# Pyrite formation from FeS and H<sub>2</sub>S is mediated by a novel type of microbial energy metabolism

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### 23 Abstract

24 The exergonic reaction of FeS with H<sub>2</sub>S to form FeS<sub>2</sub> (pyrite) and H<sub>2</sub> was postulated to have 25 operated as an early form of energy metabolism on primordial Earth. Since the Archean, 26 sedimentary pyrite formation played a major role in the global iron and sulfur cycles, with direct 27 impact on the redox chemistry of the atmosphere. To date, pyrite formation was considered a 28 purely geochemical reaction. Here, we present microbial enrichment cultures, which grew with FeS, H<sub>2</sub>S, and CO<sub>2</sub> as their sole substrates to produce FeS<sub>2</sub> and CH<sub>4</sub>. Cultures grew over 29 periods of three to eight months to cell densities of up to 2–9×10<sup>6</sup> cells mL<sup>-1</sup>. Transformation of 30 FeS with H<sub>2</sub>S to FeS<sub>2</sub> was followed by <sup>57</sup>Fe Mössbauer spectroscopy and showed a clear 31 biological temperature profile with maximum activity at 28°C and decreasing activities towards 32 33 4°C and 60°C. CH<sub>4</sub> was formed concomitantly with FeS<sub>2</sub> and exhibited the same temperature 34 dependence. Addition of either penicillin or 2-bromoethanesulfonate inhibited both FeS<sub>2</sub> and CH<sub>4</sub> production, indicating a syntrophic coupling of pyrite formation to methanogenesis. This 35 36 hypothesis was supported by a 16S rRNA gene-based phylogenetic analysis, which identified 37 at least one archaeal and five bacterial species. The archaeon was closely related to the 38 hydrogenotrophic methanogen Methanospirillum stamsii while the bacteria were most closely related to sulfate-reducing Deltaproteobacteria, as well as uncultured Firmicutes and 39 40 Actinobacteria. We identified a novel type of microbial metabolism able to conserve energy 41 from FeS transformation to FeS<sub>2</sub>, which may serve as a model for a postulated primordial iron-42 sulfur world.

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### 43 Significance statement

44 Pyrite is the most abundant iron-sulfur mineral in sediments. Over geological times, its burial controlled oxygen levels in the atmosphere and sulfate concentrations in seawater. Its 45 formation in sediments is so far considered a purely geochemical process that is at most 46 47 indirectly supported by microbial activity. We show that lithotrophic microorganisms can directly 48 transform FeS and H<sub>2</sub>S to FeS<sub>2</sub> and use this exergonic reaction as a novel form of energy 49 metabolism that is syntrophically coupled to methanogenesis. Our results provide insights into 50 a syntrophic relationship that could sustain part of the deep biosphere and lend support to the 51 iron-sulfur-world theory that postulated FeS transformation to FeS<sub>2</sub> as a key energy-delivering 52 reaction for life to emerge.

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### 53 Introduction

54 With an annual formation of at least 5 million tons, pyrite (FeS<sub>2</sub>) is the thermodynamically 55 stable end product of iron compounds reacting with sulfide in reduced sediments, with the latter being produced mainly by microbial sulfate reduction. Consequently, pyrite is the most 56 57 abundant iron-sulfur mineral on Earth's surface (1). Over geological times, burial of pyrite was tightly intertwined with organic matter preservation in reduced sediments (2). These massive 58 reservoirs of reduced sulfur and carbon are being counterbalanced by the photosynthetically 59 60 produced oxygen in the Earth's atmosphere (2). Despite this importance of pyrite for Earth's 61 iron, sulfur, and carbon cycles as well as Earth's surface redox state, the mechanism of pyrite formation in natural environments is still being debated (1). Currently, two mechanisms are 62 63 discussed to drive pyrite formation in sediments, which both preclude dissolution of 64 precipitated iron(II) monosulfide (FeS) to an aqueous FeS intermediate. In the polysulfide pathway, FeS<sub>aq</sub> is attacked by nucleophilic polysulfide to form FeS<sub>2</sub> (equ. 1). Alternatively, 65 66 pyrite may form from the reaction of  $FeS_{aq}$  with  $H_2S$  (equ. 2), which is known as the  $H_2S$ pathway or the Wächtershäuser reaction (1). 67

68 Equ. 1: FeS +  $S_n^{2-} \rightarrow FeS_2 + S_{n-1}^{2-}$   $\Delta G_r^{0'} = -64 \text{ to } -75 \text{ kJ mol}^{-1}$ 69 Equ. 2: FeS +  $H_2S \rightarrow FeS_2 + H_2$   $\Delta G_r^{0'} = -41 \text{ kJ mol}^{-1}$ 

Using inorganic experimental systems, abiotic pyrite formation has been observed at temperatures of 25–125°C (e.g. 1, 3). However, already trace amounts of organic compounds containing aldehyde groups were reported to inhibit pyrite formation in such experiments (1, 4). Absence of stringent control for such compounds may explain why many other studies with abiotic systems did not observe pyrite formation at environmentally relevant temperatures (5-

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75 11). On the other hand, pyrite formation is known to take place in the presence of organic 76 matter and especially alive microorganisms in sediments (5, 12). Indeed, pyrite formation could 77 be observed as a side reaction in pure and enrichment cultures of heterotrophic sulfate-78 reducing and chemolithotrophic sulfur-dismutating bacteria, with the assumption that these 79 microorganisms provide mainly H<sub>2</sub>S as a substrate for concomitant abiotic pyrite formation (13-80 15). In addition, a more complex involvement of microorganisms has been postulated that 81 includes the support of nucleation of FeS minerals on bacterial cell surfaces (16). However, in 82 all these studies biogenic pyrite formation has so far been understood only as an indirect 83 abiotic process.

