellent preservation of microfossils (Mendelson & Schopf, 1982; Sergeev et al., 1997). Direct age constraints are limited, but on the basis of microfossils and stromatolites, Russian stratigraphers have generally recognized the boundary between the Sukhaya Tunguska and overlying Derevnya formations as the Middle-Upper Riphean boundary, implying an age slightly >1,000 Ma (Petrov et al., 1995; Semikhatov et al., 2000). Carbonate carbon isotope stratigraphy is consistent with this estimate (Bartley et al., 2001), as is a Pb-Pb carbonate date of 1.035 ± 65 Ma (Ovchinnikova et al., 1995).

The Draken Formation is exposed in the glaciated highlands of Spitsbergen. Part of the nearly two thousand meter thick carbonate Akademikerbreen succession (Knoll & Swett, 1990), the 150- to 250-m-thick Draken Formation consists of tidal flat and lagoonal dolomites, including microbial laminites and low domal stromatolites, oolites, oncolites, and intraformational conglomerates, along with arenitic and micritic beds (Knoll et al., 1991; Fairchild, Knoll, & Swett, 1991). Although Draken carbonates are almost entirely dolomitic, fabric retention is excellent, suggesting penecontemporaneous dolomitization (Fairchild et al., 1991). Desiccation cracks occur sporadically throughout the succession, and both tepees and fenestrae are common, underscoring the peritidal setting of most Draken carbonates. Early diagenetic silicification is also common, with excellent preservation of microfossils (Knoll et al., 1991). Sample P4705-A comes from the uppermost part of the formation, directly beneath the lowermost oolitic beds of the Backlundtoppen Formation; it consists of interlaminated grainstones and intraclasts of microbial laminites, with fabric specific silicification of the mat fragments. The age of the sample is constrained by the ca. 800 Ma Bitter Springs carbon isotope anomaly in the underlying Svanbergfjellet Formation (Halverson, Hoffman, Schrag, Maloof, & Rice, 2005) and by the overlying 717–660 Ma Sturtian tillites (Rooney, Strauss, Brandon, & Macdonald, 2015). Consistent with this age, Draken samples contain vaso-shaped protistan microfossils comparable to those in successions from the Grand Canyon (Porter, Meisterfeld, & Knoll, 2003) and northwestern Canada (Strauss, Rooney, Macdonald, Brandon, & Knoll, 2014). The assemblage within the Draken is likely correlative with the older vaso-shaped microfossils from both localities. If correct, the Draken Formation is older than 740 Ma and well below the Islay Anomaly, which is preserved in the overlying Russøya Member in Svalbard (Karstrom et al., 2000; Hoffman et al., 2012; Strauss et al., 2014).

One reason for selecting samples from these two formations is that both sites have avoided significant post-depositional alteration. While both formations have been buried, involved in collisional tectonic events, and exhumed, the maximal temperatures experienced were similar, with no evidence for significant metasomatic fluid flow events. The Sukhaya Tunguska Formation of the Turukhansk Uplift experienced a collisional event following Mesoproterozoic deposition but prior to the Ediacaran (Nikishina, Sobornov, Prokopiev, & Frolov, 2010). This faulting and uplift resulted in at least 3–4 km of erosion and maximum burial temperatures of 150–200°C (Frolov et al., 2015). The source rocks that have been studied from the Mesoproterozoic succession are post-mature but TOC-rich (Frolov et al., 2015). This is in contrast to the overlying Ediacaran stratigraphy whose source rocks are immature to early mature indicating the early burial and uplift of the Sukhaya Tunguska Formation was the only significant deformational event in its history (Frolov et al., 2015). The geochemical character of Sukhaya Tunguska carbonates (both dolomites and limestones) has low Mn/Sr and Fe/Sr ratios and display 87Sr/86Sr ratios, δ13C values, and δ18O values similar to other rocks of their age (Gorokhov et al., 1995; Pope, Bartley, Knoll, & Petrov, 2003), which is consistent with minimal post-depositional alteration of the carbonates.

The Draken Formation is overlain by a significant overburden of Neoproterozoic to Ordovician rocks (~3 km; Knoll et al., 1991; Fairchild et al., 1991; Halverson, 2011). Additionally, deformation from the Caledonian Orogeny resulted in shortening and shear zones across Svalbard. However, this and more recent tectonic events appear to have had minimal thermal overprinting effects on the northern Ny Friesland succession (Fairchild et al., 2016). Thermal estimates of the Ordovician-aged strata indicate immature to early mature organic material from conodonts with CAI 1 (Lehnert, Stouge, & Brandl, 2013). Fine-scale observations suggest early alteration of the samples during shallow burial (Knoll et al., 1991; Knoll, Fairchild, & Swett, 1993; Fairchild et al., 1991). Geochemical analyses of carbonates from the Akademikerbreen Group are also similar to other carbonates of their age (Halverson, Wade, Hurtgen, & Barovich, 2010; Fairchild et al., 2016). As both sites have avoided significant burial and tectonic overprinting, they allow us to explore trends in sulfur isotopes preservation from samples of different ages with minimal bias imposed by differences in the metamorphic and metasomatic history.

An additional motivation for selecting samples from these two sites is their similar peritidal depositional setting and early diagenetic history including fabric-retentive dolomitization and selective early silicification in both cases. The mat fabrics from each unit are almost always preferentially silicified, and both sites contain exceptional microfossil preservation of both filamentous forms and other types of mat dwellers (Mendelson & Schopf, 1982; Sergeev et al., 1997; Knoll et al., 1991, 1993; Fairchild et al., 1991). Silicification preceded substantial decay of mat building and dwelling microorganisms. Insofar as pyrite formation preceded silica emplacement, this sharply constrains the timing of pyrite deposition.
2.2 | Bulk geochemical analyses

Pyrite was extracted from homogenized samples using the chromium reduction technique (Canfield, Raiswell, Westrich, Reaves, & Berner, 1986) in the Isotope Geobiology Laboratory at Harvard University. Briefly, powdered samples (~1 g) were loaded into vessels that were subsequently purged of O₂ by N₂ gas. Pyrite-S was extracted as H₂S by reaction with in boiling 1 M chromium chloride and 6N hydrochloric acid for 2 hr and driven by an N₂ carrier gas to a trap vessel containing 0.3 M zinc acetate and 0.6% acetic acid and converted to zinc sulfide. An aliquot of the trap vessel was removed for concentration determination by the Cline method using a Thermo Scientific GENESYS 10S UV/Vis Spectrophotometer (Cline, 1969). The remaining ZnS was converted to Ag₂S and purified by rinsing with 1M ammonium hydroxide, which removes all silver salts besides Ag₂S (Firsching, 1961), and deionized water. For δ³⁴S measurements, 0.35–0.45 mg of Ag₂S was weighed in tin boats along with 2–4 mg of vanadium pentoxide. Measurements of δ³⁴S values were made on a Thermo Scientific Delta V Plus continuous-flow isotope ratio mass spectrometer at Harvard University by combustion of Ag₂S to SO₂. S-isotope measurements were reproducible within 0.2% based on repeat analysis of international standards (IAEA S-1, IAEA S-2, and IAEA S-3).

