Aqueous system-level processes and prokaryote assemblages in the ferruginous and sulfate-rich bottom waters of a post-mining lake

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Abstract. In the aqueous oligotrophic ecosystem of a post-mining lake (Lake Medard, Czechia), reductive Fe(II) dissolution outpaces sulfide generation from microbial sulfate reduction (MSR), and ferruginous conditions occur without quantitative sulfate depletion. An isotopically constrained estimate of the rates of sulfate reduction (SRR) suggests that despite a high genetic potential, this respiration pathway may be limited by the rather low amounts of metabolizable organic carbon. This points to substrate competition exerted by iron and nitrogen respiring prokaryotes. Yet, the microbial succession across the nitrogenous and ferruginous zones of the bottom water column also indicates sustained genetic potential for chemolithotrophic sulfur oxidation. Therefore, our isotopic SRR estimates could be rather portraying high rates of anoxic sulfide oxidation to sulfate, probably accompanied by microbially induced disproportionation of S intermediates. Near and at the anoxic sediment-water interface, vigorous sulfur cycling can be fuelled by ferric and manganic particulate matter and redeposited siderite stocks. Sulfur oxidation and disproportionation then appear to prevent substantial stabilization of iron monosulfides as pyrite but can enable the interstitial precipitation of small proportions of equant microcrystalline gypsum. This latter mineral isotopically fingerprints sulfur oxidation proceeding at near equilibrium with the ambient anoxic waters, whilst authigenic pyrite-sulfur displays a 38 to 27 ‰ isotopic offset from ambient sulfate, suggestive of incomplete MSR and likely reflective also of an open sulfur cycling system. Pyrite-sulfur fractionation decreases with increased reducible reactive iron in the sediment. In the absence of ferruginous coastal zones today, the current water column redox stratification in the post-mining Lake Medard has scientific value for (i) testing emerging hypotheses on how a few interlinked biogeochemical cycles operated in nearshore paleoenvironments during redox transitional states; and (ii) to acquire insight on how similar early diagenetic redox proxy signals developed in sediments affected by analogue transitional states in ancient water columns.
1 Introduction

The biogeochemical reactions governing the distinctive redox structure of modern meromictic lakes have been studied, for the most part, in natural settings featuring high concentrations of dissolved iron and with common sulfate deficiency (Swanner et al., 2020). Improved by insights from laboratory experiments (e.g., Konhauser et al., 2007; Rasmussen et al., 2015; Jiang and Tosca, 2019), geochemical and microbiological analyses made in such lacustrine ferruginous systems have provided us with an empirical framework to interpret modern iron biomineralization mechanisms. Natural meromictic lakes are also relevant to decipher specific chemical and isotopic signatures of ferruginous type sediments that are meaningful for paleoenvironmental interpretations. For example, these settings have significantly informed our Precambrian ocean redox stratification models (Crowe et al., 2008; Walter et al., 2014; Koeksoy et al., 2016; Lambrecht et al., 2018; Swanner et al., 2020).

Ferruginous bottom water columns that also contain elevated dissolved sulfate concentrations are not uncommon in shallow pit lakes (e.g., Denimal et al., 2005; Trettin et al., 2007), and have also been reported from the post-mining Lake Medard in NW Czechia (Petrash et al., 2018; Fig. 1a). This newly formed, lacustrine system with a 60 m depth features oligotrophic and meromictic conditions, and given its recent water filling history it can be considered as a large-scale incubation experiment reflecting an imbalanced transitional state between ferruginous and euxinic aquatic redox states (Scholz, 2018; van de Velde et al., 2021). These states are defined by the presence of water column divalent iron ($\text{Fe}^{2+} > \text{H}_2\text{S}$ and $\text{Fe}^{2+} > \text{NO}_3^-/\text{NO}_2^-$) or dissolved sulfide ($\text{H}_2\text{S}/\text{HS}^-$), respectively.

Here we combined spectroscopic hydrochemical analyses of the sulfate-rich, dysoxic (i.e., 2.0 to 0.2 mg·L$^{-1}$ O$_2$), nitrogenous and ferruginous (suboxic) and ultimately anoxic (< 0.03 mg·L$^{-1}$ O$_2$) bottom water column of Lake Medard. We interpret system-level processes that can be linked to the presence of specific planktonic prokaryote functionalities within its redox stratified water column. Isotope ratios of carbon and oxygen in dissolved inorganic carbon, sulfur and oxygen in dissolved sulfate, and concentration profiles of bioactive ions and volatile fatty acids (VFAs) were measured together with a 16S rRNA amplicon gene sequence profile. To complement our interpretations, in the upper anoxic sediments we also conducted mineralogical analyses and a mineral-calibrated wet chemical speciation study of the reactive Fe and Mn pools. Using these data, we developed a mechanistic model that assesses the potential regulatory roles of prokaryotes over the geochemical gradients detected in the water column, and their influence over iron (Fe) mineral transformations and biogeochemical cycling of sulfur (S), carbon (C), nitrogen (N) and manganese (Mn) across the redoxcline, and near the anoxic sediment-water interface (SWI). The consumption and replenishment of these elements are presented as a set of biogeochemical reactions that differentiate the bottom waters in distinctive niches where a phylogenetically and metabolically diverse microbial community induce an active elemental internal cycling involving reactive minerals in the sediments.

Our observations in this unique artificial lacustrine system are thought relevant since analogue aqueous-level system processes would have also punctually operated in ancient ferruginous coastal settings. Lake Medard could therefore offer valuable information to further understand the roles played by comparable microbial aqueous ecosystems in mineral
authigenesis in the past. In addition, our research furthers understanding of the cryptic S cycle under ferruginous conditions unaccompanied by a quantitative sulfate exhaustion, and for refining our proxy-based palaeoceanographic reconstructions of nearshore redox stratified water columns.

Figure 1. The area now occupied by the post-mining Lake Medard was previously an open cast coal mine near Sokolov, NW Czechia. Upon mine abandonment, the deepest parts of the open cast mine became shallow acidic pit lakes and are now the post-mining lake depocenters. These feature ferruginous and sulfate-rich aqueous conditions. The star marks the central sampling location in a recent lake imagery superimposed on the 2005 mine-pit imagery. The mine-pit had important fluxes of solutes linked to pyrite oxidation in exploited coal seams and their associated pyrite-bearing lithologies. These fluxes may still affect the hydrochemistry of the present-day lacustrine system, i.e., solutes are now sourced from now submerged lithologies (b-c). Imagery dates 5/19/2020 (©CNES/Airbus) and 1/1/2004 (©GEODIS Brno). Historical photographic record by courtesy of The Czech Geological Survey.

2 Study site

Reclamation (flooding) of land occupied by the decommissioned Medard open-cast lignite mine in the Sokolov mining district of Karlovy Vary, northwest Czechia, led to the ca. 4.9 km² (~60 m max. depth) post-mining Lake Medard (Fig. 1a; 50°10’41” N, 12°35’46” E). The lake was filled with waters diverted for reclamation purposes from the nearby Eger River. During closure and abandonment of the former lignite mine, dissolved iron, and sulfate—derived from pyrite oxidation, leached towards shallow pit lakes as mining operation ceased and surficial and groundwaters filled the open-cast mine (Fig. 1b-c). In the mining-impacted brines metastable Fe(III)-oxyhydroxides and -oxyhydroxysulfates precipitated (Murad and
Runoff also affected the hydrochemistry of these shallow water bodies (Fig. 1b-c) by carrying solutes sourced from a Miocene sedimentary unit (Cypris Formation) that overlies the mined coal seam. The detrital material consists of fine-grained minerals (silt to clay in size); derived from a tuffaceous and carbonate-rich lacustrine claystone described in detail by Kříbek et al. (2017). Water-rock interactions in the underlying granitic basement also influenced the hydrochemistry of a Miocene playa lake (Rapantová et al., 2012) that extend across the area now occupied by Lake Medard. In the fractures of the crystalline basement rocks, percolation of waters from that paleolake produced epithermal mineral salt deposits in the form of localized thermadite (Na$_2$SO$_4$) efflorescences (Pačes and Šmejkal, 2004). Percolation and subsurface flow of modern meteoric water caused dissolution of the subsurface salts. Thermadite dissolution could have introduced significant loads of isotopically heavy sulfate into the present-day hydrogeological system (Pačes and Šmejkal, op. cit.). Additional details on the geological framework of the area, and its ongoing influence over the hydrochemistry of the more recently formed, post-mining lacustrine system are provided in Appendix A.

At present, Lake Medard is an oligotrophic water body exhibiting density, temperature and redox stratification. Its bottom water column is sulfate-rich and ferruginous. Both, abiotic and microbially mediated precipitation of poorly crystalline iron minerals—i.e., amorphous ferric hydroxide (Fe(OH)$_3$) and metastable nanocrystalline ferrihydrite (Fe$_2$O$_3$·(H$_2$O)$_n$)—occurs near the redoxcline from where these phases are exported to the SWI (Petrash et al., 2018). Mineral equilibrium reactions at the SWI proceed mostly within the suboxic range of redox potential (Eh) and at a circumneutral to moderately alkaline pH. Stability diagrams showcasing the predicted stability of S and Fe species in the bottom waters of Lake Medard are shown in Fig. B1 (Appendix B).

