

Phase, morphology, elemental composition, and formation mechanisms of biogenic and abiogenic Fe-Cu-sulfide nanoparticles: A comparative study on their occurrences under anoxic conditions

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ABSTRACT

We report on a systematic study on the physicochemical attributes of synthetic Fe-Cu-sulfide nanoparticles (NPs) precipitated under conditions similar to the anoxic, low-temperature aqueous, sedimentary, soil, and subsurface environments where these NPs have been repeatedly identified. Characterizing the basic attributes of these NPs is the first step in understanding their behaviors in various processes including in the bio-availability of essential and toxic metals, environmental remediation, and resource recovery. Abiotic experiments are compared to biotic experiments in the presence of the sulfate-reducer *Desulfovibrio vulgaris* to elucidate biological controls on NP formation. First, the single-metal end-member NPs are determined by precipitation in a solution containing either aqueous Fe(II) or Cu(II). Limited differences are observed between biogenic and abiogenic precipitates aged for up to one month; the Fe-only experiments resulted in 4–10 nm mackinawite (FeS) NPs that aggregate to form nanosheets up to ~1000 nm in size, while the Cu-only experiments resulted in mixtures of covellite (CuS) NPs comprised of <10 nm fine nanocrystals, 20–40 × 6–9 nm nanorods, and ~30 nm nanoplates. The crystal sizes of biogenic mackinawite and covellite are, respectively, larger and smaller than their abiogenic counterparts, indicating a mineral-specific response to biological presence. Structural defects are observable in the fine nanocrystals and nanorods of covellite in both biogenic and abiogenic experiments, indicative of intrinsic NP instability and formation mechanism via particle attachment. In contrast, covellite nanoplates are defect free, indicating high stability and potentially rapid recrystallization following particle attachment. Next, mixed-metal sulfide NPs are precipitated at variable initial aqueous Fe-to-Cu ratios (2:1, 1:1, and 1:5). With an increasing ratio of Fe-to-Cu, Fe-rich covellite, nukundamite (Cu_{5.5}FeS_{6.5}), chalcopyrite (CuFeS₂), and Cu-rich mackinawite are formed. The Fe-rich covellite NPs are larger (100–200 nm) than covellite precipitated in the absence of Fe, indicating a role for Fe in promoting crystal growth. Chalcopyrite and nukundamite are formed through the incorporation of Fe into precursor covellite NPs while retaining the original crystal morphology, as confirmed by doping a covellite suspension with aqueous Fe(II), resulting in the formation of chalcopyrite and nukundamite within days. Additionally, in the biological systems, we observe the recrystallization of mackinawite to greigite (Fe₃S₄) after six months of incubation in the absence of Cu and the selective formation of chalcopyrite and nukundamite at lower initial Fe-to-Cu ratios compared to abiotic systems. These observations are consistent with NP precipitation that are influenced by the distinct (sub)micro-environments around bacterial cells compared to the bulk solution. Comparative TEM analyses indicate that the synthetic NPs are morphologically similar to NPs identified in natural environments, opening ways to studying behaviors of natural NPs using experimental approaches.

Keywords: Metal sulfide nanoparticle, mackinawite, covellite, chalcopyrite, greigite, biomineral

INTRODUCTION

Metal sulfide nanoparticles (NPs) such as FeS, CuS, and ZnS are being increasingly recognized as important and understudied components in the biogeochemical cycling of sulfur and heavy metals in various natural settings (e.g., Luther and Rickard 2005;

Hochella et al. 2008; Sharma et al. 2015). In environments such as floodplain soils and estuaries, the precipitation and dissolution of metal sulfide NPs under variable redox conditions affect the sequestration and release of toxic metals into the ecosystem and surrounding waters (Weber et al. 2009a; Fulda et al. 2013; Hofacker et al. 2013; Niu et al. 2018). In heavy-metal contaminated riverbeds and subsurface aquifer sediments, the phase and morphology of metal sulfide NPs formed in situ greatly affect the efficiency of environmental remediation via heavy metal im-

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mobilization (Labrenz et al. 2000; Moreau et al. 2004; Hochella et al. 2005; Qafoku et al. 2014; Xu et al. 2017). The formation of metal sulfide NPs also acts as an important step for the resource recovery of precious metals from mine tailings and industrial processes (Liang et al. 2012; Falagán et al. 2017). Additionally, metal sulfide NPs released from hydrothermal vents on seafloors are important carriers of trace metal nutrients to the ocean (Yücel et al. 2011; Gartman et al. 2014). The ubiquity of metal sulfide NPs, their higher stabilities compared to aqueous phases, and their unique properties such as high surface reactivity compared to their larger counterparts lend them an important role in mediating the biogeochemical cycling of elements and in maintaining the balance of microbial ecology.

The high reactivity of metal sulfide NPs stems from several properties unique at the nanoscale that disappears once minerals reach sizes greater than several to many tens of nanometers. Nanoparticles are characterized by high surface areas and high concentration of atomic defects near the crystal surface that provide active reaction sites (e.g., Banfield and Zhang 2001; Hochella et al. 2008; Eskelsen et al. 2018). These properties may enable certain bacterial metabolisms. For example, Bosch et al. (2012) demonstrated that nano-sized (but not micrometer-sized) pyrite could be utilized by microbes as a source of sulfur for sulfur oxidation coupled to nitrate reduction. Additionally, thermodynamically metastable and potentially highly reactive metal sulfide phases may exist at the nanoscale, as demonstrated by the precipitation of wurtzite (hexagonal ZnS; Moreau et al. 2004; Xu et al. 2016) and α -NiS (Sitte et al. 2013) in low-temperature biological systems. Empirical observations and calculations demonstrate that even at low abundances NPs may be the most important component of a geochemical system as a result of their high reactivities (Hochella et al. 2012).

The reactivity and physicochemical attributes (size, morphology, phase, elemental compositions) of metal sulfide NPs are strongly affected by their formation mechanism(s). *Classical models* (e.g., Benning and Waychunas 2008) describe formation via the addition of simple ions (e.g., aqueous metals) until a sufficient density is reached to form a nucleus. This nucleus tends to be a metastable phase that will dissolve and reprecipitate to form larger and more stable mineral phases in a process termed Ostwald ripening. In the past two decades, an alternative pathway for crystal formation in the form of *particle attachment* has come into light (reviewed by De Yoreo et al. 2015). Through this pathway, crystal nucleation and growth are mediated by attachment of smaller primary units, which may range from sub-nanometer sized aqueous clusters to NPs tens of nanometers in size. The particle attachment pathway tends to be dominant at high supersaturation when nucleation events are common (De Yoreo et al. 2015), a condition that is easily satisfied for metal sulfides due to their low solubilities (Rickard and Luther 2006). The nucleation of metal sulfides most likely results from assembly of aqueous clusters (e.g., Fe_2S_2 , Cu_3S_3 , Zn_3S_3 less than 1 nm in size; Luther and Rickard 2005), followed by growth via oriented attachment of discrete nanocrystals (Pileni et al. 1997; Libert et al. 2003; Moreau et al. 2004; Schliehe et al. 2010; Mullaugh and Luther 2011; Xu et al. 2016). Misalignments during particle attachment may result in defects in the crystal structure, which contribute to the nanoparticle reactivity if the defects are not eliminated

through recrystallization (Penn and Banfield 1998; Echigo et al. 2012; Eskelsen et al. 2018).

The physicochemical attributes of naturally occurring metal sulfide NPs can greatly impact their roles in biogeochemical cycles. It is, however, challenging to characterize natural NPs due to their relatively low abundances and the current technical limitation in separating them from the environment (e.g., Lead and Wilkinson 2006; Caraballo et al. 2015). Metal sulfide NPs also tend to be altered immediately in the presence of oxygen and thus requires strict anoxic sampling and storage procedures prior to analyses. To circumvent these challenges, metal sulfide NPs can instead be precipitated under specified and controlled laboratory conditions. A few generalizations can be made to aid experimental designs that yield results relevant to natural settings. First, the majority of metal sulfides are precipitated in low-temperature anoxic zones as a result of sulfate reduction metabolism by microorganisms (Berner 1984; Picard et al. 2016). Experiments have shown that besides supplying the sulfide source, the presence of microbial cell walls and extracellular organic compounds also play major roles in modifying the formation mechanism, size, shape, phase, solubility, and aggregation state of the precipitates (Gramp et al. 2006, 2010; Moreau et al. 2007; Sitte et al. 2013; Xu et al. 2016; Picard et al. 2018). Second, natural metal sulfides precipitate in complex solution containing other inorganic metals, which may result in co-precipitation of mixed-metal sulfide NPs such as chalcopyrite (CuFeS_2 ; Hochella et al. 2005; Fulda et al. 2013; Gartman et al. 2014), Zn-Fe sulfides (Zbinden et al. 2001), and Cu-Cd-Pb sulfides (Hofacker et al. 2013). Some metal sulfides may also exist within pyrite grains as nano- to micro-sized inclusions (Large et al. 2014; Gregory et al. 2015). These mixed-metal sulfides will likely have different attributes and reactivities compared to single-metal sulfides, which highlights the need for systematic studies on their properties to better constrain their impacts to the environment. Furthermore, any characteristics that are unique to biogenic or abiogenic NPs can be used as biosignatures (Des Marais et al. 2008; Horneck et al. 2016) or to differentiate between natural (often biogenic) and engineered (often abiogenic) NPs that are increasingly being released to the environment (e.g., Nowack et al. 2012).