Pyrite formation according to equ. 2 provides reducing equivalents in the form of H<sub>2</sub> that could 84 85 be coupled to the reduction of  $CO_2$  to  $CH_4$  or more complex organic matter. Coupling of pyrite 86 formation to methanogenesis has been proposed by Jørgensen and coworkers to be part of a 87 cryptic sulfur cycle in deep marine sediments where it could support the enigmatic life forms of 88 the deep biosphere (17). Coupling of this reaction to the synthesis of organic matter is the basis of the "iron-sulfur world" theory proposed by Wächtershäuser, by which pyrite formation 89 90 is viewed as the central process that led to the transition from Fe-S surface-catalyzed 91 synthesis of organic molecules to actual life on the primordial Earth (18-20). Here, we show for 92 the first time that the reaction of FeS with H<sub>2</sub>S to form FeS<sub>2</sub> can be used by microorganisms to 93 conserve energy for lithotrophic growth if syntrophically coupled to methanogenesis.

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### 94 Results and Discussion

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### 95 Pyrite formation as a microbially catalyzed process

Mineral medium containing 5 mM FeS and 6 mM H<sub>2</sub>S as sole substrates and CO<sub>2</sub>/HCO<sub>3</sub><sup>-</sup> as 96 97 carbon source was inoculated with digested sewage sludge, freshwater or marine sediment 98 (Table S1) and incubated at 16°C or 28°C. Microbial activity was followed via methane 99 formation, and transfers were made every three to eight months, typically when the methane 100 content in the headspace approached a plateau. Of seven enrichments started, four exhibited 101 methane formation for more than ten transfers. The most active enrichment culture J5, which 102 was started from digested sewage sludge and incubated at 28°C, was characterized in more 103 detail after more than 20 transfers. On average, the methane content reached 0.7 mmol per L 104 of culture in J5. In contrast, no methane formation was observed in abiotic controls (Fig. 1A). 105 This was mirrored in the turnover of total H<sub>2</sub>S: While in culture J5 total H<sub>2</sub>S decreased over 106 time from approx. 6 mmol to 0.04-1.1 mmol per L of culture (Fig. S1), abiotic controls showed 107 a much less pronounced decline of total H<sub>2</sub>S (3.7 mmol residual H<sub>2</sub>S per L). The observed 108 decrease in the abiotic controls may be due to inorganic background reactions (see below).

109 Conversion of FeS solids was followed by <sup>57</sup>Fe Mössbauer spectrometry. After nearly seven 110 months of incubation, the Mössbauer spectrum of culture J5 was dominated by a FeS<sub>2</sub> doublet 111 (Fig. 1B), which corresponded to 53-63% of the iron-sulfur mineral phase (Table S2). In 112 contrast, no evidence of a singlet peak corresponding to FeS was present. In addition, a poorly 113 defined sextet feature was required to achieve a satisfactory fit of the Mössbauer spectrum. 114 This poorly defined sextet is best described as a metastable phase, which we have termed 115  $FeS_x$  in accordance with the notation used by Wan et al. (21), and represented 31–39% of the 116 remaining iron-sulfur minerals. Furthermore, a well-defined sextet with a hyperfine magnetic

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field of 32 T was required to fit the data, which corresponded well to greigite ( $Fe_3S_4$ ) (22) and made up 7–8% of the remaining iron-sulfur minerals. Greigite is the sulfur isomorph of magnetite and was previously observed as an intermediate phase in the FeS conversion to pyrite in abiotic studies (23, 24).

121 FeS<sub>2</sub> formation in culture J5 was confirmed by X-ray diffraction analysis, which recovered all 122 major XRD reflections of pyrite in the obtained XRD pattern (Fig. 2A). Since no indication of a 123 parallel formation of the dimorph marcasite was evident from the XRD pattern, the recovered 124 FeS<sub>2</sub> phase is referred to as pyrite from here on. Further support for pyrite formation in culture 125 J5 was provided by scanning electron microscopy (SEM) coupled to energy-dispersive X-ray 126 (EDX) spectroscopy. Here, µm-scale iron-sulfur crystals with a euhedral structure as typical of 127 pyrite could be observed (Fig. 2B), which resembled the expected Fe:S ratio of 1:2 as revealed 128 by EDX point measurements (Fig. S2).

In contrast to culture J5, the Mössbauer spectrum of the nearly seven-months-old abiotic 129 130 control was dominated by a prominent FeS singlet peak (64% of iron-sulfur minerals). In 131 addition, a poorly defined sextet corresponding to FeS<sub>x</sub> was required to achieve a satisfactory 132 fit of the obtained data (36% of iron-sulfur minerals) (Fig. 1, Table S2). The abiotic conversion 133 of FeS to FeS<sub>x</sub> most likely explains the observed decrease of total H<sub>2</sub>S in the abiotic control. 134 Absence of pyrite formation in abiotic controls was further supported by the obtained XRD 135 patterns (Fig. 2A). In addition, freshly prepared medium was also devoid of pyrite as evidenced 136 by an overall disordered iron-sulfur mineral phase in SEM-EDX images without any euhedral 137 crystals indicative of pyrite (Fig. 2B, Fig. S2).

To further support our hypothesis of biogenic pyrite formation in culture J5, we followed its iron sulfur mineral transformation over a temperature gradient of 4–60°C. The maximum of pyrite
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formation was evident at 28°C. Incubation at lower (16°C) or higher (46°C) temperatures resulted in decreased pyrite formation, with no pyrite formation at 4°C and 60°C (Fig. 3*A* and 3*B*, Table S2). While FeS was completely transformed to pyrite, greigite, and various FeS<sub>x</sub> phases at 16, 28, and 46°C, some residual FeS remained at 4°C and 60°C (Fig. 3*A*). The observed temperature profile of pyrite formation is typical of biologically catalyzed reactions centered on an optimum reaction temperature. In contrast, abiotic pyrite formation at temperatures of <100°C was shown to follow a sigmoidal temperature dependence (3).