Organic carbon concentration determinations were made on homogenized and acidified samples. To remove inorganic carbon, ~1 g of powdered samples was treated with 3N hydrochloric acid and allowed to react for 24 hr. The hydrochloric acid was removed by repeated rinses with deionized water. Calcium carbonate concentrations were determined gravimetrically based on loss during acidification. Organic carbon concentrations were determined on a Costech ECS 401 Elemental Analyzer coupled to a Delta V Plus mass spectrometer in the Stable Isotope Biogeochemistry Lab at Washington University. Analytical uncertainty is ±0.1‰ based on replicate analyses of laboratory standards.

2.3 | Petrography

Petrographic observations were made using a petrographic microscope. Images were taken using Leica EC4 Microscope camera. Scanning electron microscope images were also used for petrography (see Section 2.5).

2.4 | Grain size determinations

Grain diameters of sulfide minerals in thin sections were quantified from petrographic microscope images. These grain diameter values should be considered “apparent” grain size diameters because they were determined from the grain cross sections observed in thin section. If the thin section cut through the center of a spherical grain, then the measured diameter would represent the maximum diameter of the grain. In reality, the thin section exposes grain cross sections at random distances from the center of the grain. Therefore, the grain sizes reported here, and from similar studies of thin sections, are lower than the maximum diameter of the grain (assuming perfectly spherical grains). It is not standard procedure in many petrographic studies to perform a correction to account for this effect. However, we note that this effect should be considered when combining petrological observations with fine-scale geochemical analyses, particularly when trying to model environmental parameters giving rise to observed geochemical trends.

2.5 | Elemental analysis

Elemental mapping data and scanning electron microscope images were acquired using the JEOL-JXA-8200 Superprobe at the MIT Electron Microprobe Facility. Quantitative elemental spot analyses were carried out with an accelerating voltage of 15 kV, a beam current of 10 nA, and a beam size of 1 μm. Standards used for quantitative elemental analyses were chalcocite (S and Fe), hematite (O), and sphalerite (Zn). Detection limits were as follows: 60 ppm for S, 130 ppm for Fe, 200 ppm for O, and 500 ppm for Zn. Error estimates based on measurements of the standards (expressed as standard deviation values) were as follows: 0.23 wt% for S, 0.31 wt% for Fe, 0.47 wt% for O, and 0.70 wt% for Zn.

2.6 | Secondary ion mass spectrometry

Secondary ion mass spectrometry (SIMS) measurements were made on the CAMECA IMS 7f-GEO at the Department of Earth and Planetary Sciences at Washington University on one-inch diameter round thin sections. Prior to mounting, a small (~2 mm) hole was made with a diamond dental drill in each sample and a piece of the Washington University in-house pyrite standard (δ³⁴S = 0.13 ± 0.30‰; Meyer, Zerkle, & Fike, 2017) was inserted with epoxy. This internal standard allowed us to correct for isotopic variability between sample holders and between different analytical sessions. Thin sections were mounted and prepared with double polishing by High-Mesa Petrographics. Samples were gold coated (~35 nm) prior to analysis, in order to provide surface conductivity sufficient to sustain the stable secondary ion extraction field requisite for precise isotope ratio analysis. To further aid precision Cameca “Z-1” sample holders were used, these are specifically engineered to enhance the sample area available for precise isotope ratio analysis. Analyses were made in “spot mode.” This term indicates data were acquired via bombardment with a Cs⁺ ion beam rastered over a few micrometers, but without any specific x, y registration of data origin other than the central coordinates of the beam in relation to the sample. (In contrast, scanning ion image mode would require a larger raster and data collection would be co-registered with the temporal beam coordinates.) There were 3-week-long sessions (May 2015, November 2015, and March 2016). The primary beam was set to a current of ~1 nA and a diameter of ~2 μm and was further rastered in a 2 × 2 μm grid to produce a δ³⁴S measurement for a 4 × 4 μm area. Note that an unrastered larger spot could have been employed, however, the smaller spot plus raster mode was chosen as it allowed more flexibility to engage in close-proximity transect and/or micro-grain analyses. Experiments comparing small
spot plus raster mode with larger spot mode showed no difference in analytical precision.

Faraday cups were used to register relative ion currents for $^{32}$S (FC1: $>10^8$ c/s for 10 cycles) and $^{34}$S (FC2: $>4 \times 10^6$ c/s for 10 cycles) ions. Background levels on Faraday cups were calibrated at the beginning of each session and were $<2 \times 10^6$ c/s for $^{32}$S and $<4 \times 10^7$ c/s for $^{34}$S. Automatic peak centering, mass calibration, and pre-sputtering (10 s over a 4 μm × 4 μm area) were performed prior to each measurement. The small beam size (~2 μm) allowed for $^{34}$S measurements on small sulfide grains (i.e., ~6 μm), with an average internal relative standard deviation (RSD) of ~2.7‰ (calculated for n = 10 cycles for each sulfide analysis).

We performed a correction on sample and standard data to account for differences in the depth of $^{32}$S and $^{34}$S measurements. On the 7f/geo $^{32}$S and $^{34}$S, data are acquired sequentially, repeated over a number of cycles. Counts of $^{32}$S and $^{34}$S are acquired for 0.96 s and 4 s, respectively, with a 1.2-s wait time between analyses. This means that the time of acquisition for the $^{32}$S and $^{34}$S ions is slightly offset. This will not have an impact on measured $^{34}$S values if samples (and resulting ion counts) are homogenous; in our analyses, $^{32}$S and $^{34}$S counts each vary by <10% over the course of 10 cycles in homogeneous standards. However, $^{32}$S and $^{34}$S counts in heterogeneous pyrite grains can vary by up to 45% over the course of 10 cycles of data collection. In these cases, the timing of collection for $^{32}$S versus $^{34}$S can bias $^{34}$S results. To correct for this bias, we perform a linear interpolation on $^{32}$S count data to calculate a $^{32}$S count value that corresponds to the same time as the $^{34}$S measurement. This correction was performed on both standard and sample data.