To this end, we conducted a study to characterize the physicochemical attributes and formation mechanisms of Fe-Cu sulfide NPs precipitated under anoxic conditions in the presence and absence of sulfate-reducing bacteria. X-ray diffraction (XRD) was used in conjunction with transmission electron microscopy (TEM)-based analyses such as selected-area electron diffraction (SAED) and energy-dispersive X-ray spectroscopy (EDS) to provide data at the bulk sample and individual crystal scale, respectively. Depending on the initial aqueous Fe-to-Cu ratios, we find that either nanoparticulate mackinawite (FeS), covellite (CuS), chalcopyrite (CuFeS_2), and/or nukundamite ($\text{Cu}_{5.5}\text{Fe}_{6.5}$) is/are formed. Covellite NPs show significant variation in Fe content and morphology (fine-scale spheroids vs. nanorods vs. nanoplates) depending on the synthesis conditions and even within a single reaction system. The presence of defects in covellite with certain morphology implicates formation by particle attachment followed by potentially rapid recrystallization. Chalcopyrite formation proceeds through the incorporation of Fe into precursor

covellite NPs, which adds to the poorly understood pathway of low-temperature chalcopyrite formation. The presence of bacteria modifies the crystal sizes, the phases formed [e.g., unique biogreigite (Fe_3S_4) formation from mackinawite], and the selectivity of mineral precipitation (e.g., more chalcopyrite) compared to abiotic systems. We hypothesize that these changes are induced by precipitation within local microenvironments surrounding the bacterial cells with different chemistry compared to the bulk solution. To our best knowledge, the reported work is the first systematic study on mixed Fe-Cu sulfide NPs precipitated in low-temperature aqueous solutions that are similar to environments on the majority of the Earth's (sub)surface.

BACKGROUND: FE-CU-SULFIDE MINERAL SYSTEM

Iron sulfides such as mackinawite (tetragonal FeS), greigite (cubic Fe_3S_4), and pyrite (cubic FeS_2) are common constituents of low-temperature anoxic environments on the Earth's (sub)surface (reviewed by Rickard and Luther 2007). Multiple experiments have shown that the earliest phase precipitated during the reaction of Fe(II) with sulfide in aqueous solution is nanoparticulate mackinawite (e.g., Schoonen and Barnes 1991; Harmandas and Koutsoukos 1996; Herbert et al. 1998; Benning et al. 1999; Wolthers et al. 2003; Michel et al. 2005; Ohfuji and Rickard 2006; Csákberényi-Malasics et al. 2012; Zhou et al. 2014; Picard et al. 2018). The nucleation of mackinawite is mediated by the assembly of sub-nanometer aqueous clusters in the form of Fe_2S_2 , followed by crystal growth via aggregation and attachment of the nuclei (Luther and Rickard 2005). Iron isotopic studies support the growth of mackinawite through aggregation (Guilbaud et al. 2010), although direct observation of FeS particle attachment has not been observed to our knowledge. In the presence of either trace oxidants (e.g., oxygen, elemental sulfur, oxidized species on mineral surfaces), trace Ni, bacteria, acidic pH, or high temperatures, mackinawite can be transformed into greigite and pyrite within days to weeks (Rickard 1975; Luther 1991; Harmandas et al. 1998; Benning et al. 2000; Rickard and Luther 2007; Gramp et al. 2010; Bourdoiseau et al. 2011; White et al. 2015; Morin et al. 2017; Wan et al. 2017; Picard et al. 2018). Mackinawite may also contain and adsorb significant amounts of trace metals such as Ni, Mn, Cu, Co, and Zn, and likely play a key role in trace metal sequestration in the environment (e.g., Morse and Arakaki 1993; Parkman et al. 1999; Veeramani et al. 2013; Zavašnik et al. 2014; Ikogou et al. 2017; Wilkin and Beak 2017).

Copper sulfides, in general, are trace components in environments on the Earth's surface, except in environments exposed to effluents from Cu-rich mining sites (Weber et al. 2009b; Fulda et al. 2013; Hofacker et al. 2013). It is difficult to identify the exact phase(s) of Cu-sulfides present in nature as the Cu-S mineral system contains various phases with variable Cu-to-S ratios, redox states and crystal structures (XRD patterns for each shown in Supplemental¹ Fig. S1). Experiments in low-temperature aqueous solutions, however, indicate that covellite [hexagonal Cu(I)S] is the most common product (Gramp et al. 2006; Sampaio et al. 2009; da Costa et al. 2013; Ikkert et al. 2013). Similar to mackinawite, nucleation of Cu-sulfides is thought to result from the assembly of aqueous clusters such as Cu_3S_3 and Cu_4S_6 (Luther et al. 2002; although see Ciglenečki et al. 2005 for a debate on the existence of Cu-sulfide clusters). Copper is typically

supplied as the Cu^{2+} ion, which must undergo reduction to Cu^+ prior to covellite formation. The reduction process occurs within minutes in the presence of sulfide (Pattrick et al. 1997; Luther et al. 2002). No TEM-based studies have yet been performed to determine the formation mechanism of covellite precipitated in low-temperature aqueous solutions. Experiments done in high-temperature organic solvents indicate that aggregation and attachment of the nuclei are likely the primary mechanisms for crystal growth of various copper sulfide nanocrystals (Xu et al. 2006; Du et al. 2007; Zhang et al. 2008; Li et al. 2011).

The mixed Fe-Cu-sulfide mineral system is much less studied compared to pure Fe- or Cu-sulfide mineral system, especially under low-temperature aqueous conditions. There are several known mixed Fe-Cu-sulfide minerals ranging from the Fe-rich end-member of cubanite (CuFe_2S_3) to the Cu-rich end-member of nukundamite ($\text{Cu}_{5.5}\text{FeS}_{6.5}$) (XRD patterns for each shown in Supplemental¹ Fig. S2). Among these minerals, chalcopyrite (CuFeS_2) is arguably the most common phase on the Earth's (sub)surface. The oxidation states of Cu and Fe in chalcopyrite have been a subject of intense debate, although recent studies indicated Cu exists in the monovalent state (Cu^+) and Fe in the trivalent state (Fe^{3+}) (Boekema et al. 2004; Goh et al. 2006; Pearce et al. 2006; Klekovkina et al. 2014; Conejeros et al. 2015). Since the 1960s, there are a few studies focusing on the formation mechanism for chalcopyrite in low-temperature aqueous solutions. Roberts (1961) formed chalcopyrite by attaching thin blocks of pyrrhotite (Fe_{1-x}S ; $x < 0.2$) and covellite together, prompting the suggestion that chalcopyrite was formed via solid state diffusion of Fe into covellite. Later on, Roberts (1963) formed chalcopyrite by mixing equimolar amount of aqueous Fe and Cu with excess sulfide. On the basis of the fast reaction rate on the order of minutes, he revised his earlier hypothesis and concluded that chalcopyrite formation must proceed through aqueous-mediated recrystallization of Fe- and/or Cu-sulfides. Cowper and Rickard (1989) repeated Roberts' earlier experiments and found contradictory results. In the presence of excess sulfide, they did not form chalcopyrite either by reacting (1) solid Fe-sulfides + solid Cu-sulfides, (2) aqueous Fe + aqueous Cu, or (3) aqueous Fe + solid Cu-sulfides. The only reaction capable of forming chalcopyrite was through the addition of aqueous Cu to solid Fe-sulfides. They, therefore, concluded that chalcopyrite could only form through reactions between aqueous Cu and a precursor Fe-sulfide mineral, consistent with the formation of Fe-sulfide phases prior to chalcopyrite formation in later studies (Parkman et al. 1999; Karnachuk et al. 2008; Ikkert et al. 2013).

MATERIALS AND METHODS

The experiments were performed at (1) different initial aqueous Fe-to-Cu ratios ($M_{\text{Fe/Cu}}$) and (2) in the presence and absence of sulfate-reducing bacteria to systematically study the factors controlling the formation of mixed-metal sulfide NPs. All experiments were performed in acid-cleaned serum vials to prevent trace metal carry-overs. Sample preparation and experimental procedures were conducted under anoxic conditions unless otherwise specified. Anoxic solutions were prepared by boiling for 20 min and N_2 degassing for another 20 min, followed by storage under pure N_2 or N_2/H_2 (97%/3%) headspace. A modified metal toxicity medium (MTM) was used as the precipitation matrix, which contains (per liter): 3.8 mL 80% lactic acid, 2.23 g Na_2SO_4 , 0.06 g $\text{CaCl}_2 \cdot 2\text{H}_2\text{O}$, 1.0 g NH_4Cl , 1.0 g $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 0.05 g yeast extract, 0.5 g tryptone, and 2.83 g HEPES. Initial pH was adjusted to 7.2 by the addition of NaOH. Metal stock solutions were prepared separately as 20 mM $\text{CuCl}_2 \cdot 2\text{H}_2\text{O}$ or 10 mM $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$ (the latter in pH 2 HCl solution matrix to prevent oxidation) and then later added into the medium. Five different systems

with variable $M_{\text{Fe/Cu}}$ were tested: 1:0 (that is, 1000 μM Fe with no added Cu), 2:1, 1:1, 1:5, and 0:1 (500 μM Cu with no added Fe). The mixed metal systems were prepared either by keeping the Cu concentration constant at 500 μM while varying the Fe concentrations, or by maintaining a constant total metal concentration of 500 μM . The resultant precipitates were collected for analyses after 4 to 30 days of incubation. Long-term changes were also monitored by collecting precipitates after up to 6 months of incubation.

Bacterial cell cultures and biological precipitation

Biological precipitation experiments were conducted by utilizing the sulfate reduction metabolism of *Desulfovibrio vulgaris* strain DSM 644 (also known as ATCC 29579 or strain Hildenborough) as a gradual sulfide source. The culture was maintained at either 4 or 30 °C over a period of weeks to months in a modified Postgate medium (DMSZ medium no.63; media composition detailed in Xu et al. 2016). Prior to each precipitation experiment, the culture was inoculated at 1% v/v into MTM and allowed to grow for 72 h at 30 °C while shaking. This step was crucial to remove metals that might be carried over from the Fe-rich Postgate medium. Sulfide concentrations were monitored through the methylene blue assay (Hach Co., Colorado) at this stage and reached a maximum of 15 mM after 3 days. The cells were then inoculated at 1% v/v into 100 mL of metal-amended MTM for the actual precipitation experiments. Within a day of incubation at 30 °C while shaking, black or dark brownish metal sulfide precipitates formed in the cultures. Each sample was separated into two aliquots for XRD (~80 mL) and TEM (~5 mL) analyses. Precipitates were first pelleted by centrifugation at 15000g for 5 min. Pellets for XRD were washed once with 10 mL anoxic H₂O prior to being resuspended in ~0.8 mL of anoxic H₂O and allowed to dry on a glass slide as a thin film. Pellets for TEM were treated with 10 mL of anoxic 0.1% Triton-X solution containing lysozyme and proteinase K (100 $\mu\text{g}/\text{mL}$ of each) for 1 h at 45 °C to lyse associated bacterial cells, which were then removed from the NP fraction by centrifugation at 15000g for 15 min. The NPs were washed once with 10 mL anoxic H₂O before being set aside for TEM analyses.