### 147 Microbial pyrite formation is a syntrophically coupled process

148 Methane formation closely followed the temperature-dependent activity profile of pyrite 149 formation (Fig. 3C) suggesting metabolic coupling of both processes. This hypothesis was 150 further explored in an inhibition experiment using either penicillin G as a generic inhibitor of 151 bacterial cell wall synthesis or 2-bromoethanesulfonate (BES) as a specific inhibitor of 152 hydrogenotrophic methanogenesis (25). BES inhibited both methane and pyrite formation 153 completely (Fig. 1C and 1D). We interpret this as a direct coupling of biogenic pyrite formation 154 to hydrogenotrophic methanogenesis. In support of this hypothesis, penicillin inhibited both 155 pyrite and methane formation as well (Fig. 1C and 1D). Here, methane formation ceased after 156 an initial production of 0.15 mmol per L of culture. The latter is explained by penicillin's mode of 157 action, which inhibits growth of bacteria but not their initial metabolic activity. Interestingly, a 158 corresponding small amount of pyrite was not observed in this experiment but rather a partial 159 FeS transformation to various  $FeS_x$  phases and greigite (Fig. 1*D*, Table S2). This indicates that 160 methanogenesis could receive reducing equivalents also from these iron-sulfur mineral 161 transformations. Further support for syntrophic coupling of pyrite formation to methanogenesis 162 was provided by a third inhibition experiment in which penicillin addition was supplemented by

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163 79% H<sub>2</sub> in the headspace. Also here, pyrite was not formed (Table S2) while methane 164 production was stimulated more than 10-fold by the added  $H_2$  (>10 mmol per L of culture). This 165 clearly showed that methanogenesis could be uncoupled from pyrite formation and is essential 166 for the latter, most likely to keep H<sub>2</sub> or another electron carrier at a low level, to make pyrite 167 formation energetically more favorable as is typically observed in syntrophic processes (26). Although we could not identify the exact electron carrier, molecular H<sub>2</sub> is the most likely 168 169 candidate because it was previously observed in abiotic experiments of pyrite formation from 170 FeS and H<sub>2</sub>S (3, 27). Irrespective of the actual electron carrier, syntrophic coupling of pyrite 171 formation to methanogenesis would be exergonic and result in an expected ratio of formed 172 pyrite to methane of 4:1 (equ. 5).

173	Equ. 3:	4 FeS + 4 H <sub>2</sub> S → 4 FeS <sub>2</sub> + 4 H <sub>2</sub>	$(\Delta G_r^{\circ}) = -41 \text{ kJ/mol FeS}_2$
174	Equ. 4:	$CO_2$ + 4 $H_2$ → $CH_4$ + 2 $H_2O$	$(\Delta G_r^{\circ}) = -131 \text{ kJ/mol CH}_4)$
175	Equ. 5: 4 F	$eS + 4 H_2S + CO_2 \rightarrow 4 FeS_2 + CH_4 + 2 H_2O$	(ΔG <sub>r</sub> °' = –295 kJ/mol CH <sub>4</sub> )

176 Indeed, the ratio of pyrite to methane formed in culture J5 was 4.1:1 and 3.2:1 in two 177 independent experiments at 28°C (Table S2), respectively, thus confirming the proposed 178 overall reaction stoichiometry.

### 179 Exergonic pyrite formation supports microbial growth

Culture J5 was transferred more than 20 times on medium containing FeS,  $H_2S$  and  $CO_2$  as sole substrates. This indicates a strict dependence on syntrophic pyrite formation as the only energy-yielding reaction observed. Within incubation periods of 83–248 days, cell densities increased by more than one order of magnitude, from 2×10<sup>5</sup> cells mL<sup>-1</sup> to 2–9×10<sup>6</sup> cells mL<sup>-1</sup> (Fig. S3). Cells were typically found to be attached to mineral surfaces (Fig. 4*C*); however,

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185 there was no indication of cell encrustation (Fig. 4D). Assuming an average cell volume of 186 about 1 µm<sup>3</sup> and a dry mass content of 33% (28), our measured cell densities correspond to a maximum of ca. 3 mg dry cell mass  $L^{-1}$ . If formation of ATP requires 60–70 kJ mol<sup>-1</sup> (29) and if 187 188 - under ideal growth conditions - 10.5 g biomass (dry weight) can be synthesized at the 189 expense of 1 mol ATP (30), a complete conversion of 5 mM FeS + 5 mM H<sub>2</sub>S according to 190 equ. 5 could yield 4–5 mM ATP or ca. 50 mg dry cell mass  $L^{-1}$ . Of course, the conditions of 191 lithoautotrophic growth in our enrichment cultures are entirely different from those used in the 192 growth experiment by Bauchop and Elsden, with heterotrophic growth in an organically rich 193 medium. Moreover, formation of pyrite from FeS is an extremely slow process catalyzed at or 194 close to mineral surfaces with very slow substrate turnover, which implies that major amounts 195 of metabolic energy have to be invested in cell maintenance without concomitant growth (31). 196 Thus, it is not surprising that our cell yields are lower than estimated above. If every partial 197 reaction in equ. 5 (5 in total; 4 × FeS transformation, 1 × CH<sub>4</sub> formation) uses about 20 kJ for 198 synthesis of a minimum of 1/3 of an ATP equivalent (29), the total process could synthesize 199 5/3 mmol ATP equivalents per liter. Considering that in culture J5 a maximum of 62.5% FeS 200 conversion to  $FeS_2$  was observed (Table S2), the expected cell yield would – optimally – be 12 mg cell dry matter  $L^{-1}$ , which is close to the measured cell yield. 201

202 The community composition of culture J5 was analyzed by 16S rRNA gene clone libraries. All 203 16S rRNA gene sequences derived from amplification with a universal archaeal primer set 204 belonged to the same species-level operational taxonomic unit (OTU, 99% sequence identity) 205 Methanospirillum stamsii (Fig. and showed 97.6% sequence identity to 4*B*), a 206 hvdrogenotrophic isolated methanogen from а low-temperature bioreactor (32). 207 Methanogenesis in culture J5 is likely performed by this OTU. Using a universal bacterial

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208 primer set, we detected five bacterial OTUs (Fig. 4A). The majority of clones (76%) belonged 209 to OTU 3, which showed 99.9% sequence identity with Desulfomicrobium baculatum, a sulfate 210 reducer within the class Deltaproteobacteria (33, 34). Further OTUs related to 211 Deltaproteobacteria were OTU 2 (12% rel. abundance) and OTU1 (6% rel. abundance). OTU 2 212 was closely related to Desulfovibrio sulfodismutans (97.9% sequence identity), which is 213 capable of dismutating thiosulfate or sulfite (35), while OTU 1 was distantly related (96.2% 214 sequence identity) to Smithella propionica, which is known to degrade propionate in syntrophy 215 with methanogens (36). The remaining two bacterial OTUs were either distantly related (<91% 216 sequence identity) to cultured members of the Firmicutes (OTU 4, 3% rel. abundance) or 217 Actinobacteria (OTU 5, 3% rel. abundance). Interestingly, all OTUs belonging to the 218 Deltaproteobacteria and Firmicutes fell into larger clusters that include cultured representatives 219 with a sulfur-related energy metabolism. Therefore, it is tempting to speculate that enzymes 220 operating in the respective sulfur transformation pathways might be involved in the microbial conversion of FeS to FeS<sub>2</sub>. 221