The embedded standard was typically measured 2–4 times between every 9 and 15 sample measurements. We used the bracketing standard data to correct our sample data to minimize the effect of instrumental bias imposed by drift over the course of the analytical session (Valley & Kita, 2009). Replicate analyses of the embedded Washington University in-house standard ($\delta^{34}$S = 0.13 ± 0.30‰; Meyer et al., 2017) and a separately mounted Balmat standard ($\delta^{34}$S = 15.1‰) indicated that the average standard deviation (SD) of $\delta^{34}$S values measured on isotopically homogenous samples is 0.3‰. However, due to the greater isotopic heterogeneity in our samples compared to the standards, we report $\delta^{34}$S uncertainty for the unknowns as the SD of individual sulfide grain data (n = 10 cycles). This SD value is used to express the spread in the data over 10 cycles, but does not necessarily imply that the data were normally distributed because over 10 cycles the data were either randomly distributed around the mean or trended in a negative or positive direction. Data are provided in order of analysis in Table S1.

Sphalerite (ZnS) grains were also analyzed by SIMS. Isotope ratios for sphalerite grains were calculated using a correction factor that was determined from the analysis of a Washington University in-house sphalerite standard with a known $\delta^{34}$S value ($\delta^{34}$S = 3.5 ± 0.3‰ [2SD]; determined by gas-source isotope ratio mass spectrometry). The count rate for sphalerite grains was on average 1.5× the count rate of pyrite grains. Differences in the crystal orientation of sphalerite grains with respect to the incident primary Cs+ beam have been shown to introduce an instrumental bias that can result in an inaccuracy of up to 3% if the standard grain has a different orientation from the sample (Kozdon et al., 2010); in contrast, pyrites have a maximum known orientation effect of 0.2‰ (Kozdon et al., 2010). We did not determine the extent of inaccuracy imposed by the crystal orientation effect in our analyses and therefore use the maximum value reported by Kozdon et al. (2010) as our uncertainty in Table S2.

To account for potential spatial $\delta^{34}$S variability in larger pyrite grains (i.e., >14 μm), we performed $\delta^{34}$S measurements in grain transects along the central vertical and horizontal plane, as positioned in the SIMS and pictured in petrographic images. To evaluate biases imposed by closely spaced (~5 μm) measurements, we used the same method to measure vertical and horizontal transects on the Balmat standard. The close spacing (i.e., 5 μm) of measurements along the horizontal plane resulted in $\delta^{34}$S variability of up to 5.4‰ on standards. This increased isotopic variability is attributed to differential ionization effects introduced by interactions between the incident primary Cs+ beam and adjacent pits made by previous measurements; this effect occurs because the incident primary Cs+ beam approaches the surface of the sample at angle along the horizontal plane. Additional experiments determined that, for these analytical parameters, data were no longer compromised when a horizontal spacing >8 μm was used. This effect could also be avoided using diagonal steps for closely spaced measurements in orthogonal transects. The maximum $\delta^{34}$S variability was much lower (i.e., 1.7‰) on transects along the vertical plane (i.e., orthogonal to the angled direction of incident Cs+ beam), where the impact of adjacent spots was minimal. Thus, here we only report $\delta^{34}$S data for transects along the vertical plane. The $\delta^{34}$S measurements from each grain transect were averaged to calculate $\delta^{34}$S values for each pyrite grain, and we compare these values directly to the single $\delta^{34}$S measurements made on individual grains.

## RESULTS

The studied thin sections include centimeter-scale microbial mats, mat clasts, and clastic layers, which contain smaller mat fragments (Figure 2). Petrographic observations using both transmitted and reflected light, supplemented with elemental analysis data, indicate that portions of the samples that appear dark in Figure 2 contain finely disseminated, small (<25 μm) pyrite grains. The small pyrite grains represent ~1%–2% of these dark mat textures. The mat textures appear to be dark in both cross-polarized and plane-polarized light and are interpreted to be organic-rich (Figures 2, 4, and 5). In the lighter portion of the samples (Figure 2), localized populations of larger (>25 μm) sulfide grains were found on the edges of silicified void spaces. Thus, the general pattern is that small pyrite grains (<25 μm diameter) tend to occur in dark, organic-rich portions of the samples, while larger sulfide grains are found in organic-poor, silicified regions that appear transparent in plane-polarized light (Figures 2, 4, and 5). In what follows, we will describe each sample individually.
Sulfide grains in this sample are pyrite (Figures 3, 4a–c and 5a,b; Table S2). Most pyrite grains in this sample are small (<25 μm; only one of the 18 samples analyzed by SIMS was >25 μm), spherical, range from subeuhedral to euhedral grain textures, and cover <1% of the area of organic-rich mat textures (Figures 4a–c and 5a,b). Both the light and dark portions of the sample are silicified, with dolomite grains (<50 μm) comprising at most ~2% of the area in organic-rich mat textures of K95-103 (Figure 4f). Small pyrite (<25 μm) comprise ~1% of the area of organic-rich mat textures and have subeuhedral to euhedral spherical shapes (Figures 4d–f and 5c,d,g). Large sulfide grains in K95-103 occur on the edges of silicified fenestrae and are predominantly sphalerite (Figure 5e–h; Table S2). The shapes of the sphalerite grains are highly irregular and may result from the aggregation of smaller particles (Figures 5e–h).

In the Draken Formation sample P4705-A, the concentrations of S and Fe in sulfide grains range from the weight percent predicted from the stoichiometry of pyrite (FeS₂: S ≈ 53.5 wt%, Fe ≈ 46.5 wt%) to more oxidized forms resembling iron oxides with Fe = 60 wt% and O = 30–35 wt% (Figures 3 and 5; Table S2). Only pyrites with low oxygen signal (generally <1 wt% O) were analyzed for δ³⁴S values. Small (<25 μm) pyrite grains are subhedral to anhedral and comprise 3%–10% of the area of organic-rich mat textures (Figures 4g–i and 5m,n). In contrast to the Sukhaya Tunguska Formation samples, these organic-rich sections are heavily dolomitized in sample P4705-A, with silica only representing ~5% of the area (Figures 4i and 5o,p). Large pyrite grains (>25 μm) occur along the boundary of silicified vugs or channels (Figure 5m–p). These grains can be euhedral and primarily pyrite (e.g., Figure 5o) or framboidal grains that show evidence for both overgrowths and differential oxidation of both framboid boundaries and interiors (e.g., Figure 5p). Large pyrite grains were also found in a silica-rich portion of a mat fragment (Figure 5i,j). These grains have morphologies consistent with formation by aggregation of smaller particles and have either anhedral (e.g., Figure 5i) or overgrown, subhedral framboidal shapes (e.g., Figure 5k).