Abiotic precipitation

Abiogenic metal sulfide NPs were precipitated through titration of 50 mL of 15 mM Na₂S·9H₂O (in MTM matrix, pH 7.2) into 50 mL of metal-amended MTM at a rate of 2 mL/h inside the anaerobic chamber. The solutions were kept well-mixed using a polytetrafluoroethylene-coated magnetic stir bar. At the end of the titration, the stir bar was removed to prevent nucleation/growth effect due to added surface areas. Aggregates of NPs were separated by filtration through a 0.2 μm pore size mixed cellulose ester membrane; the solution remained clear during this step. The NP residues were then resuspended in 1 mL anoxic H₂O through repeated mixing by a pipettor. Approximately 0.8 mL of each sample was dried onto a glass slide for XRD analyses, while the rest was stored in H₂O until being prepared for TEM analyses.

Characterization

X-ray diffraction patterns of precipitates were collected using a Rigaku Mini-flex II equipped with a CuK α source. Individual scans were collected from 10 to 60° 2 θ angle with a step size of 0.05° and scan speed of 0.25 °/min, totaling a collection time of ~3.3 h per sample. Background correction was performed manually in Jade software using a scan collected from a blank glass slide as a reference. Average crystal size (L) along a plane was estimated using the Scherrer equation:

$$L = K\lambda/(\beta \cos \theta) \quad (1)$$

where K is the Scherrer shape constant (assumed to be 0.9; Langford and Wilson 1978), λ is the X-ray wavelength (0.15418 nm for CuK α), β is the full-width half-maximum (FWHM) of a peak corresponding to a particular plane and θ is the angle position of a peak. Intrinsic instrument broadening can contribute to error in β . This error was estimated by comparison to a commercial copper oxide nanoparticle (product no.544868 from Sigma Aldrich) with known XRD-derived crystal sizes of 18 nm (Triboulet et al. 2015). From this analysis, measurement errors contributed by β are expected to be <1 nm for crystal sizes under 10 nm.

Samples for TEM analyses were prepared in the anaerobic chamber by diluting in anoxic H₂O followed by sonication for up to one hour to disperse any aggregates. Between 2.5 and 10 μL of dispersed samples were dried onto Ni-coated grids (HC200-Ni, Electron Microscopy Sciences, Pennsylvania, U.S.A.) and shipped in an anoxic container (SampleSaver™ Storage Container, model SS-100/200, South Bay Technology Inc., California, U.S.A.) to the NanoEarth facility at Virginia Tech

(U.S.A.). Samples were imaged within two weeks of shipping using a JEOL-2100 TEM operating at 200 kV. There were no observable differences between samples that were dried immediately on the TEM grids and samples that were stored in anoxic H₂O for a few months prior to drying.

Elemental ratios (e.g., sulfur/metal) were measured with energy-dispersive X-ray spectroscopy (EDS) with a silicon drift detector and quantified using the standardless analysis method (Newbury and Ritchie 2014). EDS spot analysis was conducted with a beam size of 20–500 nm in diameter, while mapping analysis was conducted in scanning-TEM (STEM) mode with a beam size of 3–10 nm. Samples exhibiting oxygen peaks that were higher than the grid background (i.e., indicative of oxidation) were discarded from interpretation. The d -spacing values for NPs were measured on selected-area electron diffraction patterns (SAED) in DigitalMicrograph software. In high-resolution images where lattice fringes were clearly visible, d -spacing values were also measured directly by averaging several interplanar distances or in reciprocal space, based on the patterns derived after applying fast Fourier transform (FFT) in Digital Micrograph. These methods gave d -spacing values that were consistent within ± 0.05 Å.

RESULTS AND DISCUSSION

The metal sulfide NPs precipitated in this study display significant variations in their phase, morphology, and elemental compositions, primarily as a function of the initial aqueous Fe-to-Cu ratios ($M_{\text{Fe/Cu}}$). In the following sections, we first describe the physicochemical attributes of these NPs under each of the tested conditions. The formation mechanism(s) associated with these NPs in single-metal systems, and how such mechanisms may be modified in mixed-metal systems, are discussed. Additionally, the subtle differences in biogenic and abiogenic NPs are detailed, and the specific biological mechanisms causing these differences are proposed. Finally, the physicochemical attributes of the synthetic NPs in this study are compared to those of naturally occurring NPs to ascertain if experimental approaches can be utilized to gain insight into NP-mediated reactions in the natural biogeochemical cycling of metals and sulfur.

Initial Fe-to-Cu ratios as the primary control on the physicochemical attributes of biogenic and abiogenic nanoparticles

At 1:0 $M_{\text{Fe/Cu}}$ (Fe with no added Cu), mackinawite is the only phase detectable by XRD within 30 days of incubation. Both biogenic and abiogenic precipitates display a prominent XRD peak at 2 θ angle of ~17° corresponding to the (001) plane of mackinawite (Fig. 1a). Estimated crystal sizes based on the (001) XRD peak indicate slightly larger biogenic mackinawite (7 to 9 nm) than abiogenic mackinawite (4 to 6 nm) (Table 1). All other peaks are either small or broad, consistent with the nanoparticulate nature of the precipitates. Under TEM, both abiogenic and biogenic mackinawite are observed to adopt wrinkled nanosheet morphology ~300 to 1000 nm in size (Fig. 2a). This morphology is similar to mackinawite aggregates precipitated in other experimental studies (Ohfujii and Rickard

TABLE 1. XRD-derived parameters for the (001) plane of mackinawite

System	d -spacing (Å)		Size (nm)	
	Biogenic	Abiogenic	Biogenic	Abiogenic
1:0 Fe:Cu	4.91 to 5.09	4.94 to 5.02	7 to 9	4 to 6
1:0 Fe:Cu ^a	5.04 to 5.07	4.98 to 5.24	9 to 26	3 to 9
2:1 Fe:Cu	4.72 to 4.73	5.10 to 5.17	≤3	≤3
1:1 Fe:Cu	–	5.25 to 5.32	–	≤3

Note: Samples from this study were measured at least in duplicates.

^a Fe only experiments from Picard et al. (2018). Biogenic mackinawite data from experiments containing microbes (live or dead) and abiogenic mackinawite data from experiments performed in the absence of microbes.

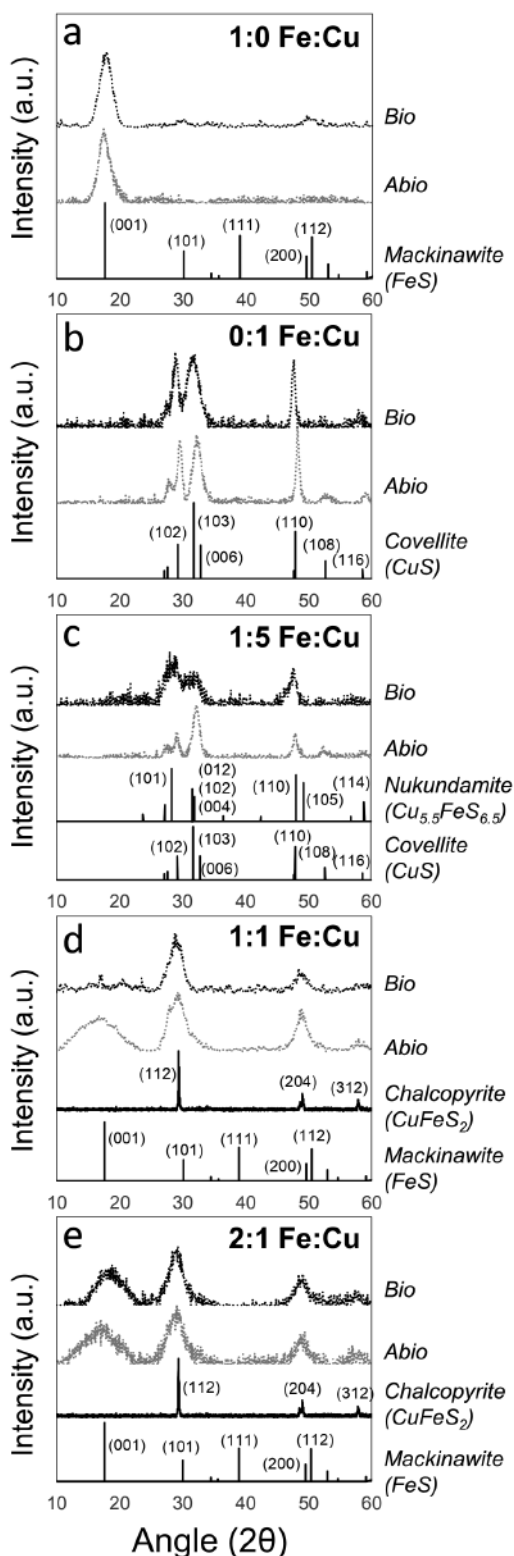


FIGURE 1. XRD patterns of biogenic (Bio) and abiogenic (Abio) precipitates, with different panels corresponding to the initial aqueous Fe-to-Cu ratios. Reference patterns of relevant phases are shown at the bottom of each panel.

