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Viable cold-tolerant iron-reducing microorganisms in geographically diverse subglacial environments

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Abstract. Subglacial environments are known to harbour metabolically diverse microbial communities. These microbial communities drive chemical weathering of underlying bedrock and influence the geochemistry of glacial meltwater. Despite its importance in weathering reactions, the microbial cycling of iron in subglacial environments, in particular the role of microbial iron reduction, is poorly understood. In this study we address the prevalence of viable iron-reducing microorganisms in subglacial sediments from five geographically isolated glaciers. Iron-reducing enrichment cultures were established with sediment from beneath Engabreen (Norway), Finsterwalderbreen (Svalbard), Leverett and Russell glaciers (Greenland), and Lower Wright Glacier (Antarctica). Rates of iron reduction were higher at 4°C compared with 15°C in all but one duplicated second-generation enrichment culture, indicative of cold-tolerant and perhaps cold-adapted iron reducers. Analysis of bacterial 16S rRNA genes indicates Desulfosporosinus were the dominant iron-reducing microorganisms in low-temperature Engabreen, Finsterwalderbreen and Lower Wright Glacier enrichments, and Geobacter dominated in Russell and Leverett enrichments. Results from this study suggest microbial iron reduction is widespread in subglacial environments and may have important implications for global biogeochemical iron cycling and export to marine ecosystems.

1 Introduction

Despite cold, dark and oligotrophic conditions subglacial environments worldwide harbour diverse microbial communities. Such communities have been documented at the ice–bed interface of valley glaciers (Sharp et al., 1999; Skidmore et al., 2005; Buzzini et al., 2005; Bhatia et al., 2006; Kashtovská et al., 2007; Mitchell et al., 2013; Hamilton et al., 2013), ice sheets (Sheridan et al., 2003; Mikucki and Priscu, 2007; Lanoil et al., 2009), and in subglacial lakes (Priscu et al., 1999; Gaidos et al., 2004; Marteinsson et al., 2013; Christner et al., 2014). Given the absence of light in these environments, microbially mediated redox reactions fuel heterotrophic and autotrophic microbial metabolism and drive chemical weathering reactions of the underlying bedrock (Sharp et al., 1999; Bottrell and Tranter, 2002; Tranter et al., 2005; Wynn et al., 2006; Hodson et al., 2008; Montross et al., 2012; Boyd et al., 2014). To date, subglacial microbiology research has focused on the microbial cycling of carbon (Boyd et al., 2011; Stibal et al., 2012), nitrogen (Skidmore et al., 2000; Boyd et al., 2011) and sulfur (Boyd et al., 2014; Harold et al., 2016). The role of microbial iron cycling, and in particular iron(III) reduction, in subglacial biogeochemical cycling is poorly understood. This is surprising, given the importance of iron in (a) catalysing the weathering of bedrock sulfides and promoting solute production (Bottrell and Tranter, 2002), (b) influencing phosphorus availability to microorganisms via adsorption–desorption reactions on Fe(III) oxide surfaces (Gunnars et al., 2002) and (c) the potential for subglacially derived iron to impact global biogeochemical cy-
icles by fertilizing productivity in downstream ecosystems, including those in marine environments (Death et al., 2014).

Recent research has shown the flux of bioavailable nanoparticulate iron associated with glacial runoff from ice sheets to be an important source of nutrients to the surrounding oceans (Hawkins et al., 2014). These iron oxyhydroxide particles are thought to originate from the oxidation of dissolved ferrous iron from anoxic subglacial environments. The source of this Fe(II) remains unknown, though microbial iron reduction is a plausible but untested explanation (Hawkins et al., 2014).

Although a number of studies have demonstrated the presence of iron-reducing microorganisms in subglacial environments using culture-independent methods (Skidmore et al., 2005; Mikucki and Priscu, 2007; Lanoil et al., 2009; Yde et al., 2010; Marteinsson et al., 2013; Mitchell et al., 2013), few studies have addressed their viability (Foght et al., 2004; Mikucki and Priscu, 2007) or their physiological characteristics. Our hypothesis is that subglacial sediments harbour active microorganisms that are capable of carrying out microbial iron reduction. Specifically, our aims in this study are (1) to assess whether subglacial sediments harbour active iron-reducing microorganisms and (2) whether these microorganisms are adapted to low temperatures. We present data from enrichment cultures initiated from subglacial sediments entombed in basal ice, sampled at the glacier margins of five glaciers geographically isolated from one another: Engabreen, Norway; Russell and Leverett glaciers, Greenland; Finsterwalderbreen, Svalbard; and Lower Wright Glacier, Antarctica. We demonstrate that viable iron-reducing microorganisms are present in each of these systems and that they are tolerant of, and may even be adapted to, low temperatures.