### 3 Methods

A water quality monitoring and profiling probe was used to measure density, temperature, and redox potential in the stratified bottom water column of Lake Medard prior to sampling. On aliquots of our water samples, we performed (i) MiSeq Illumina 16S rRNA gene amplicon sequencing, (ii) mass spectroscopic determinations of ion and cation concentrations, (iii) dissolved inorganic carbon and methane concentrations, (iv) $\delta^{13}$C analysis of total dissolved inorganic carbon and methane, (v) $\delta^{34}$S and $\delta^{18}$O of dissolved sulfate, and (vi) VFAs concentration analyses. We also sampled the upper anoxic sediment column to a depth of ~8 cm. The mineralogy of these sediments was qualitatively and semi-quantitatively assessed via X-ray diffraction (XRD). The $\delta^{34}$S and $\delta^{18}$O isotope ratios of gypsum (CaSO$_4$·2H$_2$O), $\delta^{13}$C of siderite (FeCO$_3$), and $\delta^{34}$S of pyrite (FeS$_2$) from these sediments were also measured. Scanning electron microscopy aided by electron dispersive spectrometry (SEM-EDS) was used for textural analyses focused on the S- and/or Fe-bearing phases. In addition, a sequential extraction scheme (after Poulton et al., 2004; Goldberg et al., 2012) was conducted to characterize the sedimentary partitioning of reactive Fe and Mn fractions. Details on the methods and protocols applied for sample collection and preparation, instrumentation, eDNA extraction and 16S rRNA gene-based clone libraries preparation are provided in Supplement 1. The
metagenomic data for this study (total 1,799,339 read base pairs, mean lengths ≥ 402 bp) were deposited in the European Nucleotide Archive (ENA) at EMBL-EBI under accession number PRJEB47217 (Accessible from 2022).

4 Results and discussion

4.1 Bottom water column stratification and dissolved oxygen (O$_2$) levels

Figure 2a shows the relevant physicochemical parameters measured in the dysoxic to anoxic bottom waters at the time of sampling (November 2019). Profiling of these parameters is consistent with several previous and subsequent probe monitoring measurements in the meromictic post-mining lake. Conductivity exhibits a steep gradient at ca. 48 m depth that flattens with increasing depth. Temperature increases gradually towards the bottom. The zone in the water column where these gradients concur is referred to as the hypolimnion. Increased conductivities within the hypolimnion of post-mining lakes, such as examined here, could result from the legacy of the former mine drainage and/or from groundwater inflow (e.g., Denimal et al., 2005; Schultze et al., 2010). Salinity was estimated by using the measured conductivity values (after Hambright et al., 1994). It increases three-fold from the hypolimnion downwards (Fig. 2a). This could result from recharge of groundwater carrying high loads of dissolved salts or results from the lack of mixing of the legacy mine-impacted pit lake waters with those comprising the mixolimnion. The temperature gradient results from limited seasonal vertical heat exchange between the bottom density stratified water column and the mixolimnion (Boehrer and Schultze, 2008).

Molecular oxygen (O$_2$) from the mixolimnion cannot be replenished below the density and thermally stratified bottom waters, where deoxygenation very likely proceeds through biotic and abiotic reactions involving Fe-oxidation (e.g., Boehrer et al., 2009). Dissolved O$_2$ levels thus drop rapidly from about 8.1 to ~0.2 mg·L$^{-1}$ within the 48 to 49 m depth-interval of the water column, and the deepest part of the lake is anoxic (Fig. 2a). At this level, the Eh shifts from >100 mV at the lower mixolimnion to negative values down to ≤ -230 mV near the SWI. The suboxic to anoxic zone of the water column is referred to as the monimolimnion, with the sharp redox boundary zone being referred to as the redoxcline (Fig. 2a). The hydrochemically different monimolimnion persists in the deepest depressions of the lakebed throughout the year; although with slight variations in the monitored Eh range that could be accompanied by minor (±1 m) shifts in the vertical position of the redoxcline. In this study, we focus on the central part of the lakebed as it exhibits the broadest Eh range in its bottom water column (Fig. 2a). Details on the eastern and western sampling locations are available in a descriptive study by Petrash et al. (2018). Short-lived changes in redox potential of the bottom water column were recently considered by Umbria-Salinas et al. (2021). These changes have effects on the partitioning of several redox sensitive metals that are bound to reactive iron phases in the upper sediments.
Figure 2. Physicochemical parameters in the bottom dysoxic to anoxic waters of Lake Medard in its central sampling location (a) and concentrations range of discrete VFAs (acetate, formate and oxalate) quantified in the dysoxic (n= 4) and anoxic (n= 3) waters of Lake Medard (b). The arrow in “a” shows the position of the redoxcline at the time of sampling. For VFAs detection limits were better than 60 ppb for lactate and oxalate, and 200 ppb for butyrate, formate, and acetate.

4.2 Dissolved carbon concentrations and δ¹³C isotope values

4.2.1 Dissolved Organic Carbon (DOC)

The DOC concentrations in the bottom waters sampled were 1.05 ± 0.5 mM. VFAs were at nanomolar concentrations that are reflective of the general scarcity of labile organic substrates in the Lake Medard oligotrophic system. A six- to ten-fold increase in concentrations of acetate, oxalate, and formate occurred towards the increasingly saline and O₂-depleted bottom waters. Concentrations of lactate, propionate, and butyrate can be detected at similar nanomolar magnitudes in the
mixolimnion (not shown), but in the bottom waters these VFAs are exhausted and were below our limits of quantification (<LQ) (Fig. 2b).

4.2.2 Total dissolved inorganic carbon

The concentrations of total dissolved inorganic carbon (i.e., $\Sigma$CO$_2$ = H$_2$CO$_3$ + HCO$_3^-$ + CO$_3^{2-}$) ranged from 1.9 to 9.8 mM and increased downwards (Fig 4a). The concentrations of this parameter ($\Sigma$CO$_2$) were positively correlated with alkalinity, which ranged from 1.8 to 2.9 meq·L$^{-1}$ (measured as HCO$_3^-$). The pH in the hypolimnion was ~8.2 and decreased moderately downwards, reaching 7.4 ± 0.2 units near the anoxic SWI. Simultaneous reactions involving dissolution, anoxic reoxidation and (re)precipitation of reactive minerals could be responsible of this moderate decrease in pH (see Soetaert et al., 2007). These reactions are further considered in subsequent sections of this work.

Total dissolved inorganic carbon exhibited lower $\delta^{13}$C values at the anoxic monimolimnion (Table 1, Fig. 3a), and $\Sigma$CO$_2$ were inversely correlated with the $\delta^{13}$C values. The concentrations of dissolved sulfate and $\Sigma$CO$_2$ were inversely correlated too (Fig. 3b-c). From these observations, a contribution of organically derived CO$_2$ is evident, but it could be also argued that in the bottom water column, sulfate reduction has a moderate impact on alkalinity. Although speculative, it is possible that microbial sulfate reduction (MSR) is responsible for lactate depletion in the bottom waters. It could also be a factor contributing to the slight decrease in pH in the monimolimnion (see Gallagher et al., 2014).

The CO$_2$ source flux at the lake floor can be constrained given that the $\delta^{13}$C values in sedimentary carbonates and organic matter are known. A two-component mixing model thus permits estimation of the relative contribution of these carbon sources to the bottom water $\Sigma$CO$_2$ reservoir. We plotted the $\delta^{13}$C signatures of water vs. the inverse of the $\Sigma$CO$_2$ concentrations, corrected by a mixing factor ($\delta^{13}$C = f $\Sigma$CO$_2^{-1}$ + b; after Keeling, 1961; Fig. 3d). The intercept “b” of the linear regression can then be assumed as the isotopic carbon signature of the combined CO$_2$ flux. This approach works, in our case, under the postulate that the $\Sigma$CO$_2$ flux at the lake floor displays an isotope signature resulting from the admixture of CO$_2$ evolved from both remineralization of the sedimentary organic matter ($\delta^{13}$C = -27.9 ± 0.1 ‰, n=6), and (bi)carbonate ions derived from the dissolution of carbonate phases near the SWI and below. A minor contribution of $\Sigma$CO$_2$ evolved from the oxidation of methane diffusing throughout the anoxic sediments to the bottom water column (mean $\delta^{13}$C$_{CH4}$ ≈ -67 ‰; Table 1) might also be possible and was considered. To account for the reactive C of the sedimentary mineral phases, we used the carbonate $\delta^{13}$C mean values in the anoxic sediments (+6.4 ± 0.3 ‰), which is within the range reported for carbonates in lignite-associated lithologies ($\delta^{13}$C range: +1.7 to +13.4 ‰; median = +9.8 ‰; Šmejkal, 1978, 1984). Our sediment’s $\delta^{13}$C mean value thus fingerprints mostly siderite, which is the only carbonate phase detected via XRD. Yet, other relatively more soluble carbonate phases, such as dolomite and calcite, might be present in small proportions at the lake floor because they occur with siderite in the Miocene Cypris Fm. These would account for only ≤ 0.2 wt. % (i.e., the LQ of our semi-quantitative XRD analyses).
Table 1. Measured concentrations and isotopic ratios in the O2 depleted bottom water column of the central sampling location (from 47 to 55 m depth below the surface), Lake Medard.