TABLE 2. Compiled d -spacing values (in Å) and corresponding planes for selected minerals based on the American Mineralogist Crystal Structure Database (<http://rruff.geo.arizona.edu/AMS/amcsd.php>; accessed date January 9, 2019; Downs and Hall-Wallace 2003)

	d -spacing (hkl)				
	Mackinawite	Greigite	Covellite	Chalcopyrite	Nukundamite
	5.03 (001)	5.70 (111)	8.18 (002)	3.04 (112)	11.20 (001)
	2.97 (101)	3.49 (220)	4.09 (004)	2.64 (200)	5.60 (002)
	2.56 (110)	2.98 (311)	3.29 (100)	2.61 (004)	3.73 (003)
	2.52 (002)	2.47 (400)	3.22 (101)	1.87 (220)	3.28 (100)
	2.31 (111)	2.27 (331)	3.05 (102)	1.86 (204)	3.14 (101)
	1.84 (200)	2.02 (422)	2.82 (103)	1.59 (312)	2.83 (012)
	1.81 (112)	1.90 (511)	2.73 (006)	1.58 (116)	2.83 (102)
	1.73 (201)	1.90 (333)	2.32 (105)		2.80 (004)
	1.68 (003)	1.75 (440)	2.10 (106)		2.46 (013)
	1.56 (211)	1.56 (620)	2.05 (008)		2.24 (005)
			1.90 (107)		2.13 (104)
			1.898 (110)		1.89 (110)
			1.74 (108)		1.85 (015)
			1.65 (200)		1.85 (105)
			1.64 (201)		1.79 (112)
			1.62 (202)		1.68 (113)
			1.59 (109)		1.64 (200)
			1.57 (203)		1.62 (016)
			1.56 (116)		1.62 (106)
					1.62 (021)
					1.60 (007)
					1.57 (022)
					1.57 (202)
					1.567 (114)

Note: Bolded values indicate major diffraction peaks, defined as having a relative intensity >0.2 compared to the most intense peak when analyzed by XRD.

2006; Csákberényi-Malasics et al. 2012; Veeramani et al. 2013; Picard et al. 2018). Close observations of the nanosheets reveal electron dense, fibrous-like structures with d -spacing of 5.04 Å (Figs. 2b and 2c), corresponding to the (001) planes of mackinawite (Lennie 1995; Table 2). These areas are interpreted to reflect regions where the nanosheet can be viewed edge-on, in cross section, revealing the (001) plane. The SAED patterns derived from the nanosheets show diffuse rings with d -spacing values that corresponded to the (200), (111), (002), and (101) planes of mackinawite (Fig. 2d). The (001) diffraction ring, if present, is difficult to see due to its proximity to the high intensity (000) center spot. After six months of incubation, mackinawite remains as the sole material detectable by XRD with no significant change in the crystal sizes. Interestingly, TEM observations reveal minor amounts of 30–50 nm nanoplates in the biological systems that are absent in abiotic systems. Analysis of the SAED patterns derived from these nanoplates show that they are single crystals of greigite (Figs. 3a and 3b) or greigite aggregated with mackinawite (Figs. 3c and 3d).

At 0:1 $M_{Fe:Cu}$ (Cu with no added Fe), covellite is the only phase detectable by XRD for both biogenic and abiogenic precipitates, with higher crystallinity associated with the latter (Fig. 1b). Three morphologies of NPs are observed for both abiogenic and biogenic samples: fine nanocrystals <10 nm in size (Fig. 2e), ~35 nm long nanorods (Fig. 2f) and ~40 nm wide nanoplates with poorly defined edges (Figs. 2f and 2g). The SAED patterns collected from aggregates of the fine nanocrystals and nanoplates show diffuse rings as well as distinct diffraction spots with overlapping d -spacing values that are consistent with covellite (Fig. 2h; Table 2). Similarly, elemental S-to-Cu ratios (via EDS) of the nanorods and nanoplates indicate a ratio of 1.0 ± 0.1 ($n = 2$ each), consistent with covellite. However, the fine

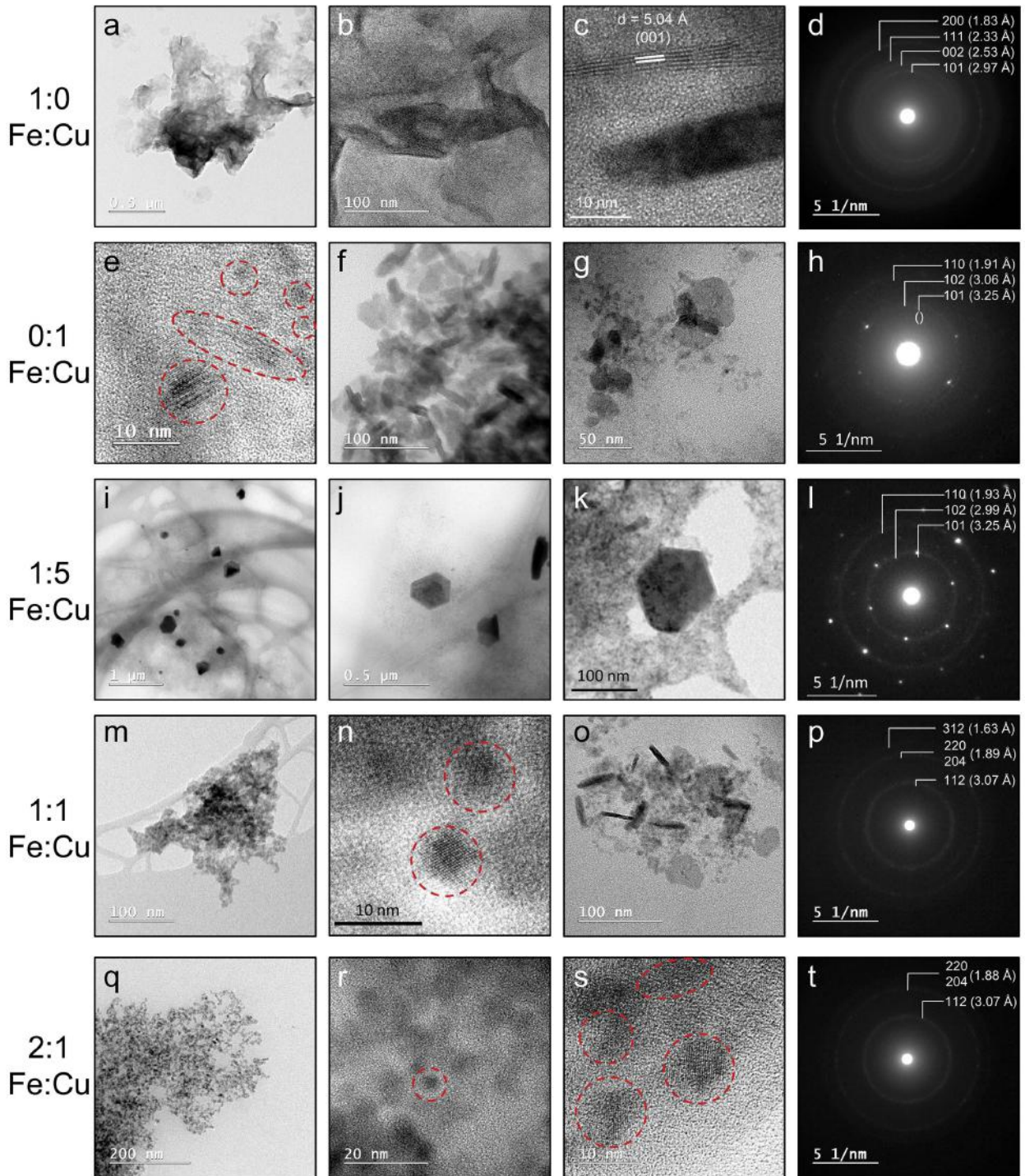


FIGURE 2. TEM images of nanoparticles precipitated at different initial aqueous Fe-to-Cu ratios as arranged by row. The final column shows representative wide-view (>50 nm spot size) SAED patterns for the samples, with d -spacings that are consistent with (from top to bottom row) mackinawite, covellite, covellite, chalcopyrite, and chalcopyrite. (a–d) Wrinkled nanosheet composed of mackinawite aggregates, with electron-dense regions exhibiting fibrous-like structures. (e–h) From left to right: <10 nm fine covellite nanocrystals, nanorods, and nanoplates. The SAED pattern is collected from aggregates containing all three morphologies. (i–l) Images focusing on Fe-rich euhedral covellite nanoparticles. The SAED pattern is collected from a single euhedral crystal (distinct diffraction spots) that is surrounded by fine covellite nanocrystals (diffraction rings). (m–p) Chalcopyrite aggregates composed primarily of <10 nm fine nanocrystals, with corresponding diffraction rings in the SAED pattern. In the same sample, minor amounts of nanorods and nanoplates [with variable Fe/(Fe+Cu) ratios] are also visible (panel o). (q–t) Chalcopyrite aggregates composed primarily of <10 nm fine nanocrystals, with corresponding diffraction rings in the SAED pattern. (Color online.)