In all three samples, mean δ³⁴S values of individual sulfide grains range around the sample bulk δ³⁴S values (Figure 6), which were determined from sulfide extracted via the chromium reduction method (Canfield et al., 1986). While generally assumed to reflect pyrite, the chromium reduction method recovers sulfide that is operationally defined as the chromium reducible sulfide (CRS) pool because it recovers sulfide from both pyrite and sphalerite (as well as other metal sulfide...
minerals) if no acid pre-treatment step is included. Individual pyrite grain $\delta^{34}$S values in Sukhaya Tunguska Formation sample K95-101 range from $-11.6 \pm 3.1\%$ to $11.7 \pm 3.2\%$, with a mean SIMS $\delta^{34}$S value within the sample of $-4.5\%$ ($\sigma = 5.7\%, n = 18$). The bulk $\delta^{34}$S$_{CRS}$ value for K95-101 is $0.6 \pm 0.2\%$. The other sample from the Sukhaya Tunguska Formation, K95-103, has a slightly lower bulk $\delta^{34}$S$_{CRS}$ value of $-5.1 \pm 0.2\%$. Correspondingly, the mean individual grain $\delta^{34}$S value for all sulfide grains in K95-103 is slightly lower at $-8.6\%$ ($\sigma = 6.7\%, n = 24$), with a range of $\delta^{34}$S values between $-22.6 \pm 1.5\%$ and $7.6 \pm 2.1\%$. The Draken formation sample, P4705-A, has a bulk $\delta^{34}$S$_{CRS}$ value of $-28.0 \pm 0.2\%$, a bimodal distribution of individual grain $\delta^{34}$S values, a mean SIMS $\delta^{34}$S value of $-34.6\%$ ($\sigma = 15.9\%, n = 46$), and an individual grain $\delta^{34}$S range of $-61.0 \pm 4.4\%$ to $-6.4 \pm 1.9\%$. The concentrations of CRS are $230 \pm 20$ ppm S in K95-101, $301 \pm 20$ ppm S in K95-103, and $516 \pm 20$ ppm S in P4705-A. Total organic carbon concentrations are $0.6 \pm 0.1$ wt% in K95-101, $0.4 \pm 0.1$ wt% in K95-103, and $0.2 \pm 0.1$ wt% in P4705-A.

4 | DISCUSSION

4.1 | Fine-scale $\delta^{34}$S trends

These two sample suites were selected for their similar diagenetic histories of early dolomitization, silicification, and exceptional mat
fabric preservation, as well as their moderate temperature history and lack of evidence for metasomatic alteration. Although separated in age, they sample comparable environments that underwent similar early diagenetic silicification and preservation of early pyrite. Fine-scale $\delta^{34}S$ analyses and petrographic observations of sulfide minerals reveal a well-defined relationship between populations of small versus large sulfide grains and $\delta^{34}S$ values (Figure 7). Smaller pyrite grains (<25 µm), which are more common in organic-rich portions of mat fragments (Figures 4 and 5a–d,m,n), tend to have higher $\delta^{34}S$ values. Larger pyrite and sphalerite grains (>25 µm), found along the boundaries of silicified pore spaces (Figure 5e–f,m,n) and in mat fragments that have been almost completely replaced by silica (Figure 5i,j), tend to have lower $\delta^{34}S$ values. These two groups of pyrite grains are reflected in the bimodal distribution of $\delta^{34}S$ values in the histogram of P4705-A (Figure 6c). The distribution of $\delta^{34}S$ values in the Sukhaya Tunguska Formation samples (Figure 6a,b) does not show a bimodal pattern because the number of large sulfide grains in the sample is small (i.e., one >25 µm pyrite grain was analyzed in K95-101 and three >25 µm sphalerite grains were analyzed in K95-103). We interpret the relationship between
δ³⁴S values in small versus large sulfide grains to reflect different generations of sulfide formation.

The small pyrite grains (<25 μm) with relatively high δ³⁴S values represent one generation of sulfide formation. If we consider δ³⁴S_sulfide trends from modern microbial mats and consortia, where δ³⁴S_sulfide values tend to be higher near redox boundaries due to high average csSRR and low εSR values (i.e., the differential metabolic activity trend in Figure 1; Fike et al., 2008, 2009; Wilbanks et al., 2014), we could infer that these small pyrite grains represent a first generation of pyrite formation near the surface of the mat (i.e., location #1 in Figure 1). However, high δ³⁴S_sulfide values also occur deeper within the sediments of many modern marine sedimentary environments (i.e., location #2 in Figure 1) as a result of closed-system Rayleigh isotope effects during early diagenesis. Closed-system Rayleigh isotope fractionation trends have not been documented in modern marine microbial mats because sulfate concentrations are high (i.e., >28 mm) and only a small portion of the sulfate reservoir is consumed (<1%; Fike et al., 2009). Similarly, incubations with low sulfate concentrations (0.2–1 mm) using mat samples from modern marine environments did not show Rayleigh isotope fractionation trends. The lack of closed-system isotope effects was attributed to low rates of sulfide production (Fike et al., 2009), likely because the microbial communities in the mats were not adapted to low sulfate concentrations (Bradley et al., 2015). However, it is possible that the microbial mat communities associated with mat fragments in the studied samples were adapted to low sulfate conditions that were common in Proterozoic oceans (Canfield, 2004; Kah, Lyons, & Frank, 2004) and therefore could consume a sufficiently large portion of the sulfate reservoir to generate a Rayleigh fractionation δ³⁴S_sulfide trend; such a pattern has been documented in late Archean sediments (Fischer et al., 2014). Thus, high δ³⁴S_sulfide values in the small pyrite grains could either capture the signature of sulfide near the redox boundary at the surface of the mat or at greater depths where a large fraction of the sulfate reservoir was consumed. It is also possible that high δ³⁴S values in the small pyrite grains reflect local microniches with variable but localized influences of Rayleigh isotope fractionation on δ³⁴S_sulfide values, which could occur at any depth in the mat.

Using petrographic context and knowledge of pyrite formation mechanisms, we can attempt to differentiate between these two potential locations of small (<25 μm) pyrite formation. First, the small pyrites are evenly disseminated in organic-rich groundmass with microtextures such as wavy crinkled laminae (Figures 4d,e, and 5a–d) sponge pore fabrics (Figure 4a,b,g,h) and adjacent shrinkage cracks (Figure 5a–f) that are characteristic of microbially induced sedimentary structures (Noffke, 2009). Thus, although samples K95-101 and K95-103 come from a different formation than P4705-A, all of the studied samples show a petrographic association between small pyrite grains and organic-rich fragments with mat textures. Similar types of small pyrite grains in kerogen-rich matrix material in Archean microbial carbonates were interpreted to be early diagenetic pyrites (Kamber & Whitehouse, 2007). The early diagenetic origin of the small pyrite grains in our samples is further
supported by the excellent fabric retention and preservation of microfossils in both the Sukhaya Tunguska and Draken microbial mats, which suggests early diagenetic silicification and/or dolomitization (Mendelson & Schopf, 1982; Fairchild et al., 1991; Knoll et al., 1991; Sergeev et al., 1997).