| Depth (m) | pH (mV) | Eh [mg-L⁻¹] | O₂ [µS cm⁻¹] | Cond. | ΣCO₂[^a] | [CH₄] | [NO₃⁻] | [NH₄⁺] | [Fe²⁺] | PO₄³⁻ | [Mn²⁺] | [SO₄²⁻] |
|----------|--------|-------------|-------------|-------|---------|-------|-------|-------|-------|-------|-------|-------|
|          |        |             |             |       | µM     | µM    | µM    | µM    | µM    | µM    | µM    | µM    |
| 47       | 8.1    | 85.9        | 8.0         | 1394  | n.d.   | 24.2  | 3.4   | <0.07 | 9.1   | 9.6   | 10.9  | 2.4   |
| 48       | 8.1    | 88.4        | 8.0         | 1409  | n.d.   | 24.9  | 5.2   | <0.07 | 9.1   | 9.6   | 10.9  | 2.4   |
| 48.5     | 7.8    | -36.4       | 3.7         | 3143  | n.d.   | 26    | 15.7  | <0.07 | 19.9  | 8.3   | 11.3  | 2.4   |
| 49       | 7.8    | -145.1      | 0.9         | 4871  | n.d.   | 19.8  | 3.4   | <0.07 | 20.1  | 9.6   | 11.5  | 3.9   |
| 50       | 7.7    | -159.9      | 0.1         | 5197  | n.d.   | 17.5  | 68.7  | 22.8  | 30.6  | 12.8  | 12.1  | 3.5   |
| 52       | 7.7    | -185.3      | 0.07        | 6661  | n.d.   | 17.5  | 68.7  | 22.8  | 30.6  | 12.8  | 12.1  | 3.5   |
| 54       | 7.7    | -204.6      | 0.06        | 7440  | n.d.   | 18.3  | 134.5 | 32.7  | 13.7  | 16.8  | 13.3  | 4.0   |
| 55       | 7.7    | -214.8      | 0.03        | 7618  | n.d.   | 17.9  | 127.7 | 28.1  | 28.1  | 11.1  | 16.0  | n.d.  |

[^a]: ΣCO₂ = H₂CO₃ + HCO₃⁻ + CO₂⁻.

Precision of the isotopic values reported—based on repeated measurements of analytical standards (better than; % ± 3σ): (a) 0.05; (b) 0.4; (c) 0.9; (d) 0.1; (e) 0.4

The estimated isotopic C signature of the CO₂ flux at the anoxic sediments is between -3.0 and -4.2‰ (Fig. 3d). This predicted δ¹³C range of values for CO₂ diffused into the overlying monimolimnion is within the range observed in emanations regionally releasing magnetically derived CO₂ (Weinlich et al., 1999; Dupalová et al., 2012). In other words, the variable contribution of CO₂ derived from OM degradation, carbonate mineral dissolution and any plausible methanotrophic activity produces an isotopic signature in the ΣCO₂ of the lake bottom water that matches that of the magmatic-derived CO₂ emissions. The mixing factor in the simple linear mixing model accounts for the proportional contributions of the sources to the ΣCO₂ admixture entering the bottom waters (e.g., Phillips and Gregg, 2001). Accordingly, dissolution of sedimentary carbonates accounts for 70 ± 5% of the dissolved inorganic carbon, with the remaining fraction being the product of organic matter heterotrophy. The contribution of isotopically light CO₂ derived from the oxidation of diffused methane is negligible. Any contribution of CO₂ from the magmatic source cannot be estimated because of the similar isotopic signatures. The implication for environmental/early diagenetic interpretations of this approach is that if siderite is formed in the lake sediments, it appears to display a significant δ¹³C offset from the signature of the ΣCO₂ reservoir of the lake’s floor. We will revisit the siderite origin under Sect. 4.6.3.
4.3 Nitrogen, iron and sulfur species in water column with functional annotations on the planktonic prokaryote community

4.3.1 Nitrogen species transformations and nitrogen-utilizing prokaryotes

Dissolved nitrate (NO$_3^-$) concentrations across the dysoxic hypolimnion are approximately 25 µM and decrease about 28 % towards the anoxic monimolimnion. This decrease is accompanied by a significant increase in ammonium from 16 µM to up to 142 µM (Table 1; Fig. 4a). Similar behaviour of reactive N species were described in other ferruginous water columns (e.g., Michiels et al., 2017; Lambrecht et al., 2018).
The relative abundance of 16S rRNA gene sequences that can be ascribed to N-utilizing planktonic prokaryotes indicates that *Nitrosomonas*-like species (95 to 98 % gen similarity) are in the dysoxic hypolimnion at a low normalized abundance which increases significantly at the redoxcline. Here *Nitrosomonas*-like species may conduct the first and rate-limiting step in nitrification, i.e., NH$_3$ oxidation (Lehtovirta-Morley, 2018). The second nitrification step, nitrite oxidation to NO$_3^-$, could be exerted predominantly by species exhibiting similarity (98% gene sequence) to *Ca. Nitrotoga*. *Ca. Nitrotoga* was sequenced in all our bottom water column samples but exhibited a higher normalized abundance (up to 9 %) at the redoxcline (Fig. 5a).

Among the relatively abundant, NH$_3$-oxidizing microbes sequenced is an archaeon related to *Nitrosarchaeum koreense* (97-98 % gen similarity). This archaeon appears to thrive better in the ferruginous waters below the redoxcline (Fig. 5a, also Supplement 2). Its distribution across the redox gradient is at odds with the fact that *N. koreense* has been previously suggested to be an aerobe (Jung et al., 2018). Our observation could make the case for niche differentiation linked to high loads of dissolved metal concentrations conferring a competitive advantage to the *N. koreense*-like microorganism (e.g.,

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**Figure 4.** Measured dissolved concentrations of nitrate and ammonia (a), manganous manganese and ferrous iron (b), and (c) sulfate.
Gwak et al., 2019). Alternatively, the archaeon sequenced predominantly in the suboxic waters possesses tolerance to anoxia (see Mußmann et al., 2011). The abundance peak in *Azospira*-like microorganism (95 % similarity) coincides with the peak of relative abundance of members of the Gallionellacea family at 49 to 50 m depth (Fig. 5a, Supplement 2). Like *Gallionella, Azospira* also possess dissimilatory N and Fe-based metabolisms capable of yielding dinitrogen (N$_2$). N$_2$ production can thus be thought to account for the apparent nitrogen loss observed when the dissolved reactive NH$_4^+$ and NO$_3^-$ levels are compared across their counter gradients (Table 1; Fig. 5a).

Comparison of the gradients of abundances of the N-utilizing prokaryotes with the gradients of other bioactive species is presented in Fig. 4. This suggests that while metabolizing nitrogen the prokaryote community also impacts the cycles of Fe and S (e.g., Jewell et al., 2016, 2017; Starke et al., 2017). These cycles in the aqueous system under consideration are interlinked throughout microbial mediation in the following Rcs (1-3):

\[
\begin{align*}
10\text{Fe}^{2+} + 2\text{NO}_3^- + 24\text{H}_2\text{O} & \rightarrow 10\text{Fe(OH)}_3^- + \text{N}_2 + 18\text{H}^+ \\
\text{NO}_3^- + 8\text{Fe}^{2+} + 21\text{H}_2\text{O} & \rightarrow \text{NH}_4^+ + 8\text{Fe(OH)}_3^- + 14\text{H}^+ \\
5\text{HS}^- + 8\text{NO}_3^- & \rightarrow 5\text{SO}_4^{2-} + 4\text{N}_2 + 3\text{OH}^- + \text{H}_2\text{O}
\end{align*}
\]

Reaction 1 proceeds mixotrophically, usually requiring a favourable organic co-substrate, whereas reactions 2 and 3 likely proceed under the influence of chemolithotrophic Fe(II) and/or S oxidizing nitrate reducers. Due to energetic considerations, these microorganisms are known for having metabolic advantages under ferruginous conditions over solely denitrifying organisms (see Robertson and Thamdrup, 2017). Reaction 3 is known to proceed at rather low sulfide levels (Brunet and Garcia-Gil, 1996; Barnard and Russo, 2009), such as those characterizing the monimolimnion of our study site (≤ 0.3 µM).