### 222 Conclusion

223 Pyrite is produced in massive quantities in today's sediments (2). However, its formation in 224 nature is far from understood, especially because its nucleation is kinetically hindered (1). The 225 presence of sulfide-producing microorganisms as passive pyrite nucleation sites indicated 226 support for abiotic pyrite formation (13-15), but could not be reproduced in every bacterial 227 model system (6). We show that pyrite formation can be mediated by microorganisms to 228 overcome the kinetic hurdle of nucleation, but as an essential part of their energy metabolism 229 and not just as a mere abiotic side reaction on their cell surface. This may help to explain the 230 ambiguous results published so far on the role of microorganisms in this process. Since we

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found only one archaeal species closely related to methanogens in our enrichments, it is likely that one or several of the bacterial partners actually catalyze FeS transformation to pyrite. An exciting question currently remaining is whether these bacteria utilize internally the H<sub>2</sub>S pathway (equ. 2) or a combination of sulfide oxidation to zero-valent sulfur coupled to the polysulfide pathway (equ. 1) to finally produce pyrite (Fig. S4).

236 Our results further showed that the reducing equivalents released from FeS transformation to 237 pyrite can be transferred to methanogenesis. This opens an interesting perspective on the 238 metabolic versatility sustaining the vast deep biosphere inhabiting the Earth's subsurface (37). 239 While recalcitrant organic matter or H<sub>2</sub> released by radiolytic cleavage of water (38, 39) have 240 been proposed to sustain the enigmatic life forms of the deep biosphere, there is also 241 experimental evidence of a cryptic sulfur cycle within deep sediments that would include pyrite 242 formation coupled to methanogenesis to be functional (17). Our results show that this missing 243 link could indeed be mediated by microorganisms and supply energy to support microbial 244 growth. Since the redox potential (E<sub>h</sub>°') of the FeS/FeS<sub>2</sub> couple is –620 mV at circumneutral pH 245 (40), it is well suited to provide reducing equivalents also for CO<sub>2</sub> fixation to acetate 246  $(E_h^{\circ})^{\circ} = -290 \text{ mV}$  and more complex organic matter in the pyrite-forming microorganism. 247 Wächtershäuser proposed in his "iron-sulfur world" theory that exactly this mechanism was the 248 basis for an autocatalytic metabolism and the resulting evolution of life at hydrothermal vents 249 on primordial Earth (e.g. 20, 41). Our enrichment cultures may serve as a model to understand 250 the enzymatic background of this hypothesis.

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### 251 Methods

### 252 Cultivation

253 Enrichment cultures were initiated and maintained in carbonate-buffered, sulfide-reduced 254 (1 mM) freshwater mineral medium (42) supplemented with selenite-tungstate solution (43), 7-255 vitamin solution (42), and trace element solution SL10 (44). The medium was prepared and 256 stored under a N<sub>2</sub>/CO<sub>2</sub> atmosphere (80:20). The final pH was adjusted to 7.2 to 7.4. From a 257 CO2-neutralized sulfide stock solution (45), 350 µmol H2S was added to 70 mL mineral 258 medium in 180 mL serum bottles that were sealed with butyl rubber stoppers. Since 1 mM H<sub>2</sub>S 259 was already present as reducing agent in the mineral medium, the final amount of H<sub>2</sub>S was 420 µmol. FeS was prepared from anoxic solutions of 0.4 M FeCl<sub>2</sub> and 0.4 M Na<sub>2</sub>S. The 260 261 resulting FeS precipitate was washed at least once and re-suspended in oxygen-free distilled water. For Mössbauer spectroscopy analysis, FeS was prepared from a FeCl<sub>2</sub> solution that 262 contained 10% <sup>57</sup>FeCl<sub>2</sub> to enhance signal quality. 350 µmol FeS was added to 70 mL mineral 263 264 medium. Enrichment cultures were incubated in the dark at 28°C if not indicated otherwise. For 265 inhibition experiments, cultures were supplemented with either penicillin-G (1,000 U mL<sup>-1</sup>) or 2-266 bromoethanesulfonate (10 mM). Abiotic controls were run without inoculum.

### 267 Monitoring of substrate turnover

For total dissolved sulfide measurements [ $\Sigma(H_2S_{aq}, HS^-, S^{2-})$ ], 100-µL samples were taken from liquid cultures without disturbing the precipitated iron sulfide minerals, and directly transferred to 100 µL of an anoxic 0.2 M NaOH solution. From the alkalinized sample, 10– 20 µL were fixed in 100 µL of a 0.1 M zinc acetate solution, and sulfide was quantified by the methylene blue method (46). The corresponding amount of H<sub>2</sub>S in the headspace was calculated using Henry's law and a temperature-adjusted k-value of 0.093 (28°C, 47). CH<sub>4</sub> was

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measured by gas chromatography with a flame ionization detector (SRI Instrument SGI 8610C) using a consecutive arrangement of a Porapak (80/100 mesh; 1 m × 2 mm) and a Hayesep-D packed column (80/100 mesh; 3 m × 2 mm). The injector and column temperatures were 60°C, and the detector temperature was 135°C. The carrier gas was N<sub>2</sub> at a flow rate of 3.2 ml min<sup>-1</sup>.

278 Chromatograms were recorded with the PeakSimple v4.44 chromatography software.

Iron-sulfide minerals were analyzed by <sup>57</sup>Fe Mössbauer spectroscopy. Within an anoxic 279 280 glovebox (100% N<sub>2</sub>), the enrichment culture was passed through a 0.44-µm filter and then 281 sealed between two pieces of air-tight Kapton tape. Samples were transferred to a Mössbauer 282 spectrometer (WissEl, Starnberg) within an airtight bottle filled with 100% N<sub>2</sub> that was only opened immediately prior to loading the samples inside the closed-cycle exchange gas 283 284 cryostat (Janis cryogenics). Measurements were collected at a temperature of 5 K with a constant acceleration drive system (WissEL) in transmission mode with a <sup>57</sup>Co/Rh source and 285 calibrated against a 7  $\mu$ m thick  $\alpha$ -<sup>57</sup>Fe foil measured at room temperature. All spectra were 286 287 analyzed using Recoil (University of Ottawa) by applying the Voight Based Fitting (VBF) routine (48). The half width at half maximum (HWHM) was fixed to a value of 0.138 mm s<sup>-1</sup> for 288 289 all samples.