Second, empirical and theoretical studies of pyrite formation suggest that pyrite is formed near redox boundaries (Rickard, 1975; Wilkin, Barnes, & Brantley, 1996; Wilkin, Arthur, & Dean, 1997) and that small pyrite grains, in particular, are formed in locations with rapid redox oscillation (Wilkin et al., 1996). Therefore, it is likely that the early diagenetic, small pyrite grains were formed near the surface of the mat where diurnal cycles promote redox oscillation via vertical migration of the chemocline and the presence of intermediate sulfur species promotes pyrite formation. Biotic or abiotic (i.e., by reaction with H₂S) reduction in iron oxides would also promote pyrite formation near the redox interface. It is possible that pyrite growth continued during burial upon the grains nucleated near the mat surface, and this may account for some of the inter- and intragrain range in the SiMS δ³⁴S values of the <25 μm pyrite grains. The intragrain δ³⁴S range could also reflect Rayleigh isotope effects occurring in microenvironments. We cannot completely rule out pyrite formation in deeper portions of the mat where δ³⁴S_sulfide values were high due to significant consumption of the sulfate reservoir. However, finely disseminated pyrite textures are believed to be the result of nucleation bursts and subsequent decreases in pyrite saturation (Rickard, 2012), which likely occurs in locations with frequent redox oscillations. Thus, based on these insights about how pyrite formation processes impact pyrite textures and the association with organic-rich mat fragments (Figures 4 and 5a–d,m,n), we posit that the small pyrite grains (<25 μm) with high δ³⁴S_pyrite values were formed near a redox boundary (i.e., location #1 in Figure 1). In this location, low apparent S-isotope fractionations between sulfate and sulfide are a result of either high average csSRR associated with the respiration of photosynthetically produced labile organic matter and low sulfide concentrations due to active sulfate oxidation or Rayleigh isotope fractionation in microniches.

Large pyrite grains (>25 μm) represent a different generation of sulfide formation in P4705-A, with lower δ³⁴S values than the small pyrites. The δ³⁴S value of the single >25 μm pyrite grain in K95-101 is also slightly lower (~10.5 ± 2.0‰) than the mean δ³⁴S value of the <25 μm pyrite grains (~4.2‰, σ = 5.7‰, n = 17). Only a single >25 μm pyrite grain was observed, and therefore, we will focus this discussion on P4705-A, which has a more robust data set; however, we note that grain size and δ³⁴S trends in K95-101 are consistent with those observed in P4705-A. These large (>25 μm) grains in P4705-A are located along the boundaries of silicified vugs or channels (Figure 5m–p) as well as channelized portions of silicified mat fragments (Figure 5i–l). The pore spaces are thought to form either as a result of primary porosity in the microbial mats or from the dissolution of precursor materials under the influence of meteoric or deep-burial fluids (Choquette & Pray, 1970). While we cannot completely rule out the possibility that these grains formed during deep burial from hydrothermal (or other) fluids, their location along the edges of silicified void spaces and mat fragments implies their formation predates silicification.

It is also unlikely that these grains formed from hydrothermal fluids because the δ³⁴S values of the large (>25 μm) pyrite grains are lower than hydrothermally derived sulfide (i.e., 1‰ to 3‰; Arthur, 2000) as well as the small pyrite grains, which could have served as a source of sulfide from dissolution in hydrothermal fluids. It is also possible that dissolution of pyrite in adjacent sediments could be a source of sulfide. However, fractures cross-cut the large pyrite grains (Figure 5i,j) and silicified pore spaces (Figure 5m,n) and therefore postdate the formation of the large pyrite grains and silicified pore spaces. This implies that, if sulfide from adjacent sediments was transported through these fractures, it did not contribute to the formation of the large pyrite grains. Instead, it is likely that the large pyrite grains formed from early diagenetic pore fluids flowing through the vugs or channels at depth within the mat (i.e., location #3 in Figure 1) or, in the case of reworked mat fragments, after the mats had been ripped up by storms and redeposited in more offshore settings. Additional support for a diagenetic rather than hydrothermal sulfide source is the excellent fabric retention and preservation of microfossils associated with these samples that indicates early dolomitization and silicification (Knoll et al., 1991; Fairchild et al., 1991). The refractory nature of organic carbon deep within the mat and/or entrained and redeposited mat fragments could lead to low average csSRR, resulting in high ε₃⁴S values and low δ³⁴S values. Nutrient limitation or co-limitation may also play a role in suppressing average csSRR (Bradley et al., 2015), but is not necessarily required to explain the fine-scale δ³⁴S patterns. In sum, the large pyrite grains tend to have lower δ³⁴S values because they record δ³⁴S values of sulfide produced at low average csSRR deep within sediments in locations exposed to fluids promoting pyrite formation.

The large (>35 μm) sphalerite grains in K95-103 represent yet another stage of sulfide mineral formation. It is important to note that sphalerite grains were only detected in one sample, and therefore, we have only observed evidence for this stage of sulfide mineral growth in sample K95-103. In K95-103, the sphalerite grains are located along the edges of pore spaces (Figure 5e–h) and have low δ³⁴S values, while the smaller pyrite grains are evenly distributed in organic-rich mat fragments (Figures 4d,e and 5c,d) and have higher δ³⁴S values. Pyrite and sphalerite have different thermodynamic relationships, reaction kinetics, and redox reaction pathways that influence how we interpret the δ³⁴S values of the sphalerite grains. In particular, sphalerite is thought to form prior to pyrite due to faster water exchange reaction kinetics associated with Zn (Morse & Luther, 1999). Thus, we might expect that the sphalerite grains formed before both the small and the large pyrite grains. In fact, precipitation of sphalerite has been documented in modern biofilms (Labrenz et al., 2000) and these sphalerite grains share some characteristics with the sphalerite grains in K95-103. Sphalerite grains in the modern biofilms have microtextures formed by aggregation of smaller particles, which are a few tens of micrometers in size, have spatially heterogeneous distribution patterns, and are found in organic-rich microenvironments (Druschel, Labrenz, Thomsen-Ebert, Fowle, & Banfield, 2002). Yet, sphalerite grains are not found within the organic-rich mat fragments in K95-103. Instead, the sphalerite grains are primarily found along the boundary between organic-rich mat fragments and laminoid fenestrae (Figure 5e–h).
Laminoid fenestrae are commonly formed in intertidal to supratidal settings from wetting and drying of microbial mats and/or carbonate mud that leads to separation of the mat surface from the adjacent sediment (Shinn, 1968; Davies, 1970; Logan, Hoffman, & Gebelein, 1974; Hardie & Ginsburg, 1977). Thus, the sphalerite grains likely formed as a result of diagenesis that occurred after the wet-dry cycles that led to the development of the laminoid fenestrae.