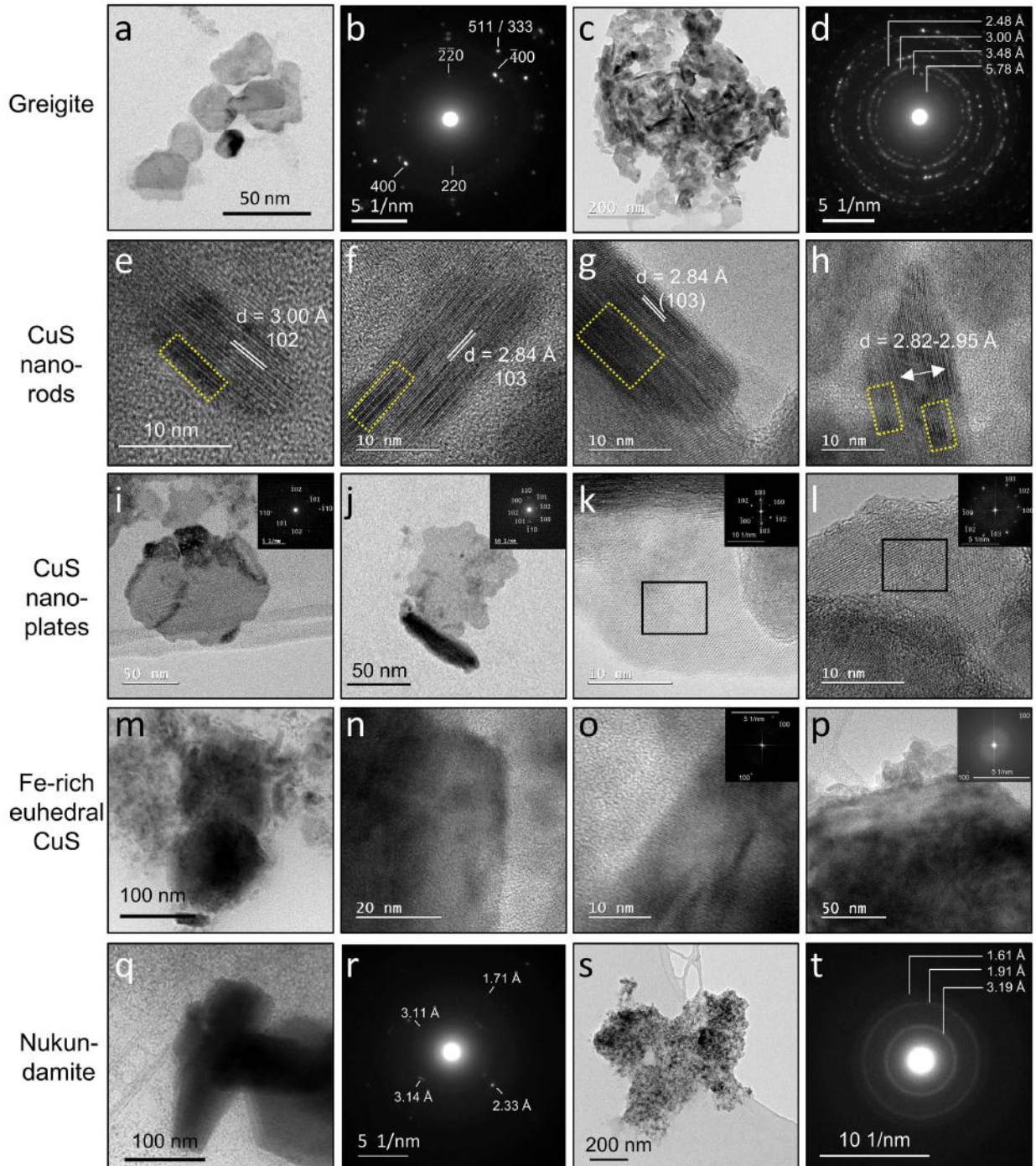


FIGURE 3. TEM images focusing on the different nanoparticle morphology. (a–d) Greigite nanoplates with corresponding SAED pattern derived from multiple greigite crystals. (e–d) Greigite-mackinawite aggregates with corresponding SAED patterns. The 5.78 Å *d*-spacing is characteristic of greigite (111) reflection, while the rest of the diffraction rings can be attributed to either greigite or mackinawite (e–h) Covellite nanorods with predominantly (102) or (103) plane perpendicular to the direction of elongation. Defects within the crystal structures (electron dense regions or kinks) are highlighted in yellow boxes. (i–l) Covellite nanoplates with poorly defined edges. Insets show either the whole-area SAED (i and j) or FFT-derived diffraction patterns from partial areas defined by the black square outline (k and l). Note that the diffraction patterns may reflect multiple overlapping crystals. (m–p) Euhedral nanoparticles of Fe-rich covellite. These NPs are larger and thicker compared to the thin nanoplates in i–l. (q–t) Large tabular NPs and nanoparticle aggregates in mixed metal Fe-Cu systems ($M_{\text{Fe/Cu}} = 1:5$ and 1:1). The SAED patterns display *d*-spacing values of ~3.15 Å, consistent with the (101) reflection of nukundamite ($\text{Cu}_{3.5}\text{Fe}_{6.5}$). Other *d*-spacing values are ambiguous and can be assigned either to covellite, chalcopyrite, or nukundamite. (Color online.)

nanocrystals occasionally display excess S relative to Cu (up to S/Cu = 1.5), which could be due to sulfur adsorption to the NPs' surface or trace quantities of villamaninite (CuS₂).

High-resolution TEM images of both biogenic and abiogenic covellite nanorods reveal lattice fringes perpendicular to the elongation direction, with *d*-spacing values that correspond to either the (102) or (103) plane (Figs. 3e–3g). All of the imaged nanorods display electron dense regions that we interpret to indicate randomly located defects within the crystal structures. Depending on the orientation of the nanorods with respect to the electron beam, we can observe that some nanorods display clear defects in the form of atomic dislocations (Fig. 3h). For these defect-containing nanorods, the measured *d*-spacing (perpendicular to the elongation direction) vary from 2.82–2.95 Å; this value range falls in between the (102) and (103) plane (optimal *d*-spacing of 3.05 and 2.82 Å, respectively; Table 2). Meanwhile, covellite nanoplates in the same sample are observed to be single crystals with no visible defects within the crystal structure; *d*-spacing analyses indicate a prominence of the (110), (102), (103), and (100) planes depending on the viewing orientation (Figs. 3i–3l).

In mixed-metal systems, the physicochemical attributes of the precipitates are dependent on the $M_{\text{Fe/Cu}}$ but are not influenced by the total metal concentrations within the studied range. At 1:5 $M_{\text{Fe/Cu}}$, the XRD data for the abiogenic precipitates indicate covellite as the sole precipitate. In contrast, the XRD data for the biogenic precipitates indicate the presence of covellite along with another phase that resulted in a more intense peak at 28° 2θ angle (Fig. 1c). TEM analyses for both abiogenic and biogenic samples indicate the presence of NPs with morphological similarity to fine covellite nanocrystals, nanorods, and nanoplates, along with the additional occurrence of larger (100–200 nm) and thick euhedral crystals with well-developed edges (Figs. 2i–2l). Based on EDS, the large euhedral crystals contain Fe incorporated within its structure (Fig. 4a) and display average Fe/(Fe+Cu) ratio of 0.11 ± 0.05 and S/(Fe+Cu) ratio of 0.92 ± 0.09 (*n* = 8). The elemental ratios are consistent with either Fe-rich covellite [Fe/(Fe+Cu) ratio of 0.01–0.18; Clark 1971; Maydagán et al. 2013; Melekestseva et al. 2017] or nukundamite [hexagonal Cu_{5.5}FeS_{6.5}, Fe/(Fe+Cu) = 0.15; Rice et al. 1979]. Comparison of SAED patterns collected from the biogenic and abiogenic euhedral crystals indicates one major difference between the two. While abiogenic crystals display diffraction patterns with *d*-spacing values that are consistent with covellite (Fig. 2l), the biogenic crystals display an additional diffraction spot/ring corresponding to a *d*-spacing value around 3.15 Å (Figs. 3q–3t); this *d*-spacing corresponds to the (101) plane of nukundamite with a corresponding 28° 2θ angle in XRD spectrum (Sugaki et al. 1981). Thus, abiogenic precipitates are composed of Fe-rich covellite while biogenic precipitates are composed of mixtures of Fe-rich covellite and nukundamite.

At 1:1 $M_{\text{Fe/Cu}}$, XRD data indicate the formation of chalcopyrite (CuFeS₂) in both biological and abiotic systems. In addition, smaller XRD peaks indicative of mackinawite and possibly nukundamite are also present in the abiotic system (Fig. 1d). High-resolution TEM analysis of both biogenic and abiogenic precipitates show aggregates of 3 to 12 nm crystals with SAED-derived *d*-spacing values of 3.07 and 1.89 Å (Figs. 2m, 2n, and 2p), which can be attributed to either chalcopyrite's (112) and (220)/(204) planes or to covellite's (102) and (103) planes (Table 2).

Elemental mapping of the nanocrystal aggregates indicates overlapping distributions of Fe, Cu, and S with approximate ratios of 1:1:2 (Fig. 4b). Based on the XRD, EDS, and SAED data, we, therefore, identify the nanocrystal aggregates as chalcopyrite. Close observations also reveal the presence of nanorods (in both the biological and abiotic systems) and nanoplates (in biological system only) that are morphologically similar to covellite precipitated in the absence of Fe (Figs. 2o and 5). Based on EDS, the nanorods and nanoplates display a wide range of Fe/(Fe+Cu) ratios (0.04 to 0.43; average 0.26 ± 0.15) and S/(Fe+Cu) ratios (range of 0.74 to 1.11; average = 1.00 ± 0.11). The SAED patterns derived specifically from the nanoplates and nanorods yield *d*-spacing values that are consistent with chalcopyrite, nukundamite, and covellite; these crystals likely represent various Fe-Cu-S minerals with variable crystal chemistry and elemental compositions similar to chalcopyrite at one end and covellite at the other.

At 2:1 $M_{\text{Fe/Cu}}$, chalcopyrite is the main precipitate detected by XRD along with lesser amounts of mackinawite in both biological and abiotic systems (Fig. 1e). The chalcopyrite nanocrystals are similar in morphology to the nanocrystal aggregates observed at 1:1 $M_{\text{Fe/Cu}}$ (Figs. 2q–2s). No nanorods or nanoplates are observable. Collected SAED patterns from the nanocrystal aggregates display diffuse rings with *d*-spacing values of 3.07 and 1.88 Å corresponding to the (112) and (204) planes of chalcopyrite (Fig. 2t). Although the presence of mackinawite is indicated by XRD data, no images of mackinawite are captured by TEM.