290 X-ray diffraction (XRD) patterns were recorded with the D8 Discover system (Bruker) with I $\mu$ S 291 radiation source (2 mm in diameter), and a Lynxexe XE detector. Samples were dried for 2 h 292 under a continuous stream of 100% N<sub>2</sub> and measured within 48 h under air as described 293 previously (49). Measurements were done using CuK $\alpha$  rays in angles ranging from 10–70° 2 $\Theta$ 294 in 0.02° steps with 2,880 sec measuring time and a total measuring time of 12 h and 47 min. 295 The resulting spectra were compared with spectra provided in the international crystal structure

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database (ICSD), FIZ Karlsruhe (version 2016/2) using the software DIFFRAC.EVA (version
4.1.1, Bruker).

### 298 Scanning electron microscopy coupled to energy dispersive x-ray spectroscopy (SEM-EDX)

299 For SEM-EDX analysis, 1 mL of culture was centrifuged at 4,500 ×g for 10 min. 200 µL of the resulting pellet was transferred on gelatin-coated glass slides. Samples were fixed in 1 mL 300 301 2.5% glutaraldehyde in 0.1 M HEPES-buffer containing 0.01 M KCI (HEPES-KCI) and in 2% 302 OsO<sub>4</sub> in HEPES-KCl for 60 min each. Fixed samples were dehydrated in a graded ethanol 303 series (30%, 50%, 70%, 80%, 90%, 96% and absolute ethanol) for 30 min each. Thereafter, 304 samples were critical-point dried under CO<sub>2</sub> in a Bal-Tec CPD030 (Balzers). Sputter coating of 305 6 nm platinum was done in a Quorum Q150R ES sputter coater (Quorum Technologies) and 306 micrographs were taken with a FESEM Auriga 40 (Zeiss). EDX mappings and point 307 measurements were taken at a working distance of 5 mm with an Oxford X-Max detector 308 (Oxford Instruments) and at 10 kV and 15 kV, respectively. Point measurements were 309 normalized to 10,000 counts within a K $\alpha$  energy of 6.3–6.5 keV. Sample preparation for cell 310 counts by fluorescent microscopy is described in detail in the Supporting Information.

### 311 Phylogenetic analysis

Total genomic DNA was extracted from 50 mL of a 4.5-months old culture using a phenolbased beat-beating protocol modified after Loy, Beisker and Meier (50). Subsequent amplification of bacterial or archaeal 16S rRNA genes was done using standard PCR protocols based on universal primers. Details are given in the Supporting Information. 16S rRNA gene clone libraries were constructed using the TOPO® TA Cloning® Kit (ThermoFisher Scientific). Bacterial or archaeal 16S rRNA gene fragments were aligned by use of the SINA webaligner (51) to the non-redundant 16S rRNA gene database v.123.1 available on the SILVA online

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319 platform (52, www.arb-silva.de) and imported into ARB for initial phylogenetic analysis (53). 320 OTU clustering was performed in mothur v.1.22.2 (54) using the furthest neighbor approach 321 and a 99%-identity cutoff to delineate OTUs at the approximate species level (55). For 322 phylogenetic inference of 16S rRNA gene fragments representing individual OTUs, Maximum 323 Likelihood (ML) trees were calculated using RAxML v8.2.9 (56) as implemented on the 324 CIPRES webserver (57, www.phylo.org). Using a 50% conservation filter of nucleic acid 325 positions within the domain Bacteria, a RAxML tree was inferred from 1,102 unambiguously 326 aligned nucleic acid positions for bacterial 16S rRNA genes. The reconstruction of the archaeal 327 tree followed the same outline but using 752 unambiguously aligned nucleic acid positions and 328 no conservation filter because of the close relatedness of all included sequences. Calculations 329 were based on the GTRGAMMA distribution model of substitution rate heterogeneity. MRE-330 based bootstrap analysis stopped after 204 and 102 replicates for the bacterial and archaeal 331 16S rRNA gene tree, respectively. Sequences are available from NCBI GenBank under 332 accession numbers MH665848-MH665880 and MH665881-MH665889 for Bacteria and 333 Archaea, respectively.

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### 340 Figures

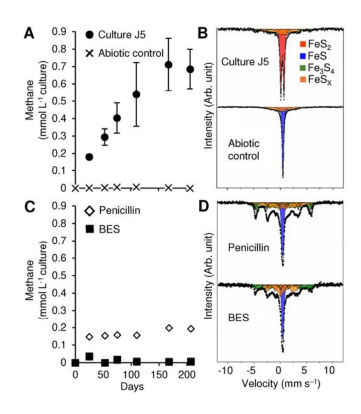
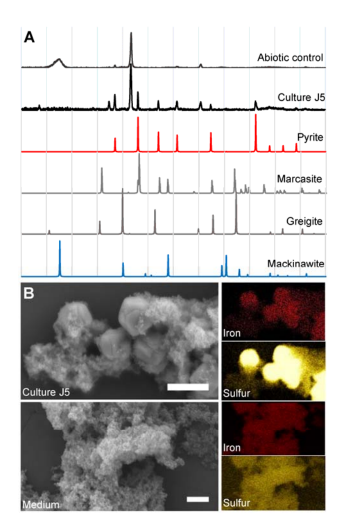


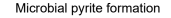
Figure 1. Time-resolved CH<sub>4</sub> formation in comparison to iron-sulfur mineral composition after nearly seven months of incubation (207 days) in culture J5 as compared to abiotic controls and incubations of culture J5 with penicillin-G (1,000 U ml<sup>-1</sup>) or 2-bromoethanesulfonate (BES, 10 mM). (A) and (C) show the mean  $\pm$  one standard deviation of CH<sub>4</sub> measurements of three independent incubations. Standard deviations are often smaller than the actual symbol size. (B) and (D) show Mössbauer spectra corresponding to the last time point in the presented time series with FeS<sub>2</sub> in red, FeS in blue, Fe<sub>3</sub>S<sub>4</sub> in green, and intermediate FeS<sub>x</sub> phases in orange.