Shale and iron formation records suggest that Proterozoic marine zinc concentrations were similar to modern marine levels (Scott et al., 2013; Robbins et al., 2013), and so seawater zinc would be available for sphalerite formation. We do not find sphalerite grains finely dispersed in the organic-rich matrix materials, which necessarily formed prior to the development of laminoid fenestrae during desiccation. It is possible that organic complexation or microbial uptake of zinc in the mats depleted the porewaters of available zinc and inhibited the formation of zinc sulfides within the organic-rich matrix. Then, during the time periods between the re-wetting of the desiccated mat and the establishment of a new phototrophic microbial mat community, diminished zinc uptake by microorganisms allowed for higher concentrations of zinc. This laid the foundation for sphalerite formation along the boundary between fenestrae and organic-rich mat fragments that served as an organic carbon substrate for MSR. This would require a large quantity of zinc and therefore repeated wet-dry cycles or introduction of new zinc by flushing with seawater under continued wet conditions over a long time period before the re-establishment of phototrophic microbial mat communities. The formation during repeated wet and dry cycles is consistent with the large range of $\delta^{34}\text{S}$ values measured in sphalerite grain transects (i.e., up to 117.4‰; Table S1) because differential rates of MSR, diffusion and advection, evaporation and wetting (which would concentrate or dilute sulfate, sulfide, and zinc), and Rayleigh isotope fractionation in microenvironments would result in heterogeneous $\delta^{34}\text{S}_{\text{sulfide}}$ values over the course of mineral growth. Despite the large range of $\delta^{34}\text{S}$ values in sphalerite grain transects, mean $\delta^{34}\text{S}$ values of sphalerite grains are similar (Figure 6b). Thus, mean $\delta^{34}\text{S}$ values of sphalerite grains capture an integrated signal of this stage of sulfide growth in this sample. The heterogeneous intragrain $\delta^{34}\text{S}$ values imply that microbial sulfate reducers were able to utilize organic carbon in the mat or entrained in fluids in previously desiccated mats to reduce sulfate to sulfide in an unsteady diagenetic regime.

Based on the hypothesized formation mechanisms for the different generations of sulfide minerals, we can draw some inferences about the environmental history of the microbialite samples and the $\delta^{34}\text{S}_{\text{sulfide}}$ signatures of each stage of sulfide formation. In this discussion, it is important to note that the lagoonal depositional setting of the microbial mats (i.e., prior to entrainment and redeposition) could have had varying degrees of connectivity with the ocean and/or rates of evaporation during the lifetime of the mat. Evaporation of water does not directly affect the isotopic composition of sulfate, but it does alter the size of the local sulfate reservoir. Enhanced evaporation of seawater would lead to higher sulfate concentrations and a decreased potential of Rayleigh isotope fractionation during MSR to drive sulfate to high $\delta^{34}\text{S}$ values. In contrast, the input of freshwater from land or precipitation would lead to lower sulfate concentrations and a greater potential for sulfate to become $^{34}\text{S}$-enriched due to MSR. Or, put another way, $\delta^{34}\text{S}_{\text{sulfate}}$ values can vary to a greater extent when sulfate levels are low due to the dilution of seawater by freshwater or precipitation. In what follows, we take S-isotopic analyses of late Mesoproterozoic and early to mid-Neoproterozoic gypsum and anhydrite $\delta^{34}\text{S}$ values of ca. 20 ± 5‰ to approximate the isotopic composition of seawater sulfate (Kah, Lyons, & Chesley, 2001; Canfield, 2004), but note that the sulfate concentration and potential for $^{34}\text{S}$ enrichment in sulfate may vary as a function of lagoon hydrology, and this also has potential to affect $\delta^{34}\text{S}_{\text{sulfide}}$ values.

In K95-101, pyrite grains are predominantly small (<25 μm) and have high $\delta^{34}\text{S}$ values (~11.6 ± 3.1‰ to 11.7 ± 3.2‰). We suggest that these grains capture the signature of sulfide near the redox boundary based on their even distribution in organic-rich mat fragments, excellent fabric retention, and preservation of microfossils in these samples that indicates early lithification (Mendelson & Schopf, 1982; Fairchild et al., 1991; Knoll et al., 1991; Sergeev et al., 1997), and theoretical models of how environmental conditions influence pyrite morphology (Rickard, 2012). Although this sample could have gone through wet-dry cycles and have been impacted by fluid flow during deep burial, we do not see petrographic evidence for sulfide formation during these stages. Thus, the bulk $\delta^{34}\text{S}_{\text{CRS}}$ value of K95-101 records the $\delta^{34}\text{S}$ signature of pyrites formed adjacent to the redox boundary in the upper portions of the mat during early diagenesis. The $\delta^{34}\text{S}$ values of the majority of the individual pyrite grains are lower bulk $\delta^{34}\text{S}_{\text{CRS}}$ value, which may indicate that the pyrite grains that were too small to be measured by our SIMS method (i.e., <5 μm) have higher $\delta^{34}\text{S}$ values. The $\delta^{34}\text{S}$ range in individual pyrite grains of 23.3‰ could reflect localized Rayleigh isotope fractionation due to significant sulfate consumption during early diagenesis or heterogeneity of $\varepsilon_{\text{SR}}$ values in early diagenetic microniches in the upper portions of the mat environment.

Considering the average of the small pyrite $\delta^{34}\text{S}$ values, the S-isotope difference between this small pyrite population and estimated coeval seawater sulfate is 24.5‰, which is similar to the S-isotope difference between the bulk $\delta^{34}\text{S}_{\text{CRS}}$ value and inferred coeval seawater sulfate (19.4‰). This implies that, in absence of significant Rayleigh isotope fractionation of the overlying sulfate reservoir, the average S-isotope fractionation between sulfate and sulfide ($\Delta^{34}\text{S}$) near the upper redox boundary was ~20‰ in this 1,000 Ma microbial mat, which should be considered a minimum value because Rayleigh isotope fractionation in microenvironments would decrease the apparent $\delta^{34}\text{S}$ offset.