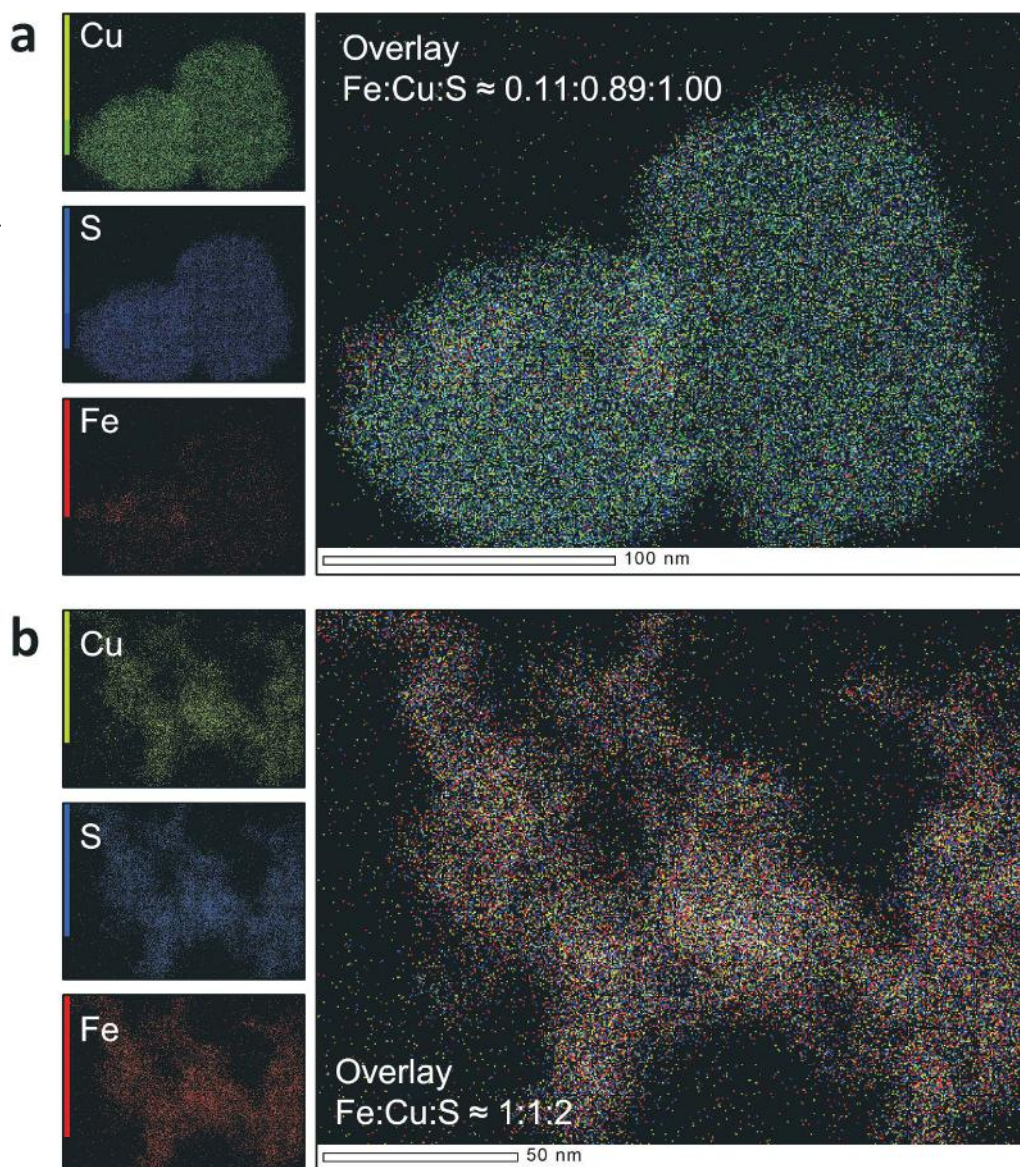
Aged precipitates from the pure Cu or mixed Fe-Cu systems are examined up to four months following the initial precipitation to determine if the NPs evolve significantly over time. Aging does not cause variation in the bulk XRD data, indicating that most nucleation and growth occurred within the first few days and stopped after the aqueous metal supply is exhausted due to excess sulfide production or addition. Under TEM, small changes in crystal sizes over time are discernible. Specifically, biogenic covellite nanoplates precipitated at 0:1 $M_{\text{Fe/Cu}}$ decrease slightly in size over a period of 50 days, while the size of the nanorods remains unchanged (Supplemental¹ Fig. S3). These changes are not studied systematically for all samples and are beyond the scope of the current study.

Formation mechanism(s) of Fe-Cu-sulfide nanoparticles

The final size, morphology, and elemental composition of the metal sulfide NPs, be it mackinawite, covellite, or chalcopyrite, vary significantly depending on the initial aqueous Fe:Cu ratios, both in the absence and presence of *D. vulgaris*. These variations in physicochemical attributes likely stem from fundamental changes in the formation mechanism(s) of the respective NPs caused by precipitation in the presence of other metal ions, as discussed below.

Mackinawite: Size modification in the presence of Cu. In our study, we find that smaller mackinawite NPs are formed in both biogenic and abiogenic systems when precipitation occurred in the presence of Cu (Table 1; compare $M_{\text{Fe/Cu}}$ = 2:1 and 1:1 vs. $M_{\text{Fe/Cu}}$ = 1:0). This result is in contrast to the study by Zavašnik et al. (2014) who found larger mackinawite NPs when precipitation occurred in the presence of Cu. Density functional theory simulations by Kwon et al. (2015) support the results of Zavašnik et al. (2014) and further attribute the change to the substitution of Cu for Fe in mackinawite's crystal structure, leading to an increase in

FIGURE 4. EDS maps of (a) euhedral Fe-rich covellite crystals precipitated at 1:5 Fe:Cu ratio, displaying fairly homogenous Fe distribution within their structures and (b) aggregates of chalcopyrite precipitated at 1:1 Fe:Cu ratio. (Color online.)



d-spacing along the (001) plane. Thus, we also measure the (001) *d*-spacing based on our XRD data and find that while abiogenic mackinawite display increased (001) *d*-spacing (consistent with previous results), biogenic mackinawite actually displays smaller (001) *d*-spacing in the presence of Cu. This implies that differences in the experimental setup and/or biological effects can possibly modify the exchange mechanism between Fe and Cu ions in mackinawite as well as the incorporation of Cu into the space between the (001) planes, leading to differences in the particle size. Such changes may have implications to mackinawite's reactivity and stability (Kwon et al. 2015), and further investigation is necessary to illuminate the specific mechanism.

Covellite: The role of particle attachment and Fe in promoting growth. In this study, covellite show remarkable variation in terms of morphology, from fine nanocrystals, nanorods, and nanoplates, to larger and thicker Fe-rich crystals of up to 200 nm in size. Because of covellite's exceedingly low solubility (Shea

and Helz 1989) and rapid nucleation kinetics (Morse and Luther 1999), crystal growth via particle attachment was proposed to be its dominant growth mechanism (Luther and Rickard 2005; Rickard and Luther 2006). This is in part supported by previous TEM analyses showing oriented attachment of covellite nanorods (Xu et al. 2006) and nanoplates (Du et al. 2007) and other Cu-sulfide phases (Li et al. 2011). In the current study, direct observations of oriented attachments for covellite nanostructures are however uncommon. One example of such observation is seen for precipitates at day 4, whereby ~3–5 nm nanocrystals are oriented to form a larger pseudo-nanorod structure (Fig. 6). Aside from this example, TEM analyses also indicate aggregation of nanorods and stacking of smaller nanoplates and nanocrystals on top of larger nanoplates (Supplemental¹ Fig. S4), although it is unclear if these are the natural state of the NPs or an artifact of sample preparation. The absence of strong evidence for oriented attachment may implicate either classical ion-by-ion growth or fast recrystalliza-

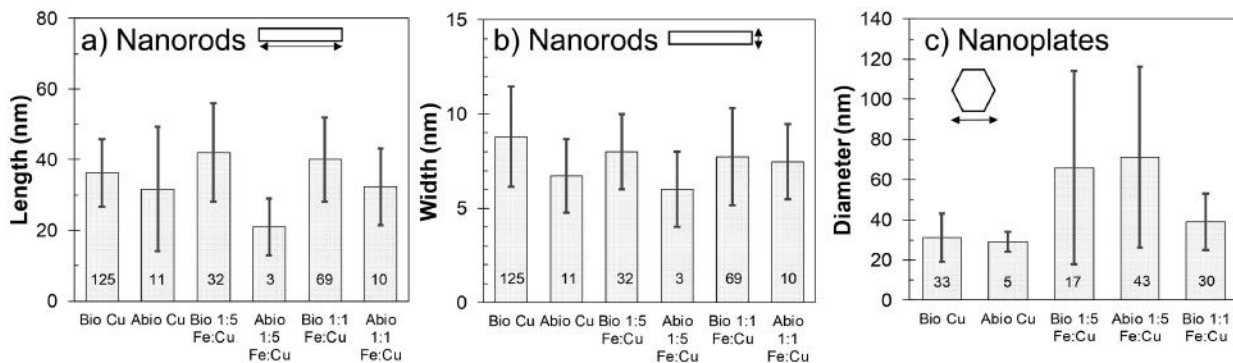


FIGURE 5. Size dimensions for (a–b) nanorods and (c) nanoplates. Numbers in the bar charts indicate the number of crystals that were measured from TEM images.

tion of covellite NPs following initial particle attachment. For the latter mechanism, we note that previous studies performed TEM analysis within 24 h of initial covellite precipitation (Xu et al. 2006; Du et al. 2007; Li et al. 2011), compared to the longer (i.e., ≥ 4 days) duration in the current study. It is possible for covellite recrystallization to occur within this period based on its fast nucleation kinetics, leading to the erasure of any evidence for oriented attachment prior to TEM analysis of the samples.