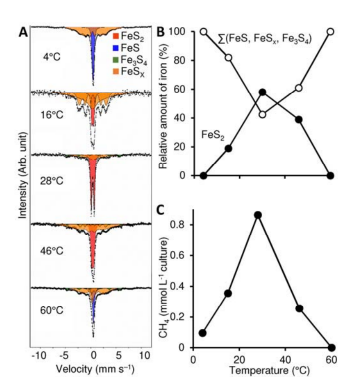


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350 Figure 2. (A) Representative X-ray diffractograms of mineral precipitates formed in culture J5 351 and an abiotic control setup after 9 months of incubation (281 days). Diffraction data of the two 352  $FeS_2$  dimorphs pyrite and marcasite as well as of  $Fe_3S_4$  (greigite) and FeS (mackinawite) are given as reference. (B) Scanning electron microscopy images of a nearly 7-months old (211 353 354 days) culture J5 in comparison to freshly prepared medium without inoculum. The scale bar 355 represents 2 µm. Images to the right show the corresponding results from energy dispersive X-356 ray spectroscopy (EDX). Besides atoms from medium salts, iron and sulfur were the only 357 elements discovered in the mineral phases.

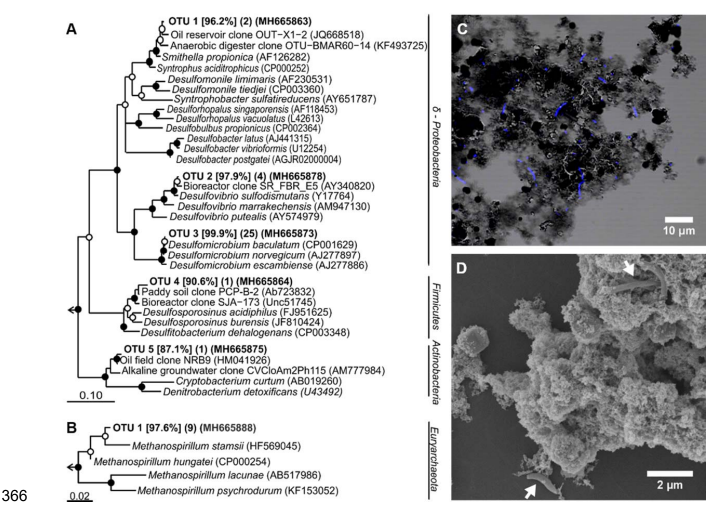




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Figure 3. Temperature-dependent pyrite and methane formation in culture J5 after nearly 7 months of incubation (207 days). A) Mössbauer spectra showing the temperature-dependent iron-sulfur mineral composition (FeS<sub>2</sub> in red, FeS in blue, Fe<sub>3</sub>S<sub>4</sub> in green, and intermediate FeS<sub>x</sub> phases in orange). B) Relative abundance of pyrite (FeS<sub>2</sub>) in comparison to all other measured iron-sulfur minerals plotted against temperature as the explanatory variable. Details are provided in Table S2. C) Average amount of methane (n=2) plotted against temperature as the explanatory variable.

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367 Figure 4. Bacterial and archaeal community composition of enrichment culture J5. RAxML 368 trees based on bacterial (A) and archaeal (B) 16S rRNA gene sequences obtained from culture 369 J5. Representative sequences of OTUs at the approximate species level (99% sequence 370 identity) are shown. Sequence identity to the next cultured relative is given in percent in square 371 brackets. Numbers of clones from the same OTU are presented in parenthesis followed by the 372 GenBank accession number of a representative sequence. Bootstrap support is indicated by 373 closed (≥90%) and open (≥70%) circles at the respective branching points. The scale bar 374 represents 10% (Bacteria) and 2% (Archaea) estimated sequence divergence. (C) Combined 375 phase contrast image and fluorescent image of DAPI-stained cells and (D) scanning electron

### Microbial pyrite formation

- 376 microscopy image with cells indicated by white arrows of culture J5 after 7.4 and 10 months of
- 377 incubation, respectively.

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# 510 Supporting Information

### 511 Supporting Materials and Methods

512 Cell counts by fluorescence microscopy. For cell counts, 0.5 mL culture was fixed overnight 513 in 9.5 mL freshly prepared paraformaldehyde solution (4%), subsequently centrifuged at 514 10,000 × g for 10 min at 4°C and re-suspended in 1 mL PBS [130 mM NaCl, 5% (v/v) 515 phosphate buffer (40 mM NaH<sub>2</sub>PO<sub>4</sub>, 160 mM Na<sub>2</sub>HPO<sub>4</sub>), pH 7.2] and 9 mL ammonium oxalate 516 solution (5.6 g ammonium-oxalate and 4.2 g oxalic acid dihydrate in 200 mL distilled water). 517 Samples were vortexed for 10 min to dissolve most of the iron sulfide minerals so that cells 518 could be collected on a 0.2 µm pore size filter (GTTP-white, Millipore). Filters were air-dried and stored at  $-20^{\circ}$ C. Filter sections were stained with a 1 µg mL<sup>-1</sup> 4',6-diamidino-2-519 520 phenylindole (DAPI) solution and incubated for 10 min in the dark. Thereafter, filters were washed for 5 min in distilled water, followed by two 1-min washing steps in 80% ethanol. Dry 521 522 DAPI-stained filters were mounted on microscope slides using CitiFluor™ AF1. For 523 fluorescence microscopy, an inverted microscope (AxioObserver, Zeiss) with a 40x/0.60 LD-524 PlanNeofluar objective was used. Z-stacks were acquired with a distance of 0.28 µm. Image 525 processing involved 3-dimensional deconvolution of each stack using a theoretical PSF with 526 ZEN Black (Zeiss AG). Cells were counted using an image processing workflow set up in 527 KNIME 3.4.0 (58) using orthogonal projections of the de-convoluted input stacks. The workflow is available at https://github.com/bic-kn/cell-counting-workflow. 528