In the following section, to further investigate details on the microbial ecology of the bottom ferruginous waters of Lake Medard, we consider the concentration profiles of dissolved Fe and Mn along the redoxcline. Concentrations of these dissolved metals are operationally defined as the combined ionic and colloidal fractions that passed the 0.22 µm cut-off of membrane filters. By co-evaluating the dissolved Fe and Mn concentration trends, we pursue further insight on the mechanism procuring and/or consuming these metals in the stratified water column (Davidson, 1993). A 16S rRNA gene abundance profile of known iron-utilizing prokaryotes also permitted inferences on what members of the microbial community could be exerting a direct dissimilatory (catabolic), or indirect (via electron transfer) control over the concentration trends of these metals across the redox gradient.
4.3.2 Dissolved manganese and iron and the iron-utilizing prokaryotes

Dissolved manganese concentrations ([Mn]) peaked at about 50 m-depth (Table 1). Below this depth, [Mn] showed a steady decrease (Fig. 4b). This trend indicates that in the water column the 50-m depth acts as a point source of Mn(II) (Davison, 1993). Fe is also present at a similar concentration magnitude at this depth (Fig. 4b, Table 1), and it can readily act as a reductant of most particulate Mn(IV) settling down from the mixolimnion (Lovley and Phillips, 1988):  

$$2\text{Fe}^{2+} + \text{MnO}_2 + 1.5\text{H}_2\text{O} \rightarrow \text{Fe}_2\text{O}_3 \cdot 0.5\text{H}_2\text{O} + \text{Mn}^{2+} + 2\text{H}^+$$  

(3)

Accordingly, a significant fraction of the Fe(II) diffusing upwards from the monimolimnion is re-oxidized or cycled back to Fe(III) within the peak zone of Mn(IV)-reduction at 50 m-depth (Fig. 4b). Mn(II) yielded during iron oxidation can then be transported both upwards and downwards away from the 50 m-depth source point by eddy diffusion (Fig. 4b; Davidson, 1993). The internal bottom water column cycling of iron also reflects on the concentration gradient of dissolved phosphate (Table 1). Solubilization of this oxyanion in the bottom water column is regulated by its complexation with particulate Fe(III) sinks that are in turn being reductively dissolved in the monimolimnion and SWI. Dissolved phosphate is re-complexed back onto nanocelluline and amorphous ferrihydrite-like phases precipitating at the redoxcline.

Contrary to Mn, dissolved [Fe] increased steadily downwards, and its global maximum is reached at about 54 m-depth in the monimolimnion (Table 1). Immediately below this depth, [Fe] decreases by about 14%. This decrease can be consistently
observed in other anoxic zones of the lake (Petrash et al., 2018), and hints to Fe(II) and reduced S co-precipitation as metastable acid volatile monosulfide (FeS; e.g., mackinawite). The dissimilar distribution of divalent Fe and Mn in the bottom water column (Fig 4b) reflects reductive dissolution being much more effective for the sinking manganic particulate than for ferric particulate matter.

Our planktonic prokaryote analysis shows that above the redoxcline the diversity of known iron-respiring prokaryotes was low and dominated by species closely related to the β-Proteobacterium *Rhodoferax* (99-100 % gene similarity) (Fig. 5b, Supplement 2). Other sequences that can be functionally affiliated to Fe(III)-reduction in the dysoxic hypolimnion include a bacterium with some similarity to *Aciditerrimonas* (91% similarity), and also operational taxonomic units (OTUs) exhibiting between 92 and 100% gen similarity to unclassified *Pseudomonas* spp. (Fig. 5b). Bioutilization of manganese by *Pseudomonas* species—both in oxidation and reduction reactions—has been reported (e.g., Tebo et al., 2005; Geszvain et al., 2011; Lovley, 2013; Wright et al., 2018).

As previously mentioned, we detected a sharp increase in both diversity and abundance of microaerophilic Fe(II)-oxidizing *Gallionella* species at the redoxcline and immediately below it. It accounted for up to ~24% of the total normalized gene reads (Fig 5b). The increase in relative abundance of *Gallionella* spp. coincides with an increase in sequences related to *Sideroxydans* spp. (Fig. 5b). These latter microaerophiles can also use Fe(II) as an energy source for chemolithotrophic growth with CO₂ as the sole carbon source (Emerson and Moyer, 1997). Other different physiological groups of Fe(II)-oxidizing microorganisms sequenced above and near-redoxcline samples included anoxicogenic phototrophic and nitrate-reducing species (*Magnetspirillum* and *Ferrigenium*; Fig. 5b, Supplement 2), and *Azospira*-like species. Prokaryotes that can adapt their metabolic strategies to the less pronounced geochemical gradients prevailing at the monimolimnion become predominant below the redoxcline. Among them is a bacterium distantly related (89% identity in 399 bp) to *Candidatus* Magnetobacterium (after Lin et al., 2014), which relative abundance substantially increases at the 50-m depth (Fig 5b). At this level, our gene sequence reads also included an OTU distantly related (78% gene similarity in 419 bp) to *Deferrisoma*. This genus has been designated as a mixotroph also functionally affiliated to dissimilatory Fe(III) reduction (e.g., Pérez-Rodríguez et al., 2016), as well as an OTU closely related to *Georgfuchsia toluolica*. *G. toluolica* is a strictly anaerobic β-Proteobacterium capable of degrading aromatic compounds with either Fe(III) or NO₃⁻ as electron acceptors (Weelink et al., 2009).

Towards the SWI, important members of the Fe-respiring community are those from the family Geobacteracea, which can use insoluble Fe(III) and/or Mn(IV) as electron acceptors, and acetate, formate, alcohols, aromatics, and dihydrogen (H₂) as electron donors (Weber et al., 2006; Lovley and Holmes, 2021). The abundance of *Geobacter* species peaked around the maximum of Fe(III) reduction within the monimolimnion. Here, acetate availability is also relatively high (Fig. 2b). The relative proportion of *Geobacter* spp. increases in parallel with that of their phylogenetically associated *Pelobacter propionicus*, which is a fermentative acetogen that can only indirectly mediate Fe(III) reduction. A possible ecological interaction between *P. propionicus* and *Geobacter* species at the interface of redox boundaries in sedimentary environments has already been reported by Holmes et al. (2007) and Butler et al. (2009).
4.3.3 Dissolved sulfate and the sulfur-utilizing prokaryotes

The dissolved sulfate concentration ([SO₄²⁻]) changes significantly at the redoxcline, where it increases from 6.0 to 16.8 mM (Fig. 4c). At the lower monimolimnion, a decrease in [SO₄²⁻] coincides with a decrease of [Fe(II)] (Table 1, Fig. 4b-c). We detected an increase in both diversity and relative abundance of sulfate reducers in the lower monimolimnion (Fig. 5c). Their by-product sulfide, however, does not accumulate in the ambient waters ([H₂S + HS⁻] ≤ 0.30 μM). The lack of substantial dissolved sulfide towards the SWI and the similar hydrochemical responses of both Fe(II) and [SO₄²⁻] could be considered circumstantial evidence for FeS precipitation. Additional insight on this and other mechanisms of sulfate turnover operating in the water column was sought by evaluating the distribution of S-utilizing prokaryotes.

Our 16S rRNA gene analyses (Fig. 5c; also Supplement 2) revealed a rather low diversity of sulfur-respiring bacteria at the dysoxic hypolimnion. Here OTU assignments show mostly a few uncultured members of the newly proposed order Desulfobulbales of the phylum Desulfovibrateria (previously δ-Proteobacteria, Waite et al., 2020; Ward et al., 2021))(Fig. 5c). Some species within Desulfobulbales require intermediate S or thiosulfate for heterotrophic growth but can also gain energy from pyruvate fermentation (Flores et al., 2012). Desulfobulbus spp. can perform dissimilatory sulfate reduction via the incomplete oxidation of lactate, but *D. propionicus* is known for efficiently conducting disproportionation of elemental sulfur (Lovley and Phillips, 1994). Pyruvate, as lactate, was found below our detection limits across the bottom water column where sequences distantly related to *D. propionicus* (91% similarity in 428bp) appeared to be particularly abundant (Fig. 5c; Supplement 2). Probably important for the microbial sulfur cycling at this level of the water column is also a γ-Proteobacterium from the order Chromatiales that has 92% gene identity in 424 bp to Thiobacillus paradoxus (Fig. 5c). *T. paradoxus* is a chemolithoautotrophic sulfur-oxidizing bacterium that can use both reduced and intermediate S compounds for C fixation (Berben et al., 2015).

There are gene sequences that could be confidently ascribed to the facultative S-utilizing autotroph *Sulfuritalea hydrogenivorans* (3 OTUs with ≥ 97 identity in 424 bp) at the redoxcline. The abundance of *S. hydrogenivorans* increases in parallel to a decrease in the *T. paradoxus*-like bacterium, which suggests that the latter may be at a disadvantage and limited by organic C fixation under the specific hydrochemical conditions prevailing at the redoxcline. Such conditions may include, for instance, an abundance of aqueous intermediate S species. Under such conditions, *S. hydrogenivorans* can outcompete the *T. paradoxus*-like bacterium by oxidizing, under denitrifying conditions, either thiosulfate, S³⁻ and/or H₂ for C fixation (Kojima and Fukui, 2011; Kojima et al., 2014). Interestingly, at the redoxcline, the relative abundance of the species distantly related to fully sequenced Desulfobulbasles also increases to ~1.7% (Fig. 5c).

Below the redoxcline, our genomic data revealed successional development of a more diverse sulfur-respiring bacterial population (Fig. 5c). This was dominated by many relatively rare taxa and a few abundant lineages (Supplement 2), and with a punctuated dominance of species distantly related to *Desulfobacca acetoxidans* (90% identity in 432 bp). *D. acetoxidans* oxidizes acetate using either sulfate, sulfite (SO₃²⁻) or thiosulfate (S₂O₃²⁻) as electron acceptors, but not S⁰ (Oude Elferink et al., 1999). The *D. acetoxidans*-like prokaryote first appeared at 49 m depth but became dominant towards the SWI, together
with Desulfomonile-related species (96% identity in 432 bp). Desulfomonile-related species could be also responsible for the previously noticed pyruvate depletion in the bottom waters, but here they may be also thriving chemolithoautotrophically with either SO$_4^{2-}$ or S$_2$O$_3^{2-}$ as terminal electron acceptors (DeWeerd et al., 1990; Sun et al., 2001). Other prokaryotes probably gain energy out of intermediate S disproportionation in the anoxic monimolimnion. These may include uncultured species distantly related to Desulfatibacillum and Dissulfurirhabdus (2 OTUs with 87% identity in 428 bp). The prevalence of Sulfitobacter across the aqueous redox gradient and into the monimolimnion (Fig. 5c) points to a sustained genetic potential for chemolithotrophic sulfur oxidation in the entire bottom water column.