K95-103 contains both small (<25 μm) pyrite grains and large (>35 μm) sphalerite grains. Applying the criteria given in the previous paragraph for the formation of small pyrite grains in K95-101 to this sample, we similarly suggest that the small pyrite grains formed first near the redox boundary close to the mat surface. Subsequent wet-dry cycles lead to the establishment of laminoid fenestrae (Figure 5e–h) and subsequent formation of large sphalerite grains. The bulk $\delta^{34}\text{S}_{\text{CRS}}$ value represents a mixed signal of two sulfide populations formed during different stages of diagenesis and is offset from the coeval seawater sulfate $\delta^{34}\text{S}$ value by 25.1‰. On its own, this bulk value may not be that meaningful because it does
not represent a single stage of sulfide mineral formation. However, the fine-scale analyses allow us to demonstrate that the mean small pyrite $\delta^{34}S$ value is 27.1‰ lower than the estimated $\delta^{34}S$ value of coeval seawater sulfate. This value, which approaches the $\Delta^{34}S$ value near the upper redox boundary, is similar to the $\Delta^{34}S$ value of 24.5‰ determined for the other 1,000 Ma sample (K95-101). The mean sphalerite $\delta^{34}S$ value is 39.4‰ lower than the estimated $\delta^{34}S$ value of coeval seawater sulfate. It is possible that the bulk $\delta^{34}S$ signature of these two sulfide populations could be determined from sequential extraction of sphalerite by acid extraction and pyrite by chromium reduction (see Morse & Luther, 1999 for discussion). Unfortunately, there is no corresponding way to chemically separate different generations of pyrite grains. Nonetheless, sequential separation of these operationally defined sulfide pools could aid in distinguishing populations of sulfide minerals with distinct $\delta^{34}S$ values. Supplementing the sequential extraction with fine-scale analyses has the added benefit of connecting the different generations of sulfide mineral formations to the environmental history of the sample. Lastly, P4705-A contains two generations of pyrite formation represented by small ($<$ 25 $\mu$m) and large ($>$ 25 $\mu$m) pyrite populations. Similar to the Sukhaya Tunguska samples, even distribution in mat fragments, retention of mat textures, and excellent microfossil preservation (Knoll et al., 1991) along with pyrite formation theory (Rickard, 2012) imply that small pyrite $\delta^{34}S$ values capture the $\delta^{34}S$ signature of sulfide adjacent to the redox boundary at the surface of the mat. In contrast, the observation that large pyrite grains occur along the edges of silicified void spaces implies that large pyrite $\delta^{34}S$ values reflect the $\delta^{34}S$ value of sulfide in pore fluids during later stages of burial or after entrainment of mat fragments and redeposition. These two populations are easily distinguished in the frequency distribution plot of SIMS $\delta^{34}S$ values (Figure 6c), and the bulk $\delta^{34}S_{Crs}$ value represents an average of these two populations. While the bulk $\delta^{34}S_{Crs}$ value is 48.0‰ lower than the estimated $\delta^{34}S$ value of coeval seawater sulfate, this $S$-isotope difference is 46.4‰ and 66.2‰ for the small and large pyrite populations, respectively. The difference in apparent $S$-isotope fractionation between the small and the large pyrite grains could indicate two different paleoenvironmental trends: (i) If small pyrite grains were precipitated in the upper portions of the mat and large pyrite grains in deeper portions, then that would imply the $\delta^{34}S_{sulfide}$ gradient was similar to what is observed in modern microbial mats (Fike et al., 2008, 2009); (ii) If small pyrite grains were precipitated in the upper portions of the mat and large pyrite grains were precipitated after entrainment and redeposition, then the difference in apparent $S$-isotope fractionation could be due to different average $csSRR$, different amounts of localized system openness, or differences in $\delta^{34}S$ values of the sulfate reservoir in lagoonal versus deeper water depositional settings. At this time, we are unable to differentiate between these options. This 750 Ma sample has a higher $\Delta^{34}S$ value near the upper redox boundary than the 1,000 Ma samples, inferred from the mean $\delta^{34}S$ value of the small ($<$ 25 $\mu$m) pyrite grains and assuming minimal influence of Rayleigh isotope fractionation on the overlying sulfate reservoir. This could be due to higher ambient sulfate concentrations, increasing $\epsilon_{MSR}$ and/or depressing closed-system behavior during MSR, enhanced oxidative $S$ cycling, or differences in organic carbon and nutrient availability.

The inferred environmental histories of sulfide formation in these microbialite samples represent our best attempt to synthesize the information provided by petrography, elemental mapping, and fine-scale $\delta^{34}S$ analyses. We cannot completely rule out other processes. For example, closed-system Rayleigh isotope effects are likely influencing $\delta^{34}S$ values on a local scale, which can explain some of the $\delta^{34}S$ variability within both individual crystals and the small and large pyrite pools. However, we do not see evidence for significant pyrite formation in where $\delta^{34}S_{sulfide}$ values are very high due to consumption of the sulfate reservoir in a closed system (i.e., location #2 in Figure 1) because petrographic evidence (Figure 4, 5a–d, m–n) suggests that small pyrite grains are formed near the surface (i.e., location #1 in Figure 1) and large pyrite grains (Figure 5i–p) are formed deeper in the mat where average $csSRR$ are low (i.e., location # 3 in Figure 1). Given that fluid inclusion data indicate marine sulfate levels, which were 50%–90% lower in the Proterozoic than modern sulfate concentrations (Lowenstein, Hardie, Timofeeff, & Demico, 2003; Spear et al., 2014), and organic carbon availability was high in these mats, we would expect near-complete consumption of the sulfate reservoir deep in the mat. If near-complete consumption of the sulfate reservoir did lead to high $\delta^{34}S_{sulfide}$ values at depth, this signature was not captured in the pyrite grains, perhaps because iron limitation inhibited significant pyrite formation in this location. Late-stage alteration also has the potential to affect $\delta^{34}S$ values. However, three observations imply that the effects of late-stage (i.e., catagenetic) alteration are minimal in our samples: (1) the lack of pyrite in veins or other alteration textures, (2) petrographic evidence that veins cross-cut sulfide grains and mat fragments (Figure 5), and (3) the excellent fabric retention and preservation of microfossils indicating early lithification. In sum, the textural and geochemical information provided by these techniques gives us confidence in our reconstructions of the evolution of sulfur cycling in these ancient microbial environments.

4.2 Implications for the sedimentary sulfide sulfur isotope record

The $\delta^{34}S$ record of sedimentary sulfides provides important information about the oxygenation of the ocean and atmosphere through Earth history (e.g., Canfield & Farquhar, 2009; Figure 8). In particular, the increase in the range of $\delta^{34}S_{pyrite}$ values toward the present has been linked to oxygenation of Earth surface environments. There are two intriguing transitions: (i) at ~2.4 Ga the range of $\delta^{34}S$ values of sedimentary pyrites increases from $\sim 20$‰ to $\sim 50$‰ and (ii) at roughly 635 Ma there is another increase in the $\delta^{34}S_{pyrite}$ range. The first transition has been linked to the Great Oxidation Event and an associated increase in sulfate concentrations due to oxidative weathering of pyrite on land and/or a decrease in the efficiency of the pyrite burial sink due to diminished iron availability of the ocean. The second transition has been linked to the expansion of well-oxygenated environments through the oceans and associated biogeochemical changes that would increase the range of preserved $\delta^{34}S$ values, including
enhanced oxidative sulfur cycling and microbial disproportionation (Canfield & Teske, 1996), decreased average cSRR, and/or increased system openness due to expanded aerobic respiration of more labile organic matter (Fike et al., 2015), or the physiologically related onset of bioturbation (Canfield & Farquhar, 2009; Aller, 2014). These interpretations are consistent with other types of geological evidence that point to transitions in Earth surface oxygenation at ~2.4 Ga and ~635 Ma. Yet, to a first order, this interpretation rests on the assumption that bulk δ34SCRS values represent an environmentally meaningful average of discrete δ34S values from individual pyrite grains.