529 DNA extraction. The DNA extraction protocol was adopted from (50). A 4.5-month old 50-mL
 530 culture was harvested after CH<sub>4</sub> concentrations reached a plateau of 2.1% in the headspace
 531 (corresponding to 55 µmol produced CH<sub>4</sub>). Harvesting was done by 10 min of centrifugation at
 532 6,000 × g. The pellet was re-suspended in 400 µL autoclaved TE-Buffer (10 mM Tris & 1 mM
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533 EDTA in MQ water, pH 8) and stored for three hours at -20°C. Cells were thawed on ice, 534 mixed with heat-sterilized zirconium beads (0.1 mm), 600 µL phenol/chloroform/isoamylalcohol 535 (25:24:1, Carl Roth), and 150 µL of a 10% sterile-filtered SDS-solution in a screw-cap tube, 536 and vigorously shaken for 20 min using a vortexer. After centrifugation for 20 min at 20,817  $\times q$ 537 and 4°C, the aqueous supernatant was transferred to a new reaction tube. Because the 538 aqueous phase was hardly visible due to remaining iron sulfide minerals, another 10-minute 539 centrifugation step was used to remove residual phenol from the extract. DNA was precipitated 540 by incubation at -20°C overnight in 0.1 volume of 3 M Na-acetate (in MQ-water, autoclaved) 541 and 2.5 volumes absolute ethanol. Afterwards, the pellet was washed twice with 70% ethanol, 542 dried for 5 min, and re-suspended in 50 µL DNase- and RNAse- free H<sub>2</sub>O. DNA concentrations 543 were quantified fluorimetrically using Quant-iT PicoGreen (Invitrogen).

544 16S rRNA gene clone library. Amplification of bacterial 16S rRNA genes was performed with Bact8f (5'-AGA GTT TGA TYM TGG CTC-3') as forward primer (59) and 1492r (5'-N TAC 545 546 CTT GTT ACG ACT-3') as reverse primer (60). Archaeal species were targeted by AR109F 547 (5'-ACK GCT CAG TAA CAC GT-3') as forward (61) and AR915 (3'-GTG CTC CCC CGC CAA TTC CT-3') as reverse primer (62). The PCR mixture contained 0.2 mM of each dNTP, 2 mM 548 MgCL<sub>2</sub>, 20 µg BSA, 1 U of Tag DNA polymerase, and a Tag polymerase buffer with KCI 549 550 (ThermoFisher Scientific). The PCR was performed using an initial denaturation at 95°C for 551 5 min; 30 cycles of 95°C for 30 s, 50°C for 30 s, 72°C for 1.5 min; and a final elongation at 552 72°C for 7 min. For PCRs with archaeal primers, the annealing temperature was set to 55°C. 553 Amplification products were purified by use of the Zymo Research DNA Clean & Concentrator Kit (Zymo Research). 16S rRNA clone libraries were obtained with the TOPO® TA Cloning® 554 Kit (ThermoFisher Scientific). Clones were screened by M13-PCR for inserts of the correct size 555

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- 556 according to the manufacturer's instructions. Resulting PCR products of expected length were
- 557 purified by use of the Zymo Research DNA Clean & Concentrator Kit and sent for sequencing.

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#### **Supporting Tables** 558

#### Table S1. Overview of inocula used to establish initial pyrite-forming enrichment cultures. 559

Enrichment Date		Sampling area	Material	Medium†	Temp. (°C)	рН	
J5*	Sept. 1995	Sewage treatment plant Konstanz, Germany	digested sewage sludge	freshwater	28	7.2	
J2*	Sept. 1995	Sewage treatment plant Konstanz, Germany	digested sewage sludge	freshwater	16	7.2	
J7	Apr. 1993	Rio Tentor (Venice, Italy)	brackish sediment	marine	16	7.2	
J8*	Mar. 1991	Wadden sea sediment, Groningen, The Netherlands	marine sediment	marine	16	7.2	
J9*	Apr. 1993	Fish market channel (Venice, Italy)	brackish sediment	marine	16	7.2	
X1	Sept. 1995	Sewage treatment plant Tübingen-Lustnau, Germany	digested sewage sludge	freshwater	28	7.2	
X2	Sept. 1995	Lake Constance, Güll	freshwater sediment	freshwater	28	7.2	

 $\ensuremath{^+}$  after Widdel and Pfennig (42)  $\ensuremath{^+}$  CH\_4 formation observed

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Table S2. Iron mineral analysis by Mössbauer spectroscopy at a temperature of 5 K in nearly 7-month-old (207 days) culture J5 incubated at various temperatures and in the presence of various inhibitors. Mössbauer parameters were obtained through Voigt based fitting (VBF).  $\delta$  – isomer shift,  $\Delta E_Q$  – quadrupole splitting,  $\varepsilon$  – quadrupole shift, B<sub>hf</sub> – internal magnetic field, R.A. – relative area, X<sup>2</sup> – goodness of fit parameter. The absolute amount of formed FeS<sub>2</sub> was inferred from the relative area of the FeS<sub>2</sub> signal and the maximum amount of 350 µmol that could be produced if all FeS would have been converted to FeS<sub>2</sub>.