Indirect evidence for particle attachment formation pathway of covellite can be seen in the forms of defects within the structures of NPs. At 0:1 and 1:5 $M_{\text{Fe/Cu}}$, stacking faults and/or twinning are observable in the fine covellite nanocrystals and nanorods. Defects within the fine nanocrystals developed along either the (100)/(101) (measured d -spacing cannot conclusively differentiate between the two) or (102) plane (Figs. 7a–7c) while defects within the nanorods developed along the (102) or (103) plane (Figs. 3e–3h and 7d–7e). It is particularly noted that these structural defects are associated with most of the observed nanorods. The occurrence of the (102) planes in the fine nanocrystals and nanorods is unexpected given the relatively high energies of these surfaces (Morales-García et al. 2017). We predict that the surface energies of the (101) and (103) planes should be similarly high in energy, although this remains to be confirmed. Based on these, we establish a positive relationship between the occurrence of structural defects and the exposure of high-energy planes of covellite. Two possible mechanisms might have led to the occurrence of structural defects. The first mechanism is through crystallographic attachment of sub-nanoparticles along (i.e., to hide) the high-energy planes. In this case, twinning and stacking faults could have been formed due to a relatively high probability of imperfect lattice matching of two (101)/(102)/(103) surfaces coming together. The second mechanism relates to the release of the relatively high energy associated with the (101)/(102)/(103) surface through formation of structural defects (as has been explained in the case of ZnS in Xu et al. 2016). Structural defects are however absent in the hexagonal nanoplates (including in the Fe-rich euhedral plates), consistent with this morphology being the predicted equilibrium morphology of covellite (Morales-García et al. 2017). The absence of defects in these nanoplates most likely indicates rapid recrystallization to a thermodynamically stable morphology. Overall, TEM data seem to confirm the growth via attachment and re-assembly of

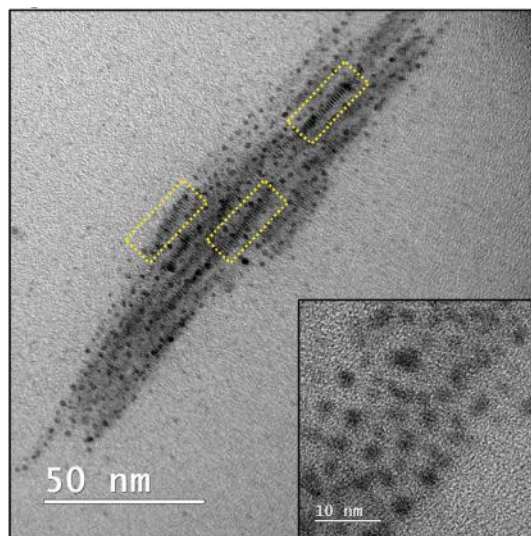


FIGURE 6. A pseudo-nanorod structure that is composed of oriented arrangement of smaller covellite nanocrystals (inset). Within the structure, smaller areas that show more coherent arrangements are highlighted in yellow. (Color online.)

sub-nanoparticles as the dominant mechanism for the formation of defects in the fine nanocrystals and nanorods.

The addition of Fe leads to the formation of euhedral covellite crystals up to 200 nm in size. To our knowledge, this is the first recognition of Fe promoting covellite's crystal growth. One explanation is that the incorporation of Fe into covellite's crystal structure results in a higher thermodynamic stability for the mineral, thus allowing growth to larger sizes at equilibrium. Another possibility, assuming growth by particle attachment, is that Fe adsorption to the covellite surface weakens the repulsion between covellite NPs and allows more attachment events to occur. Regardless of the specific mechanism, our study conclusively demonstrated that the addition of a foreign metal ion could significantly modify the size and morphology of metal sulfide NPs, which likely affects the reactivity of these NPs.

Chalcopyrite and nukundamite: Formation via Fe incorporation into precursor covellite. The formation of chalcopyrite at low temperature was previously claimed to proceed predomi-

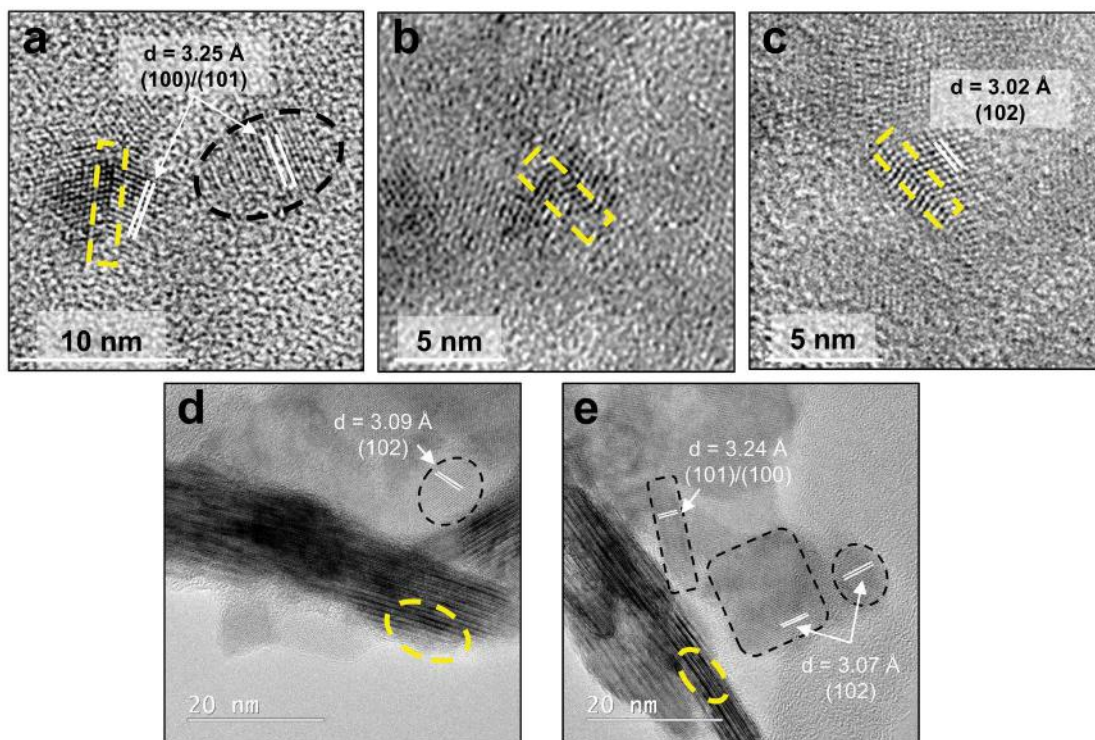


FIGURE 7. High-resolution images of fine covellite nanocrystals and nanorods, focusing on defects within the crystal structure as outlined in yellow. Boundaries of the fine nanocrystals are outlined in black to assist visualization. (a) Twinning and (b–c) stacking faults can be observed within <10 nm nanocrystals across the (100)/(101) or (102) planes. (d–e) Stacking faults and kinks can be observed within nanorods. The morphological variety of fine covellite nanocrystals with different planes exposed are also evident. (Color online.)

nantly, if not exclusively, through the incorporation of Cu into precursor Fe-sulfides (Cowper and Rickard 1989). However, in this study, we are able to form chalcopyrite by adding excess sulfide to aqueous mixtures of Fe and Cu (at 2:1 and 1:1 $M_{\text{Fe/Cu}}$), consistent with the earlier work of Roberts (1963). Cowper and Rickard (1989) performed a similar experiment and found that the “immediate” precipitates were poorly crystalline mackinawite, yarrowite ($\text{Cu}_{1.12}\text{S}$), spionkopite ($\text{Cu}_{1.39}\text{S}$) and additional XRD-invisible phases with variable Fe/Cu ratios. Aged precipitates were not analyzed in that study. We analyze the early phase abiogenic precipitates (~3 h old) produced at 1:1 $M_{\text{Fe/Cu}}$ in our study and find no XRD-detectable phases. Thus, it seems that abiogenic chalcopyrite formation requires >3 h of aging, which may explain why it was not detected in the earlier work of Cowper and Rickard (1989). In our study, this same experimental design yields mixed Fe-Cu-containing nanocrystals, nanorods, and nanoplates that are morphologically similar to covellite after four days of aging (Fig. 2o), suggesting that chalcopyrite (and nukundamite) is formed through the incorporation of Fe into precursor covellite. This is consistent with the notion that covellite should form faster than Fe-sulfides (Morse and Luther 1999). Additionally, chalcopyrite formation had been shown to proceed through the incorporation of aqueous Fe into Cu-sulfides at high temperatures (>100 °C), with the morphology of the precursor Cu-sulfides being preserved during the reaction (Liu et al. 2018). To confirm whether this same reaction pathway can proceed at low temperatures, we add 500 μM of aqueous Fe to a 500 μM covellite suspension in MTM (previously aged for

six days) and allow the reaction to proceed for four days at room temperature in the presence of excess sulfide (~7 mM). The final precipitates are identified by XRD as chalcopyrite, nukundamite, and mackinawite with no traces of covellite remaining. This result confirms that chalcopyrite can form through the incorporation of Fe into precursor covellite. The discrepancy with the earlier work of Cowper and Rickard (1989) most likely stems from the use of polished sections of highly crystalline Cu-sulfides compared to the use of covellite NPs in the current study, which highlights the stark difference in reactivity for nanoscale vs. larger-scale minerals.