2 Materials and methods

2.1 Sample sites and subglacial sample acquisition

Engabreen (E; 66°41′N, 13°46′E) is a temperate glacier of the western Svalbard Icecap in northern Norway. The underlying bedrock is metamorphic, dominated by schists and gneisses with calcite-filled cracks (Jansson et al., 1996), and contains relatively little organic carbon (O’Donnell et al., 2016). The bedrock and basal ice is accessible through a system of underground tunnels leading to the glacier bed from the Svalbard Subglacial Laboratory. This basal ice was sampled by implementing hot-water drilling, and sediment-laden ice removed from the resulting cavity 200 m below the glacier surface using a chainsaw (Stibal et al., 2012).

Russell (R; 67°03′N, 50°10′W) and Leverett (L; 67°03′N, 50°07′W) are neighbouring land-terminating outlet glaciers. Together they constitute a large discharge lobe emanating from the western Greenland Ice Sheet (GrIS); Leverett is the southern offshoot of the larger Russell Glacier. Both are polythermal (cold and warm-based regions exist at the ice–bed interface), but warmer conditions dominate, as evidenced by accelerated glacial flow at the onset of the melt season (Sundal et al., 2011). The underlying bedrock is metamorphic, dominated by Archaean gneiss, which was subsequently reworked in the Early Proterozoic (Henrikson et al., 2000) and during numerous Holocene re-advances, leading to the incorporation of organic matter (Ten Brink and Weedick, 1974). At Russell Glacier, samples were obtained from upheveled subglacial sediment near the terminus. The outermost surface of the ice was removed using a chainsaw before subsampling the remaining sediment-laden ice (Stibal et al., 2012). Two subglacial samples were collected from Leverett; L was chainsawed from a pressure ridge at the glacier margin, believed to be subglacial in origin, and FL (“fresh” Leverett) was collected from the basal sediments at the ice–bed interface within an ice cave using a flame-sterilized spatula. At the time of collection, the latter sediment (FL) was assumed to be freshly melted.

Finsterwalderbreen (F; 77°28′N, 15°18′E) is a polythermal glacier on the southern side of Van Keulenfjorden, Svalbard, Spitsbergen, Svalbard (Wadham and Nuttall, 2002). Similar to the Greenland glaciers, the thermal regime of this glacier is largely warm-based (Hodson and Ferguson, 1999). The major underlying geology is mainly sedimentary, comprising sandstone, siltstone, shale, limestone and dolomites (Dallmann et al., 1990). Basal sediment was removed using a chainsaw from a pressure ridge at the glacier terminus (O’Donnell et al., 2016).

Lower Wright (LW; 77°25′S, 163°00′E) is a cold-based glacier draining westwards from the Wilson Piedmont Glacier in the McMurdo Dry Valleys region of Antarctica. The glacier currently terminates in the permanently ice-covered Lake Brownworth (Stibal et al., 2012). The underlying bedrock is dominated by granite-gneisses (Hall and Denton, 2002); however, during the last few centuries numerous glacial advances have overridden and reworked lake sediments, evidence of which was found in the presence of algal matter in subglacial sediment (Wadham et al., 2012). The sampling site featured exposed frozen sediment sandwiched between layers of pure glacial ice at the interface of the ice-covered lake and the terminal moraines of the glacier. Blocks of this debris-laden ice were removed using a chainsaw (Stibal et al., 2012).

Samples were transported frozen from the field site to LOWTEX (University of Bristol) and stored at −20°C. Samples were prepared at the LOWTEX facility prior to their use in this study. Specifically, sediment-laden basal ice was placed in a laminar-flow hood in pre-furnace glass beakers and the outer layer of each sample was removed by washing with sterile deionized water. Samples were covered in furnace aluminium foil and transferred to an anaerobic chamber, where they were thawed under 100% nitrogen atmosphere. The liberated sediment from basal ice samples was subsampled into sterile serum vials, crimp-sealed with thick
butyl rubber stoppers and aluminium caps, and removed from the chamber. The headspace of each was immediately flushed with N2 gas for 1 min using sterile needles attached to sterile 0.2 µm syringe filters, before being transferred to the University of Edinburgh, where they were stored at 4 °C for 24 h before enrichments were initiated.