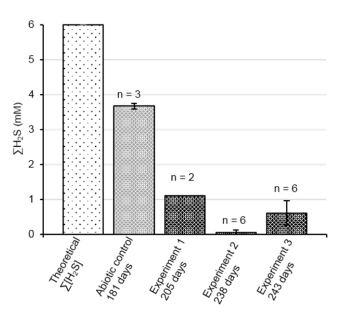
Sample	Phase	ise δ	ΔEq	٤	B <sub>hf</sub>	R. A.	<b>X</b> <sup>2</sup>	FeS <sub>2</sub>
		(mm/s)	(mm/s)	(mm/s)	(T)	%		(µmol)
Abiotic	FeS	0.43		0.15	13.1	64.3	1.67	
control	FeS <sub>x</sub>	0.50	0.13			35.7		
4 °C	FeS	0.47	0.27			37.8	6.01	
	FeS <sub>x</sub>	0.43		0.07	17.2	62.2		
16 °C	FeS <sub>2</sub>	0.41	0.38			18.5	0.78	64.8
	FeS <sub>x</sub>	0.39		0.02	19.2	48.8		
	FeS <sub>x</sub>	0.42		-0.02	16.5	32.7		
28 °C	FeS <sub>2</sub>	0.41	0.50			52.8	0.88	184.8†
Replicate I	FeS <sub>x</sub>	0.32		0.07	15.7	38.9		
	$Fe_3S_4$	0.46		0.05	32.0	8.3		
28 °C	FeS <sub>2</sub>	0.40	0.58			62.5	0.78	218.8§
Replicate II	FeS <sub>x</sub>	0.29		-0.2	16.4	31.0		
	Fe₃S₄	0.60		-0.05	32.0	6.5		
46 °C	FeS <sub>2</sub>	0.42	0.41			39.4	1.67	137.9
	FeS <sub>x</sub>	0.10		-0.09	21.1	50.5		
	FeS <sub>x</sub>	0.40		0.05	15.7	7.8		
	$Fe_3S_4$	0.64		0.03	33.2	2.3		
60 °C	FeS	0.54	0.16			27.1	1.74	
	FeS <sub>x</sub>	0.34		0.05	2.4	21.6		
	FeS <sub>x</sub>	0.55		-0.08	16.6	43.9		
	$Fe_3S_4$	0.63		-0.08	31.9	7.3		
Penicillin + 79% H <sub>2</sub>	FeS	0.51	0.16			31.6	1.12	
( headspace)	FeS <sub>x</sub>	0.38		0.15	5.3	17.4		
	FeS <sub>x</sub>	0.85		0.01	23.0	37.5		
	FeS <sub>x</sub>	0.46		-0.07	15.5	13.4		
Penicillin	FeS	0.46	0.27			26.4	1.15	
	Fe₃S₄	0.68		-0.04	33.1	8.8		
	$Fe_3S_4$	0.52		0.05	31.7	18.8		
	FeS <sub>x</sub>	0.44		0.00	14.9	15.0		
	FeS <sub>x</sub>	0.43		0.02	20.2	31.1		
BES	FeS	0.50	0.22			28.1	0.76	
	$Fe_3S_4$	0.69		-0.15	29.6	23.4		
	Fe <sub>3</sub> S <sub>4</sub>	0.62		0.02	33.1	7.1		
	FeS <sub>x</sub>	0.25		0.15	7.4	20.2		
	FeS <sub>x</sub>	0.44		-0.16	16.6	21.2		

† corresponding amount of formed  $CH_4$ : 44.9 µmol § corresponding amount of formed  $CH_4$ : 67.6 µmol



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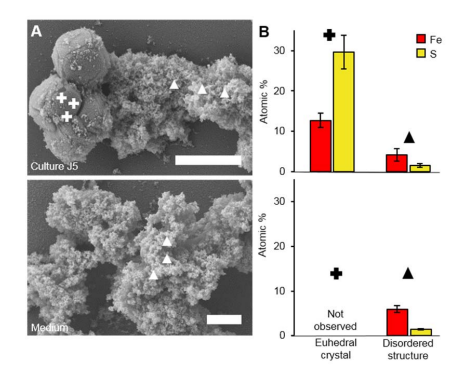
### 574 Supporting Figures



575

576 Figure S1. Total H<sub>2</sub>S as the sum of H<sub>2</sub>S<sub>gaseous</sub>, H<sub>2</sub>S<sub>aqueous</sub>, HS<sup>-</sup>, and S<sup>2-</sup> in the non-inoculated 577 medium as compared to the abiotic control and enrichment culture J5 in various independent 578 incubation experiments. The time of incubation is indicated in days. Biological replicates are 579 indicated as n.

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582 Figure S2. Fe:S ratio of different mineral phases in culture J5 and freshly prepared medium 583 without inoculum. (A) Exemplary scanning electron microscopy images used as guidance to 584 perform energy dispersive X-ray spectroscopy (EDX) point measurements of culture J5 after 585 nearly 10 months of incubation (295 days) and of freshly prepared medium without inoculum. 586 Scale bars represent 2 µm. Symbols in the SEM images indicate EDX point measurements 587 (crosses for crystals, triangles for disordered structure). (B) Atom percent ratio of iron (red) and 588 sulfur (yellow) as derived from EDX point measurements of euhedral crystals resembling pyrite 589 as well as disordered structures resembling the sum of the remaining Fe-S-mineral phase. 590 Measurements were done on eight different sampling areas with three EDX point 591 measurements each.

### Microbial pyrite formation

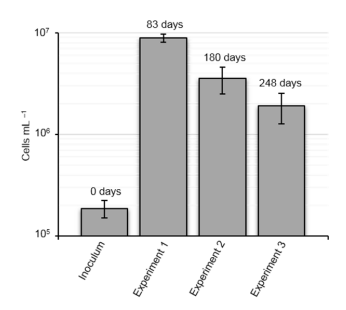
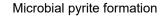


Figure S3. Average cell counts of culture J5 as based on DAPI-stained cells in three independent incubation experiments at 28°C as compared to freshly inoculated medium. The time of incubation is indicated in days. Data was obtained from biological duplicates, each measured in technical triplicates. Standard deviations are given for technical replicates.



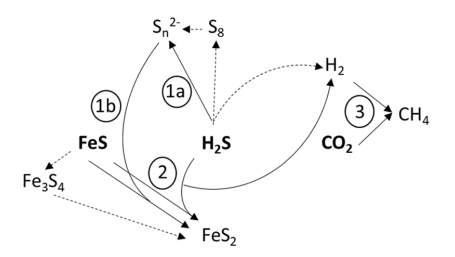


Figure S4. Schematic overview of potential  $FeS_2$  formation pathways in culture J5. The scheme illustrates pyrite formation either by a combination of sulfide oxidation to zero-valent sulfur (1a) coupled to the polysulfide pathway (1b) or by the H<sub>2</sub>S pathway (2). The released reducing equivalents are likely transferred in the form of H<sub>2</sub> to methanogenesis (3) to reduce  $CO_2$  to CH<sub>4</sub>. Dashed lines leading to *cyclo*-octasulfur (S<sub>8</sub>) and greigite (Fe<sub>3</sub>S<sub>4</sub>) represent potential alternative pathways or side reactions, respectively.