The data presented here and elsewhere indicate that the range of δ34S values measured within a single sample can be as great as or larger than the range of bulk δ34SCRS values observed globally for the same time interval (Figure 8) and may reflect different stages of sulfide mineral growth. The bulk δ34SCRS value is thus a signal that integrates the isotopic signatures imprinted at different stages of pyrite (and other sulfide mineral) formation. Apparent secular variation in bulk δ34SCRS values may thus reflect shifts in the range of individual pyrite grain δ34S values and/or changes in the proportion of sulfide grains formed at different stages of sulfide formation. In turn, these patterns will be influenced by the processes that are generally considered to impact bulk δ34SCRS values, such as changes in sedimentation rate (Claypool, 2004; Pasquier et al., 2017), depositional environment, redox conditions, iron and organic carbon availability, or the size and isotopic composition of the seawater sulfate reservoir. The data presented here are consistent with fine-scale δ34Spyrite trends in Paleoproterozoic marine sediments (Williford, Van Kranendonk, Ushikubo, Kozdon, & Valley, 2011) indicating that, on a fine scale, the increase in the range of δ34Spyrite values beyond ~50‰ occurs well before 635 Ma (Figure 8).

Yet, this range is not widely reflected in bulk δ34Spyrite values. This may be due to low marine sulfate concentrations that suppress the magnitude of the apparent sulfur isotope fractionation between sulfate and sulfide that is preserved in marine sediments (Gomes & Hurtgen, 2013). Or it may be that the types of environments that favored low average cSRR, oxidative sulfur cycling, microbial disproportionation, or perhaps even bioturbation by benthic organisms were not significant enough globally to affect the sedimentary pyrite δ34S record until after ~635 Ma (Fike et al., 2015). Ancient microbial mats and biofilms are poorly represented in the sedimentary sulfide δ34S record (Canfield & Farquhar, 2009; Figure 8). Future studies that compare bulk and fine-scale sedimentary sulfide δ34S patterns in ancient microbial mats and biofilms to continental shelf or other types of marine sediments will provide insight into the important preservational biases that influence δ34S patterns in different depositional environments. Studies of bulk and fine-scale δ34S trends in the context of petrographic observations that provide information about different generations of sulfide formation will further contribute to our knowledge of the evolution of redox conditions throughout the Proterozoic and other periods in Earth history.

More generally, on the time scale sampled in many geological investigations (i.e., tens of thousands of years to hundreds of millions of years) bulk δ34SCRS values preserve broad signals of environmental change. However, because bulk δ34SCRS values integrate many biogeochemical processes, the application of fine-scale geochemical and petrological methods to discern different generations of metal sulfide formation is key to interpreting these records. S-isotopes do suggest significant differences between our 1000 Ma and 750 Ma samples, whether viewed in terms of bulk or fine-scale analyses. Fine-scale δ34Spyrite studies serve as a useful complement to disentangle δ34Spyrite signatures associated with different stages of pyrite growth that can be used to extract additional paleoenvironmental information from geological materials. Thus, the combination of bulk and fine-scale geochemical analyses with petrographic observations allows us to better interpret the marine sedimentary record in order to explore the coupled evolution of life and the Earth surface system.

5 | CONCLUSION

Fine-scale δ34S patterns in sulfide minerals can reveal information about variable environmental conditions and differential metabolic activity of microbial communities present at different depths in ancient microbial mats. We examined microbialite samples from two
Proterozoic formations to evaluate whether fine-scale $\delta^{34}S_{\text{sulfide}}$ patterns in modern microbial mats and consortia are preserved in the geological record. We also explored how closed-system Rayleigh isotope fractionation or post-depositional alteration influence fine-scale $\delta^{34}S_{\text{pyrite}}$ patterns. We find that small (<25 $\mu$m) pyrite grains tend to have higher $\delta^{34}S$ values than large (>25 $\mu$m) pyrite and (>35 $\mu$m) sphalerite grains. Based on these patterns, elemental mapping, petrographic information, and comparison with modern microbial mats, we suggest that there are three generations of sulfide formation in the studied samples: (i) small (<25 $\mu$m) pyrite grains that formed near the redox interface with high $\delta^{34}S$ values due to high average csSRR, (ii) large (>35 $\mu$m) sphalerite grains with low $\delta^{34}S$ values formed during repeated wet-dry cycles from sulfide produced by MSR under an unsteady diagenetic regime (i.e., variable rates of csSRR, diffusion, and advection), and (iii) large (>25 $\mu$m) pyrite grains formed during later stages of burial with low $\delta^{34}S$ values due to formation from a pool of sulfide produced by slower average csSRR. Each sample contains grains from one or two of these stages of sulfide growth. Bulk $\delta^{34}S_{\text{pyrite}}$ values reflect a weighted average of the different sulfide groups. Fine-scale isotopic studies coupled with elemental mapping and petrographic observations can thus reveal important information about microbiolally mediated reactions occurring at different stages of burial. The extension of this approach to other time periods and sedimentary materials will enhance our ability to extract paleoenvironmental information from the geological record.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

ACKNOWLEDGMENTS

The authors thank the NASA Astrobiology Institute, Washington University, and Massachusetts Institute of Technology (MIT) as well as the NSF Major Research Instrumentation (EAR #1229370) and DOE/BER Bioimaging Technology (DE-SC0014613) awards to DAF for supporting this work. We are also grateful for the technical support of Nilanjan Chatterjee (MIT) for electron microprobe analyses, Erin Beirne (Harvard University) for isotope ratio mass spectroscopy measurement, Stephanie Moore (Washington University) for carbon concentration analyses, Michael Krawczynski (Washington University) for access to petrographic microscopes, and Harry-Luke McClelland (Washington University) for help constructing figures. We thank David Johnston, Andrew Masterson (Harvard), Jennifer Houghton, and Jocelyn Richardson (Washington University) for comments that improved the manuscript.

REFERENCES


SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.

How to cite this article: Gomes ML, Fike DA, Bergmann KD, Jones C, Knoll AH. Environmental insights from high-resolution (SIMS) sulfur isotope analyses of sulfides in Proterozoic microbialites with diverse mat textures. Geobiology. 2017;00:1–18. https://doi.org/10.1111/gbi.12265