2.2 Microbial enrichment cultures

Strict anaerobic culturing techniques were used throughout (Miller and Wolin, 1974). Enrichment cultures were initiated by adding sterile anoxic freshwater basal medium at pH 6.8–7.0 to serum vials containing subgelud sediments. The basal medium contained (grams per L deionized water) NaHCO3 (2.5), NH4Cl (0.25), NaH2PO4 (0.06), KCl (0.1), 10 mL vitamin solution and 10 mL trace elements solution. The vitamin solution contained (mg L–1 deionized water) biotin (2.0), folic acid (2.0), pyridoxine-HCl (10.0), riboflavin (5.0), thiamine (5.0), nicotinic acid (5.0), pantothenic acid (5.0), vitamin B-12 (0.1), 2-aminobenzoic acid (5.0), and thioctic acid (5.0). The trace elements solution contained (g L–1 deionized water) nitritriatic acid (1.5), MgSO4 (3.0), MnSO4·H2O (0.5), NaCl (1.0), FeSO4·7H2O (0.1), CaCl2·2H2O (0.1), CoCl2·6H2O (0.1), ZnCl2 (0.13), CuSO4·5H2O (0.01), AlK(SO4)2·12H2O (0.01), H3BO3 (0.01), NaMoO4 (0.025), NiCl2·6H2O (0.024) and NaWO4·2H2O (0.025). Vitamin and trace element solutions were stored at 4 °C in the dark. Enrichments were supplemented with acetate (10 mM) and lactate (10 mM) as the electron donors and poorly crystalline Fe(III) oxide (~100 mM) as the terminal electron acceptor. This combination of electron donors was chosen since the vast majority of iron-reducing microorganisms can use one or the other in the reduction of ferric iron and indeed most can draw upon a much wider array of organic compounds (Lloyd, 2003). Iron oxide was synthesized as previously described (Lovley and Phillips, 1986b). FeCl2 (1.3 mM) was added to the medium as a mild reducing agent. Enrichments were set up in triplicate and incubated at 4, 15 and 30 °C. Microbial iron reduction was determined by measuring the concentration of HCl-extractable ferrous iron with time, as previously described (Lovley and Phillips, 1986a, b). Second-generation 4 and 15 °C enrichments were initiated in duplicate using a 10% v/v inoculum from initial enrichments. No 30 °C initial enrichments were carried over to a second-generation since the focus of the study was on the presence of cold-adapted iron-reducing microorganisms. Therefore data presented here are from second-generation 4 and 15 °C enrichments and first-generation 30 °C enrichments. A production of ≥ 5 mM Fe(II) over a period of 50 days was considered positive for microbial iron reduction. This concentration was chosen as a nominal figure that greatly exceeded the typical error of the ferrozine assay, as measured in calibration procedures where triplicate ferrozine measurements are conducted on calibration standards ranging in concentration from 1 to 50 mM. Blanks were not initiated in parallel to these enrichments. However, in other experiments using the same medium and electron donor–acceptor couple, no abiotic production of Fe(II) was observed over a longer period of time (see Fig. S1 in the Supplement).

2.3 Bacterial community analysis

DNA was extracted from 5.0 mL of one replicate of each second-generation 4 °C enrichment culture using the FastDNA(R) SPIN kit for Soil (MP Biomedicals, Illkirch, France). Each 5 mL subsample was concentrated by centrifugation at 4570 × g for 15 min (Heraeus Multifuge 3SR+ with swung-out rotor, Thermo Scientific, UK) and resuspended in 500 µL sterile basal medium. DNA extraction was carried out according to manufacturer’s protocol. Early attempts to extract DNA from 4 °C LW and F enrichments failed. Subsequent attempts were successful after incorporating the protocol of Direito et al. (2012) to counteract any adsorption of DNA onto clay mineral matrices. Specifically, the FastDNA SPIN kit phosphate buffer was substituted for an equal volume of 1 M sodium phosphate buffer in 15 % molecular-grade ethanol, pH 8.0. Samples were incubated in a heating block at 80 °C for 40 min following the lysis step in the FastDNA SPIN kit protocol. All subsequent processing was as instructed in the manufacturer’s protocol.