Bacterial presence influences the crystal sizes and phase selectivity of Fe-Cu-sulfide NPs

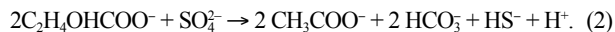
The presence and metabolic activity of sulfate-reducing bacteria (SRB) can influence Fe-Cu-sulfide NPs precipitation in our study in several ways. The cell walls of bacteria and secreted extracellular polymeric compounds (EPS) contain anionic carboxyl and phosphoryl groups, which allows for the adsorption of positively charged Fe and Cu ions (Beveridge and Murray 1976; Beveridge and Koval 1981; Ferris et al. 1989; Chen et al. 2011; Hao et al. 2013). As SRB produce sulfide, metal sulfide NPs are preferentially precipitated on the cell walls and EPS, as previously documented for Fe-sulfides (e.g., Fortin et al. 1994; Herbert et al. 1998; Donald and Southam 1999; Picard et al. 2018) and Cu-sulfides (Jalali and Baldwin 2000; Kiran et al. 2015). Once these NPs are formed, interactions with organic molecule assemblage secreted by SRB can further modify the

NPs' properties. For example, organic coatings can bridge the interaction between two or more NPs and encourage growth via aggregation, or conversely inhibit growth by coating the NPs (Ikuma et al. 2015). The properties of the biogenic NPs are therefore potentially influenced by the complex interaction between various organic molecules with the metal ions and the nanoparticle surfaces.

Mineral-specific response to biologically mediated crystal size modification. In our study, we observe a significant difference in the crystal size/crystallinity (based on XRD) of the biogenic NPs compared to abiogenic NPs. Specifically, biogenic mackinawite exhibits larger crystal sizes (Table 1; Fig. 1a) while biogenic covellite exhibits lower crystallinity compared to their abiogenic counterparts (Fig. 1b). No significant size differences are found between biogenic and abiogenic chalcopyrite, although its formation in a complex mineral mixture makes comparison difficult. Overall, the results are consistent with previous studies that compare between biogenic and abiogenic mackinawite (Gramp et al. 2010; Picard et al. 2018) and covellite (Gramp et al. 2006). Differences in precipitation rates were considered as a potential mechanism to explain these size differences, although studies on FeS and ZnS found no correlation between precipitation rates and crystal sizes (Xu et al. 2016; Picard et al. 2018). For biogenic mackinawite, its size increase was attributed to the presence of cell walls and EPS that acted as nucleation and growth sites for the mineral (Picard et al. 2018). The contrasting size response of biogenic covellite compared to biogenic mackinawite indicate a complex interaction between bacteria and NPs, with size modification effects to the NPs that are mineral specific.

Greigite and chalcopyrite formation favored via precipitation on the cell wall and EPS. In the presence of SRB, mackinawite is observed to transform to greigite after six months of incubation in the absence of Cu. This transformation process requires the oxidation of mackinawite to Fe(III)-containing mackinawite, which acts as an intermediate for greigite formation (Bourdoiseau et al. 2011; White et al. 2015). Trace amounts of oxygen introduced during sample handling may accelerate this reaction, resulting in artifactual greigite formation that is not representative of experimental conditions (Rickard and Luther 2007; Csákberényi-Malasics et al. 2012). We note that oxygen contamination is unlikely in this study given that all samples are handled anoxically as much as possible. Furthermore, greigite is only observed in the six months old biogenic sample and not for any other samples. Disregarding oxygen as a potential oxidant, H^+ is the most likely oxidant for the transformation reaction, as originally proposed by Rickard and Luther (2007). This idea is consistent with the observations that greigite formation from mackinawite proceeded slowly at near-neutral pH but was significantly accelerated at acidic pH (Bourdoiseau et al. 2011). With this in mind, we can propose that the role of SRB is to provide low-pH (sub)microenvironments that help to catalyze greigite formation. It is well established in various microbial biofilms that low-pH microenvironments around cells can be generated due to secretion of inorganic and organic acids, even in well-mixed reactors (De Los Ríos et al. 2003; Hunter and Beveridge 2005; Hidalgo et al. 2009). Specific to *D. vulgaris*, sulfate reduction

is coupled to lactate breakdown to acetate (Pankhania et al. 1986; Voordouw 2002; Heidelberg et al. 2004), which generates acidity (H^+) around the cells:



This hypothesis can be tested by comparative analysis of NPs associated with the cell surfaces vs. those located further away, possibly via: (1) sample embedding coupled to cryo-electron microscopy to preserve spatial information as much as possible or (2) pre-separation of the different NP fractions based on size and/or density. In both cases, the possibility of NPs being detached from the cell surfaces during sample preparation must be taken into account. While challenging, research into this area may provide new insights into unrecognized pathways for Fe redox cycling in anoxic environments.

In mixed-metal systems, the presence of SRB selectively favors the precipitation of nukundamite and chalcopyrite at lower $M_{Fe/Cu}$ compared to abiotic systems. Specifically, at 1:5 $M_{Fe/Cu}$, the biological system produces nukundamite at sufficiently high abundance for XRD detection while the abiotic system only produces Fe-rich covellite. Additionally, at 1:1 $M_{Fe/Cu}$, biogenic chalcopyrite is detected as the dominant phase by XRD, while the abiotic system produces a mixture of chalcopyrite, mackinawite, and nukundamite. These observations indicate that the presence of SRB accelerates the rearrangements of atoms to form mixed Fe-Cu-sulfide NPs with crystal structures that are distinct from the precursor covellite (with some Fe substituting for Cu within the crystal structure). The overall reaction for covellite [Cu(I) sulfide] transformation to chalcopyrite [Cu(I)-Fe(III)-sulfide] is given by:



As written, the sulfur atom in covellite and aqueous Fe participates in a redox reaction, with the former acting as an oxidant. The sulfur atom associated with bisulfide provides the additional sulfur to form chalcopyrite without participating in redox reactions. The overall reaction indicates that chalcopyrite formation is preferred in areas where high amounts of covellite, Fe, and sulfide are available, as typically found around bacterial cells. The transformation reaction produces H^+ as a by-product that can potentially be harvested by SRB to supplement proton gradient maintenance across cell membrane for ATP generation, similar to as proposed for the function of intracellular calcite precipitation in *Achromatium* (Mansor et al. 2015). Overall, we postulate that the presence and metabolic activities of SRB play a major role in accelerating the formation of chalcopyrite and nukundamite from covellite.

ENVIRONMENTAL IMPLICATIONS

Because of the technical challenges in isolating and characterizing naturally occurring NPs, there is a huge interest in using synthetic NPs to gain insight into the behavior of natural NPs, which may prove fruitful as long as these two types of NPs share the same physicochemical attributes. We, therefore, perform a literature search of electron microscopy-based studies on metal sulfide NPs found in low-temperature environments for comparison with our synthetic NPs. Notably, ~20 nm mackinawite

nanorods from the riverbeds of the Clarence River (Australia) (Burton et al. 2009), 2–35 nm spherical covellite, anilite (Cu_{1.75}S) and chalcocite (Cu₂S) in organic-rich soils contaminated by mining activities in Ontario (Canada) (Mantha et al. 2019), ~100 nm aggregates of <10 nm chalcopyrite NPs from riverbeds of the Clark Fork River Superfund Complex (U.S.A.) (Hochella et al. 2005), and ≤50 nm chalcopyrite spheroids in hydrothermal emissions from the Mid-Atlantic Ridge (Gartman et al. 2014) are similar in size and morphology to the synthetic NPs in our study. In contrast, 100–200 nm mackinawite nanorods and 400–500 nm greigite nanoplates found in the Yangtze Estuary sediments (Niu et al. 2018) and 10–20 μm euhedral crystals of covellite observed in Whipsaw Creek, British Columbia (Canada) (Lett and Fletcher 1980) are distinctly larger than any NPs in our study. Overall, these comparisons indicate that laboratory experiments can form metal sulfides that have similar size and morphology to natural precipitates in some environments.

It is clear from our study that under any conditions akin to complex environmental systems, a mixture of metal sulfide NPs with variable physicochemical attributes will be produced. This will be especially true in dynamic environments when the ratio of metals can vary due to external factors (variable supply rate or solubility due to changes in redox) or even as competing precipitation reactions progress at different rates. The presence of bacteria and associated extracellular compounds further act to concentrate reaction zones and to encourage the formation of mixed-metal sulfides with distinct crystal structures but do not display any overt features that can be used to distinguish biogenicity (at least for Fe-Cu-sulfide NPs). We, therefore, expect metal sulfide NPs to be extremely diverse even over small sampling scales.

The properties of the precipitated metal sulfide NPs greatly affect their reactivity, stability, and transport properties with implications relevant to heavy metal immobilization and release to surrounding waters. As a general rule for single phase (or metal) systems, smaller crystals will have the highest surface area normalized reactivity and may dominate geochemical reactions. In mixed-metal systems, one also needs to consider the formation of different phases with different solubilities (chalcopyrite vs. nukundamite vs. covellite for Cu; mackinawite vs. Cu-rich mackinawite for Fe). Additionally, the incorporation of foreign metals may cause defects and size modifications to the crystal structure that can potentially enhance or lower reaction rates, especially for NPs. Future dissolution and bio-availability studies on NPs generated from mixed-metal systems will greatly help in identifying the most reactive component(s) that may also point to the main driver of geochemical reactions in nature.

FUNDING

This work was funded by the grant DOE-BES DE-FG02-06ER15786 awarded by the Department of Energy to Mike Hochella, Mitsu Murayama, and Jie Xu and by the start-up grant to Jie Xu by the University of Texas at El Paso. This work uses shared facilities at the Virginia Tech National Center for Earth and Environmental Nanotechnology Infrastructure (NanoEarth), a member of the National Nanotechnology Coordinated Infrastructure (NNCI) network, supported by NSF (NNCI 1542100). NanoEarth is housed at Virginia Tech's Institute for Critical Technology and Applied Sciences (ICTAS).

ACKNOWLEDGMENTS

We thank the associate editor, Andrew Madden and an anonymous reviewer for helpful comments that improved the manuscript.

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MANUSCRIPT RECEIVED OCTOBER 15, 2018

MANUSCRIPT ACCEPTED JANUARY 15, 2019

MANUSCRIPT HANDLED BY ANDREW MADDEN

Endnote:

¹Deposit item AM-19-56848, Supplemental Material. Deposit items are free to all readers and found on the MSA website, via the specific issue's Table of Contents (go to http://www.minsocam.org/MSA/AmMin/TOC/2019/May2019_data/May2019_data.html).