4.4 δ$^{34}$S and δ$^{18}$O isotope values of dissolved sulfate

4.4.1 Isotopes evidence for disproportionation in the monimolimnion

Water column δ$^{18}$O$_{SO_4}$ values ranged from +2.0 to +4.0 ‰, with corresponding δ$^{34}$S$_{SO_4}$ values ranging between +10.9 and +13.4 ‰ (Table 1, Fig. 6a-b). The depth profiles of these isotopes in the water column reveal that dissolved sulfate in the anoxic monimolimnion is heavier and enriched in $^{18}$O (Fig. 6a-b) relative to the dysoxic waters. Despite the moderate decrease in [SO$_4^{2-}$] towards the SWI (Fig. 4c) no significant sulfur isotope fractionation was registered. The δ$^{34}$S$_{SO_4}$ values (Table 1) display only a weak correlation with [SO$_4^{2-}$].

The ambient bottom waters had a narrow δ$^{18}$O$_{H_2O}$ signature of -6.4 ± 0.3‰ (V-SMOW). This is consistent with ongoing meteoric water-rock interactions and rather limited evaporation effects (cf. Noseck et al., 2004; Pačes and Šmejkal, 2004; Dupalová et al., 2012). By applying the expression first proposed by Taylor et al. (1984) to relate the δ$^{18}$O values of dissolved SO$_4^{2-}$ and those of ambient waters, we deduced that the oxygen isotope effect (δ$^{18}$O$_{SO_4}$-amb. wat.) in our bottom waters ranged between +9.3 and +10.7 ‰. This range was calculated under the assumption that equilibrium of oxygen isotope exchange between cell-internal sulfur compounds and ambient water dominates over kinetic oxygen isotope fractionation (Fritz et al., 1989; Brunner et al., 2005). The estimated δ$^{18}$O$_{SO_4}$-amb. wat. is within the range experimentally derived by Brunner et al. (2005) while using similarly $^{18}$O-depleted ambient waters. It is also within the range observed in studies of S disproportionation reactions generally proceeding under anoxic conditions. Yet it is lower than δ$^{18}$O$_{SO_4}$-amb. wat. values reported by Bottrell and Newton (2006) in biotic experiments with excess reactive Fe(III) species—i.e., +16.1 to +17.5 ‰. Therefore, our δ$^{18}$O$_{SO_4}$-amb. wat. can result from the superimposition of the isotope signals of sulfate reduction, sulfide re-oxidation and intermediate sulfur disproportionation. It follows that the sulfur disproportionation in the bottom waters of Lake Medard most likely results from multiple biologically mediated reactions involving not only reactive iron, but also reducible Mn stocks in the sediments (Böttcher, Thamdrup and Vennemann, 2001). As further discussed below, the anoxic sediments contain a low—i.e., compared with Fe(III)-counterparts—yet still measurable abundance of Mn(IV)-oxyhydroxides (Table 2).
A microbially mediated/induced sulfur disproportionation mechanism that considers reactive iron forms present in the sediments, also involves Mn(IV,III) reduction, and is consistent with formation of FeS in the monimolimnion can be described by Rs. (4)-(6):

\[
3S^0 + 2\text{FeOOH} \rightarrow \text{SO}_4^{2-} + 2\text{FeS} + 2\text{H}^+ \tag{4}
\]

\[
4S^0 + 4\text{H}_2\text{O} + 3\text{FeCO}_3 \rightarrow \text{SO}_4^{2-} + 3\text{FeS} + 2\text{H}^+ + 3\text{H}_2\text{CO}_3 \tag{5}
\]

\[
3S^0 + \text{Mn}_3\text{O}_4 + 2\text{H}^+ \rightarrow \text{SO}_4^{2-} + 2\text{H}_2\text{S}^- + 3\text{Mn}^{2+} \tag{6}
\]

Although not shown in the rather simplified reaction set listed above, S\(^0\) may well be a different intermediate sulfur species such as thiosulfate and/or sulfite (e.g., Holmkvist et al., 2011). The intracellular isotope exchange of sulfite with anoxic ambient waters has been proven to produce an oxidized SO\(_4^{2-}\) product that is enriched in \(^{18}\)O relative to precursory thiosulfate and/or sulfite. This enrichment displays only a minor change, if any, in its corresponding S isotope composition (e.g., Böttcher et al., 2005; Johnston et al., 2014; Bertran et al., 2020; see Table 1).

### 4.4.2 Isotopic insight on intermediate sulfur oxidation

Figure 6c shows a cross-plot of the \(\delta^{34}\text{SO}_4\) vs. \(\delta^{18}\text{O}_4\) values along the redoxcline as well as those of all the possible geogenic sources of sulfate entering the lake system (see also Appendix B: Fig. B2). Analysis shows that the \(\delta^{34}\text{SO}_4\) values of the redox stratified Lake Medard fingerprint a mixed geogenic-sulfate source. Fig. 6d offers further detail and linear regressions of the covariation in the \(\delta^{34}\text{SO}_4\) vs. \(\delta^{18}\text{O}_4\) cross-plot. The slopes of such linear regressions can be used to roughly estimate sulfate reduction rates (SRR; after Böttcher et al., 2001; Brunner et al., 2005, among others). For assessing our SRR, it is reasonable to assume that the initial S and O isotope composition linked to dissolved sulfate was within the range of the modern nearby acidic drainage (i.e., +2.9 ± 0.1 ‰ for \(\delta^{34}\text{SO}_4\) and 0.0 ± 0.5 ‰ for \(\delta^{18}\text{O}_4\)), and similar to the initial composition of sulfate in the pit-lake prior to reclamation/flooding (Fig. 1c-d; also Fig. B2 in Appendix B). The residual isotope composition would then be that of dissolved sulfate in the bottom anoxic waters.

In agreement with the lack of accumulation of sulfide in the monimolimnion, our SRR estimation is consistent with slow gross but not net SO\(_4^{2-}\) reduction (see Böttcher et al., 2004). The SRR estimate is apparently slower at the monimolimnion (i.e., higher slope) than in the hypolimnion. This is at odds, however, with the higher abundance and diversity of sulfate reducers that we detected near the SWI (Fig. 5c). The decrease in dissolved sulfate concentration does not lower the slope of the linear regression. It means that the sulfur isotope ratio of dissolved sulfate evolves slower relative to corresponding change in oxygen isotope ratio (Antler et al., 2013). This result is likely due to sulfate regeneration through microbial sulfide oxidation; with oxygen isotope exchange with ambient water occurring via an intracellular oxidation step of intermediate sulfur (Böttcher et al., 2005; Bertran et al., 2020). Under the low organic substrate availability characterizing the bottom waters examined here (Fig. 2b), sulfate reducers capable of disproportionation (e.g., bacteria from the order Desulfobulbales)
can maintain intracellular concentrations of sulfite. This manifested geochemically as the rapid change in water column \( \delta^{18}O_{SO_4} \) (Böttcher et al., 2005; Antler et al., 2013).

4.5 Insights from solid phase analyses

4.5.1 Semi-quantitative X-ray diffraction

XRD analyses of the anoxic sediments show that most detrital minerals were sourced from the Miocene claystone lithology (Appendix A). These detrital phases include kaolinite, quartz, K-feldspar, the TiO\(_2\) polymorphs rutile and anatase, and analcime (NaAlSi\(_2\)O\(_6\)·H\(_2\)O). Minor constituents of the anoxic lake sediments that can also be quantified include gypsum, siderite, and pyrite. Gypsum and siderite are in similar abundances in the upper anoxic sediments (~3 to 4 wt. %), whereas
pyrite accounts for a maximum of 0.5 wt. % of their total mineralogy (Fig. 7a). Given that the diffraction peaks of major and minor mineral sediment constituents mask those of Fe(III)- and Mn(IV)-oxyhydroxides, the abundances of these reactive phases were determined through a sequential extraction scheme that also targets Fe(II)- and Mn(II)-bearing carbonates.

4.5.2 Sequential extractions of reactive iron

Figure 7b displays the relative concentrations of highly reactive Fe-bearing species (Fe$_{HR}$) in the upper anoxic sediment pile. The Fe$_{HR}$ sediment pool is defined as that capable of reacting (upon reductive dissolution) with dissolved sulfide to precipitate metastable FeS, which can later be stabilized to pyrite (Canfield and Berner, 1987; Canfield, 1989). We also report here Fe(II) bound to the pyrite fraction (Fe$_{py}$), and the total iron (Fe$_{T}$) in the sediments (after Poulton and Canfield, 2005).

Our Fe$_{HR}$ was dominated by poorly crystalline phases (Feh), such as ferrihydrite and/or lepidocrocite ($\gamma$-FeOOH). These Fe$_{HR}$ mineral fractions were followed in abundance by that of Fe(II)-bearing carbonates (FeC) (Fig. 7b, Table 2). Absolute Fe(III) concentrations ascribed to Feh phases increase slightly towards the bottom of our 8 cm depth core but their abundance, relative to total iron, decreases downwards. The extraction step for Feh also extracts Fe(II) bound to FeS (Kostka et al., 1995; Scholz and Neumann, 2007). These metastable phases yielded $\leq$ 0.04 wt. % according to our acid volatile sulfur (AVS) extraction. However, during sampling of the sediments possible rapid oxidation of AVS particles makes it challenging to assess their actual abundance and mineralogy (Schoonen, 2004). It thus appears then that the Feh abundance at the top of the sediments (Fig. 7b) is mostly comprised of poorly crystalline oxyhydroxide.

A significant increase in the FeC is observed with increasing depth (Table 2, Fig. 7b). This may be indicative of partial dissolution of some Fe(II)-bearing carbonates at the SWI, or the result of soluble Fe(II) being precipitated into reactive Fe-carbonates deeper through the sedimentary pile. To clarify this matter, we discuss the petrographic features and C isotope values of these sedimentary siderites in Sect. 4.6.3.

Table 2. Partitioning of reactive iron and manganese species in the lacustrine sediments (0-8 cm depth).

| Depth (cm) | Fe$_{HR}$ | Fe$_{T}$ | Fe$_{py}$ | FeC | Mn$_{HR}$ | Mn$_{T}$ |
|-----------|-----------|----------|-----------|-----|-----------|---------|
| 0-2       | 14.7      | 283.2    | 302.8     | 1024.4 | 1.0       | 3.6     |
| 2-4       | $\pm2.4$  | $\pm45.3$ | $\pm48.3$ | $\pm4.7$ | $\pm3.6$ | $\pm0.1$ |
| 4-6       | $\pm3.2$  | $\pm47.2$ | $\pm49.4$ | $\pm5.4$ | $\pm2.4$ | $\pm0.3$ |
| 6-8       | $\pm4.6$  | $\pm58.5$ | $\pm42.2$ | $\pm21.5$ | $\pm2.9$ | $\pm0.2$ |

* Sediment density is estimated in 2.71 g·L$^{-1}$ with a porosity of 40%; error of the measurement (n=4) is $\pm16\%$ [Fe] and $\pm8\%$ [Mn]; in μmol·cm$^{-3}$. 

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The iron extracted from crystalline Fe(III)-bearing phases (such as goethite) increased from 2.7 ± 0.4 % in the first 6 cm to up to 17.8 % of the FeT at the 6 to 8 cm interval (Table 2). Fe concentrations bound to pyrite (Table 2, Fig. 7b) constituted up to ~21 % of the FeT in the upper sediments (i.e., ~0.8 wt. %), and showed a general downwards decreasing trend contrasting with that of crystalline Fe(III)-bearing phases. From these observations, the 0 to 6 cm depth interval is considered as recent anoxic lake deposition, whilst below 6 cm are sediments that were deposited in the shallow pit lake now undergoing alteration under the redox dynamics of the present-day lacustrine system.

The FePy/FeHR ratio in the 8 cm sediment profile accounts for the extent to which the Fe pool was pyritized. The ratio is < 0.35 and it decreases downwards (Table 2). When considering that the corresponding FeHR/FeT ratios were consistently ≥ 0.71, the results from our sequential extraction scheme applied to iron are consistent with a persistent ferruginous but not euxinic redox state of the now anoxic sediments (Poulton and Canfield, 2011).

Variability of FePy/FeHR and FeHR/FeT with depth of the sediments reflects the redox dynamics after flooding and establishment of a persistently anoxic monimolimnion.

Figure 7. Mineralogical analyses of the upper sediment (0 to 8 cm depth), the semi-quantitative XRD data (a) shows that the sediments are dominated by aluminosilicates and contain pyrite, gypsum, and siderite. Results from sequential extraction of iron (b) and manganese (c) portray changes in partitioning of these metals in reactive oxyhydroxide, carbonate and sulfide solid phases with increasing sediment depth. SEM-EDX of rhombohedral siderite in the 0-4 (d) and 4-8 cm sediment depth intervals (e). This carbonate mineral displayed corroded surfaces near the sediment-water interface. The texture of microcrystalline equant gypsum (f) and truncated octahedral microcrystalline pyrite (g) are also shown (see text for details).

From combining results from FeHR partitioning in the sediments (Table 2) and the dissolved Mn(II) and Fe(II) concentration trends (Fig. 4b), we can now strengthen an earlier deduction that Fe(II) sourced from reductive
dissolution processes in the upper sediments diffuses upwards, where it rapidly reacts with residual \( \text{O}_2 \) in the vicinity of the redoxcline to form metastable Fe(III)-bearing particulate phases. Most of the iron in such amorphous to nanocrystalline ferrihydrite-like aggregates are deposited on the lake’s anoxic floor. From the anoxic floor they are resolubilized back into the monimolimnion, yet a significant fraction can stabilize upon burial as goethite (\( \alpha \)-FeOOH) or be bound to the surfaces of reactive carbonates. Another fraction is pyritized through reactions involving elemental sulfur and/or polysulfide near the sediment-water interface (Fig. 7b) (see Shoonen, 2004 for details). Indeed, we observed that in the upper sediment the partitioning of the reactive iron into these minerals can be swiftly altered by slight (\( \pm \) 80 mV), short-lived variations in the redox potential of the overlying bottom water column. Variations in the relative proportions of reactive iron minerals also control the distribution of siderophile redox sensitive elements in the sediment pile (Umbria Salinas et al., 2021).

### 4.5.3 Sequential extractions of reactive Mn-bearing phases

Results from our extraction scheme applied to Mn (i.e., after Slomp et al., 1997; Van Der Zee and Van Raaphorst, 2004) show that the Mn\(_{\text{HR}}\) pool in the anoxic sediment was dominated by Mn(II)-bearing carbonates (MnC) (Fig. 7c, Table 2). The carbonates were relatively more abundant at the SWI but in contrast to FeC, showed no clearly defined concentration trend in the upper sediments (Table 2, Fig. 7b-c). A declining trend downwards is clear for the proportions of easily reducible Mn(IV)-bound to poorly crystalline phases, such as \( \delta \)-MnO\(_2\). These were extracted by diluted HCl (Fig. 7c, Table 2) (Slomp et al., 1997). Reducible Mn associated with more crystalline oxyhydroxide forms are extracted by dithionite (Canfield et al., 1993), but concentrations of this fraction might be sourced from crystalline Fe(III)-oxyhydroxides that can either sorb Mn(II) or structurally incorporate Mn(III) (Namgung et al., 2020). Irrespective of its source, the highly crystalline Mn-bearing fraction in our sediment comprises \( \leq 0.2 \) wt. % of Mn\(_T\) (Table 2). The concentrations of Mn(II) bound to sulfides accounted for \( \leq 0.03 \) wt. % of the total Mn extracted (Fig. 7c, Table 2). From the analyses of the partitioning of reactive Mn species, we can thus confirm that under the anoxic conditions currently prevailing in the bottom waters and SWI of Lake Medard, a minor, yet still important fraction of reducible Mn\(_{\text{HR}}\) can be exported from the water column, and can participate, together with the reactive forms of iron, into the internal cycle of S (e.g., reaction 6).

### 4.6 Insights from siderite, gypsum, and pyrite analyses

**Siderite** accounts for up to 3.5 wt. % of the total mineralogy of the anoxic lacustrine sediment where it occurs as dispersed fine crystalline rhombohedra (Fig. 7d-e). Siderite displays corroded surfaces towards the SWI. This textural feature cannot be observed in crystals at the 4 to 8 cm depth interval. This is consistent with results from the sequential iron extraction scheme (see above) indicative of Fe-carbonate likely undergoing recrystallization and/or growth in the deeper part of the examined sediment pile but partial dissolution towards the SWI in spite of its low supersaturation in the monimolimnion waters (\( \Omega_{\text{sid}} = \log \text{IAP} \cdot (\log K_{\text{Sp}})^{-1} = 1.8 \); Supplement 3).

The siderite is enriched in \( ^{13}\text{C} \) by around \(+9\%o\) (mean \( ^{13}\text{C} \) value of siderite is \(+6.4 \pm 0.3 \%o\) relative to \( \Sigma \text{CO}_2 \) of the bottom water column (Table 1). The mean \( ^{13}\text{C} \) value is, however, within the range of \( ^{13}\text{C} \) isotope values reported by Šmejkal
(1978) for carbonates of the Cypris claystone. Also, the mean \( \delta^{18}O \) (SMOW) values (+25.7 ± 1.7 ‰) of siderite are within the range observed in Miocene carbonates which, in addition to siderite, include also dolomite and calcite (Šmekkal, 1978, 1984). From combining the average isotopic values and textural features of siderite in our anoxic sediments, the mineral can then be considered a seeded (detrital) phase also sourced from the Cypris claystone. Siderite seeds were probably redeposited in the floor of the post-mining lake, together with aluminosilicates and other major and minor phases, during the lake’s flooding stage (2008-2015) or thereafter.

Gypsum has a relative abundance of ca. 3 wt. %. It displays a microcrystalline \{010\}-dominated platy shape (Fig. 7f). This is an equilibrium morphology corresponding to a rather low supersaturation (e.g., Simon et al., 1965; van der Voort and Hartman, 1991; Massaro et al., 2010; Rodríguez-Ruiz et al., 2011). This soluble mineral is not thermodynamically stable by the aqueous-mineral equilibrium modeling of the monimolimnion water (i.e., \( \Omega_{gy} \) = -0.5; Supplement 3). However, at a low supersaturation state (0 < \( \Omega_{gy} \) < 1), gypsum would be formed within the upper sediment pore spaces where Ca\(^{2+}\) ion concentrations are increased locally, for instance by carbonate dissolution.

Gypsum precipitation under low saturation states can probably occur as the result of short-lived, climatically constrained changes in the precipitation-dissolution environment of carbonates in the upper sediment pile (see Umbria-Salinas and Valero et al., 2021). The isotope values of the sulfate moiety in the authigenic gypsum (\( \delta^{34}S_{gy} \) and \( \delta^{18}O_{gy} \)) provide further insight on the significance of this phase within the internal sulfur cycle and early diagenetic context of the system under consideration. The \( \delta^{34}S_{gy} \) isotope values ranged from -13.9 and -9.6 ‰ (V-CDT). Accordingly, gypsum shows \( ^{34}S \)-depletion of -17.8 to -11.6 ‰ relative to dissolved SO\(_4^{2-}\) in the ambient anoxic waters (Table 1). The \( \delta^{18}O_{gy} \) values range from +5.1 to +6.3 ‰ (V-SMOW). In consequence, the sulfate in gypsum is enriched by +1.4 to +2.6 ‰ as \( ^{18}O \) compared with the mean \( \delta^{18}O_{SO4} \) of the monimolimnion (Table 1). This magnitude of isotope \( ^{18}O \) enrichment of gypsum-sulfate appears consistent with the range observed when sulfate is derived from pyrite that is oxidized by ferric iron in aqueous anaerobic experiments (e.g., Taylor et al., 1984b; Toran and Harris, 1989; Balci et al., 2007).

A net O\(_2\) neutral reaction that also accounts for (i) significant monosulfide oxidation, (ii) the localized presence of corroded siderite in the upper sediment, (iii) involves chemolithoautotrophic fixation of CO\(_2\), and (iv) produces an isotopically light gypsum-sulfate, could therefore be written (Re. 7):

\[
3\text{Fe}^{32}S + 3\text{CaCO}_3 + \text{FeCO}_3 + 14\text{H}_2\text{O} \rightarrow 4\text{FeOOH} + 3\text{Ca}^{32}\text{SO}_4 + 2\text{H}_2\text{O} + 4\text{CH}_2\text{O}
\]  

(7)

Reaction 7 assumes that the acidity produced by the oxidation of pyrite and its precursors is neutralized by a 3:1 dissolution of calcium to iron carbonate phases in the upper anoxic sediments. The Ca\(^{2+}\) ions released by carbonate dissolution can then co-precipitate with the porewater SO\(_4^{2-}\) ions to form gypsum. The mineral is \( ^{34}S \)-depleted as compared to sulfate dissolved in the monimolimnion but it reflects the \( \delta^{18}O \) signature of the ambient water; with an equilibration effect (i.e., \( e^{18}O_{SO4-amb\,water} \)) which, as mentioned above, ranges between +9.3 and +10.7 ‰.

Pyrite accounted for ≤ 0.5 wt. % of the total XRD-estimated mineralogy of the sediments and occurs as finely dispersed single octahedral crystals that are up to 2 µm in size, and exhibit \{111\} and \{100\} truncations (Fig. 7g). This morphology is
often seen to develop under sulfide-limited conditions in synthetic experiments (e.g., Barnard and Russo, 2009). From the morphology of pyrite and because its $\delta^{34}$S isotope signature differs considerably from those of weathered pyrite in the coal seams-associated lithology (Bouška et al., 1997; Appendix B: Fig B2), this mineral is more probably authigenic in origin. It must have formed locally within the anoxic sediments at low supersaturation, and with nucleation itself depleting the availability of reactants (i.e., $S_2^-$ species) required for further nuclei formation (Rickard and Morse, 2005). Thus, its dispersed, fine crystalline occurrence.

The $\delta^{34}$S isotope values of the finely dispersed pyrite crystals are operationally defined as those of the bulk sediment chromium-reducible sulfide (CRS) pool (i.e., after Canfield et al., 1986). In the upper anoxic sediments, this CRS pool became $^{34}$S-enriched with depth. Accordingly, in the 0 to 4 cm-depth pyrite has $\delta^{34}$S$_{CRS}$ isotope values of -34.7 ± 0.4 ‰. At 4 to 8 cm sediment depth, however, it is relatively $^{34}$S-enriched ($\delta^{34}$S$_{CRS}$ = -23.9 ± 0.9 ‰). Pyrite captures the isotopic signature of dissolved sulfide in its local precipitation environment, and at and near the SWI this mineral appears to have recorded an isotopic offset ($^{34}$ε$_{CRS-SO_4}$) of around 38 ‰ relative to the $\delta^{34}$S$_{SO_4}$ of the monimolimnion. This magnitude of fractionation in pyrite could be ascribed to incomplete microbial sulfate reduction) also perhaps with an additional open system oxidative sulfur cycling (Johnston et al., 2005, Zerkle et al., 2010), and it may point to our biogenic pyrite results from the activity of bacteria capable of fully oxidizing the organic substrates scarcely available (Canfield, 2001; Brückert, 2004). This explains the observed depletion of lactate and pyruvate at this depth in the water column. Limited microbial sulfate reduction is consistent with the fact that pyrite in the lacustrine sediments under consideration precipitates without triggering quantitative sulfate exhaustion (Scholz, 2018; Canfield, 2001)

Approximately 10 ‰ $^{34}$S isotope enrichment in authigenic pyrite at the bottom of our section hints to an additional heavy CRS formation mechanism being more active deeper within the anoxic sediment pile. It could also be the case that the $\delta^{34}$S$_{SO_4}$ values in porewaters in equilibrium with the heavier pyrite are evolved because of variable fractionations associated with MSR (Canfield, 2001; Brückert, 2004). The $\delta^{34}$S values of pyrite from the lower part of the cores also exhibit a narrower difference when compared with those of coexisting authigenic gypsum as shown in Fig. B2 (Appendix B). We can attribute these results to a greater abundance of highly reactive Fe$_{ox}$ phases capable of oxidizing monosulfide (Table 2) in the lower part of the cores investigated.

The CRS pool also includes the sediments’ $S^0$ fraction (Canfield et al., 1986), and given that $S^0$ derived from the chemolithotrophic oxidation of sulfide is relatively $^{34}$S-enriched (e.g., Zerkle et al., 2016; Pellerin et al., 2019), we suggest that the $^{34}$S enrichment in the bottom sediments fingerprints the incorporation of some isotopically heavier $S^0$ into the CRS pool. This interpretation is not only consistent with the decreased proportions of Fe$_{py}$ in the lower part of the sediment pile (Table 2), but also with microbial disproportionation-induced fractionation (e.g., Pellerin et al., 2019).
4.7 Perspectives for deep time palaeoceanographic and/or diagenetic interpretations

The redox stratified bottom waters of Lake Medard demonstrate that ferruginous conditions can develop without substantial sulfate consumption (Scholz, 2018, and references therein). The current lake system thus provides the opportunity to investigate biogeochemical controls active under a transitional state between nitrogenous and sulfidic conditions. This state has not been observed in the scarce examples of redox stratified marine basins existing today (Lyons et al., 2009), nor in the few natural mesotrophic to eutrophic meromictic lakes also used as analogues for disentangling redox boundary conditions in ancient oceans. Similar transitional states would have been more prevalent at times with decreased seawater sulfate concentrations and diminished shuffling of Fe(II) to sediments. Together these factors would have enabled more widespread Phanerozoic ferruginous conditions (Reershemius and Planavsky, 2021) that at times also encompassed the water column of epicontinental seas (e.g., Petrash et al., 2016).

The link between the biogeochemical controls operating in the water column of our study site and the mineral equilibrium conditions prevailing near and at its anoxic SWI may also be relevant for studying elusive shallow burial diagenetic signals developed in fluid-buffered sediments. Also, to unravel long-term overprinting of redox proxies in carbonates altered in movable redox stratified coastal aquifers (Petrash et al., 2021).

In deeper geological time, the increased delivery of continental sulfate to Precambrian sediments containing not only iron oxyhydroxides but also siderite, probably triggered early diagenetic reactions similar to those reported here (e.g., Bachan and Kump, 2015). Comparable diagenetic hydrochemical conditions would have arisen as well when transgressions of ferruginous seawater affected evaporitic facies buried by coastal progradation. In this scenario, the low preservation potential of gypsum would have hindered direct interpretations of any possible isotopic offset recorded by its more stable replacive phases (e.g., silicified calcite or dolomite).

Although gypsum is rarely preserved in Proterozoic shallow-marine successions (but see Blättler et al., 2018), pseudomorphic carbonates after this mineral are volumetrically important in many Precambrian peritidal facies. In such facies, primary gypsum was often replaced by a metastable early diagenetic phase (e.g., Philippot et al., 2009). In a modern thrombolite-forming environment, Petrash et al. (2012) describes an early replacement process of gypsum that involves initial replacement by metastable aragonite. This produces Sr carbonate signals in pseudomorphic calcite replacing aragonite that depart from the Sr content of the ambient water, and, by analogy, can disguise an ancient primary gypsum mineralogy. Similarly to Sr, the structurally substituted sulfate in the carbonate lattice (CAS) of peritidal carbonates (i.e., as a putative proxy for contemporary Proterozoic seawater sulfate) can also be altered early during diagenesis, and exhibit isotope signals incompatible with those of coexisting pyrite (Blättler et al., 2020). This the $\delta^{34}$S signatures of these phases—if formed contemporaneously—shall be expected to approach one each other as per the low dissolved sulfate levels generally ascribed to Proterozoic oceans (e.g., < 400 µM, Fakhraee et al., 2019). An explanation for such a discrepancy is that the CAS and pyrite S isotope proxies recorded the pore fluid signal of diagenetically evolved sulfate in Precambrian (e.g., Rennie and Turchyn, 2014; Li et al., 2015), and some Phanerozoic (e.g., Thomazo et al., 2019) evaporitic/stromatolitic facies
Conversely, a similar inconsistency could arise when early diagenesis ensued transient out-of-equilibrium water column conditions equivalent to those currently prevailing in Lake Medard, where dissolved Fe$^{2+}$ is the dominant redox species, substantial dissolved and solid phase sulfate are present, and the upper stocks of Fe$_{HR}$ effectively buffer dissolved sulfide accumulation.

5. Conclusions

We investigated biomineralization reactions occurring, and prokaryotes thriving in the ferruginous and sulfate-rich bottom water column of a post-mining lake. For this purpose, we considered the pools and fluxes of iron, manganese, carbon, nitrogen, and sulfur in the bottom redox stratified water column and upper reactive sediments (Fig. 8). Discrete spectroscopic datasets were combined with a 16S rRNA gene-aided inference of the planktonic prokaryote community structure to unravel the mechanisms procuring and/or consuming bioactive nitrogen, iron, and sulfur species in the redox stratified ecosystem. Integration of these datasets provides evidence for niche differentiation, but despite marked redox gradients in the water column, we observed a sustained genetic potential for anoxic sulfide oxidation and intermediate sulfur disproportionation. The processes were further substantiated by using sulfate S and O isotope systematics as summarized in Fig. 8. Microbe-mineral interactions near the anoxic sediment-water interface modulate the aqueous equilibrium of both reactive authigenic and redeposited Fe- and Mn-bearing phases. A vigorous anoxic sulfide oxidation pathway is coupled to the reduction and solubilization of the ferric and manganic particulate stocks of the lacustrine sediment (Fig. 8).

Dissolved sulfate need not to be quantitatively depleted for the establishment of ferruginous conditions in the water column. The aqueous system-scale reactions currently proceeding in the redox stratified water column and upper anoxic sediments of Lake Medard are relevant for describing transient redox imbalanced stages between nitrogenous and ferruginous conditions that developed in water columns of ancient nearshore marine settings featuring decreased but not exhausted sulfate levels. These could have produced some of the conflicting signatures often described for coexisting phases of interest as peloeoredox proxies, e.g., carbonates and sulfides. The effects in the geochemical record of analogue imbalanced states are yet to be fully accounted for. This research effort has implications for untangling the deep time palaeoceanographic redox structure of continental margins. We anticipate that further studies in the ferruginous artificial lacustrine system targeted here can provide a more complete picture depicting processes recorded by conflicting proxies in several key, well-preserved Precambrian shallow marine facies.
Figure 8. Scheme summarizing the speciation and stable isotopes ranges of sulfur-bearing phases and siderite and the biogeochemical cycling mechanisms likely operating in the redox stratified Lake Medard and its SWI. A prokaryote community that is to some extent compartmentalized, mediates in the interlinked C, N, S, and Fe and Mn species transformations observed across the redoxcline. This community exerts an influence on the concentration gradients of other dissolved bioactive species, such as phosphate. Towards the hypolimnion, particulate matter formation involves a microaerophilic iron oxidizers/ nitrate reducer community (e.g., Gallionellacea). They promote a continuous amorphous iron aggregate export down to the ferruginous bottom water-sediment interface, where these aggregates may stabilize and/or be reductively dissolved by iron reducers (e.g., Geobacter spp.). In the sediment, stocks of pre-existing siderite and recently stabilized oxyhydroxide phases fuels anaerobic oxidation and disproportionation of by-product sulfide from MSR. In consequence, our coupled stable oxygen and sulfur isotope-based SRR estimate indicates no net sulfate reduction, despite an increased genetic potential for this pathway being deducted from 16S rRNA gene analysis and concomitant evidence for dissolved SO$_4^{2-}$ consumption, likely involving metastable FeS formation. The $\delta^{34}$S$_{CRS}$ values in the upper part of the sediment pile are consistent with incipient and incomplete MSR-induced fractionation, yet MSR is not accompanied by dissolved sulfate depletion, probably because of a low organic substrate availability and due to bioenergetic considerations given by the presence of dissimilatory iron reducers and an abundance of Fe(III) substrates. Importantly, the $\delta^{34}$S of the CRS pool at the lower sediment pile likely incorporates a heavier S signature from intermediate sulfur. Acidity generated by anaerobic S oxidation reactions proceeding near and at the SWI is neutralized by partial carbonate dissolution, which in turn provides Ca$^{2+}$ ions for interstitial microcrystalline gypsum precipitation. This gypsum $\delta^{34}$S values might fingerprint intermediate sulfur disproportionation. The carbonate mineral, on the other hand, may be undergoing recrystallization and growth below 4 cm sediment-depth, such as evidenced by increase FeC contents and the absence of corroded crystal surfaces in the lower part of the sedimentary section examined here.
The northwest Bohemia (Czechia) was an intracontinental lowland comprised of peatlands and isolated ephemeral lakes and peat bogs in the Late Eocene. This landscape developed and expanded in association with subsidence in the Eger rift, which was in turn related to the Alpine and Pyrenean orogeny (Dèzes et al., 2004). By the Oligocene, the lowlands extended over an area >1,000 km$^2$ along the Sokolov and Most basins (Matys Grygar et al., 2014). Thus, organic-rich peatlands now encompass humite lignite seams that correlate across the Czech-Germany boundary and toward Polish Silesia. The extended wetlands along the Eger continental rift turned, by the beginning of the Miocene—as subsidence in the rift graben increased, into a large lake affected by exhalative hydrothermal inputs (Pačes and Šmejkal, 2004) and episodically by alkaline volcanism (Ulrych et al., 2011). Deposits on this paleolake represent the last interval of the syn-rift sedimentation and consist of on 70–120 m thick carbonate-rich, kaolinitic coal-bearing claystone with several horizons of tuff material. These deposits are lithostratigraphically referred to as the Cypris Formation (Kříbek et al., 1998, 2017), and now outcrop in elevated areas of the Sokolov mining district, where they overlie the coal seams that were exploited to exhaustion in the former Medard open-cast mine.

The Miocene Cypris Formation and quaternary alluvions comprised of material largely derived from this unit functioned as the main source of sediments to the modern post-mining lacustrine system. The mineral assemblage of the stratigraphic unit includes kaolinite, K-feldspar, quartz, rutile and anatase, and gypsum. It contains also analcime (NaAlSi$_2$O$_6$), weathered pyrite, variable proportions of carbonates (calcite, Fe-dolomite, and siderite), and greigite (Fe$_3$S$_4$) (Murad & Rojík, 2003, 2005). The unit is affected by localized thenardite (Na$_2$SO$_4$) efflorescences that are associated with fluid flow along faults and fractures in subjacent rocks (Šmejkal, 1978; Pačes & Šmejkal, 2004). Organic matter in the Cypris claystone exhibits significant variability. These changes recorded discontinuous development of widespread anoxia across the intracontinental depositional system and were accompanied also by shifts in salinity and alkalinity. This promoted lacustrine carbonate deposition (Kříbek et al., 2017). Overall, the authigenic mineral assemblage, elemental concentration trends, and the heavy O and S isotopic signatures of secondary sulfate minerals of the Cypris claystone (Fig. SM2) indicate that its deposition occurred in a large saline playa paleolake in which the oxidative weathering of sulfides, volcanic exhalations, and meteoric water-rock interactions imparted a major geochemical imprint that is superimposed to that of the episodic changes in the paleolake’s redox conditions (Šmejkal, 1978; Pačes & Šmejkal, 2004). A compilation of the $\delta^{34}$S of the sulfate sourced from the Miocene Cypris claystone is shown in Figure B2 (Appendix B). As discussed in the main text, the dissolved sulfate of the modern redox stratified bottom waters appears to fingerprint these sources.
Appendix B
Figures B1 and B2

Figure B1. Pourbaix (pH–Eh) diagram of the thermodynamically stable Fe and S phases in the ferruginous and sulfate-rich bottom waters of Lake Medard.

Figure B2. A comparison of the range of reported $\delta^{34}$S values of potential sources of oxidized sulfur to Lake Medard (after Šmejkal 1978; Krs et al., 1990 for greigite), the sulfate-rich bottom water column and authigenic gypsum and pyrite in the upper anoxic sediments (i.e., this work) are also shown (filled boxes).
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