Extracted DNA from E, L, FL and R enrichments was amplified for the V3 hypervariable region of the 16S rRNA gene using primers 27F (5′-AGA GTT TGA TCC TGG CTC AG-3′) and UN1492R (5′-TAC GGT TAC CTT GTT ACG ACT T-3′) (Lane, 1991). Each 50µL PCR mixture contained 1 µL DNA template, primers (0.4 µM each), deoxynucleoside triphosphates (dNTPs; 200 µM), 1.5 mM MgCl2, 5 µL 10 X PCR buffer, 5 µL bovine serum albumin (BSA), and 0.5 µL Taq polymerase. Template DNA was initially denatured at 94 °C for 4 min, followed by 30 cycles of denaturing (30 s at 94 °C), annealing (30 s at 55 °C), and extension (60 s at 72 °C), and a final extension at 72 °C for 5 min.

DNA extracted from F and LW was amplified for the V3 hypervariable region of the 16S rRNA gene using primers 357F (5′-CCT ACG GGG AGG CAG CAG-3′) and 518R (5′-ATT ACC GCG GCT GGT GG-3′) (Muyzer et al., 1993). The PCR mix was kept as described above. For this primer set, template DNA was initially denatured at 94 °C for 4 min, followed by 35 cycles of denaturing (30 s at 94 °C), annealing (30 s at 54 °C) and extension (30 s at 72 °C), and a final extension at 72 °C for 5 min.

Amplified DNA was sequenced for bacterial 16S ribosomal RNA using the Roche 454 Pyrosequencing platform (Research and Testing Laboratories, Austin, TX, USA). Resulting sequence reads were quality checked and analysed via the Research and Testing Laboratories pipeline. This analysis is split into two stages. Firstly, sequences were quality trimmed, denoised, and checked for chimeras (Edgar, 2010; Edgar et al., 2011). Secondly, phylogenetic analysis was carried out, involving clustering sequences into oper-
3 Results

3.1 Occurrence of microbial iron reduction

Results for positive or negative detection of microbial iron reduction in 4, 15 and 30 °C enrichment cultures after 50 days are summarized in Table 1. All 4 °C enrichments tested positive for microbial iron reduction. In contrast, half of enrichments incubated at 15 °C and only one of six incubated at 30 °C were positive for microbial Fe(III) reduction. Particularly high Fe(II) concentrations were detected in 15 °C Leverett (L), in which 37.75 ± 0.002 mM was measured after 58 days.

The relative rates of microbial iron reduction in 4 °C compared with 15 °C duplicated enrichments are shown in Fig. 1 and given in Table 2. With the exception of L, significantly greater microbial iron reduction is evident at T = 58 in 4 °C compared to 15 °C enrichments (Student’s t test, two-tailed, type 2: p values < 0.005), indicative of cold-tolerant iron-reducing microorganisms. In the case of Leverett, iron reduction in the 15 °C enrichment was significantly higher (p value < 0.05).

3.2 Phylogenetic diversity of cold-tolerant enrichment cultures

Genus-level taxonomic affiliations of 16S rRNA genes amplified from DNA extracted from second-generation 4 °C enrichments are shown in Fig. 2, along with corresponding Shannon’s diversity indices (H). Overall genus-assigned diversity is highest in the second-generation enrichment initiated with L (Shannon’s H of 1.92), followed by LW (H of 1.39), whilst the enrichment initiated from F is characterized by the lowest diversity (Shannon’s H of 0.26), followed by E (H of 0.74). Enrichments from FL and R exhibit similar overall diversity (H indices of 1.00 and 0.97, respectively). Sequences affiliated with Desulfovibrio (95% sequence match) dominate the second-generation enrichment cultures originating from samples E (60.3%), F (94.3%) and LW (54.6%) and are present but not dominant in R (23.0%) and FL (3.4%). In contrast, sequences affiliated with Geobacter (95% sequence match) dominate enrichments from FL (78.2%) and R (66.0%). Sequences affiliated with Geobacter are also present in L (14.0%) but are absent in E, F and LW. Other identified genera of relevance to microbial iron reduction are Desulfotignum (23.5% of sequence reads from LW) and Rhodotherax (18.30% of sequence reads from L).

A number of the 16S rRNA gene sequences from each enrichment could be resolved to species-level taxonomic
identification. Of relevance for microbial iron reduction are *Geobacter psychrophilus* (>97% sequence match), which accounted for 46.3% of the *Geobacter* reads in FL; *Rhodoferax ferrireducens* (>97% sequence match) in enrichments L (16.2%) and FL (0.08%); and *Desulfosporosinus lacus* (>97% sequence match) in enrichments E (0.08%) and R (0.01%).

4 Discussion

4.1 Distribution and characteristics of iron-reducing microorganisms in subglacial sediments

Very limited information exists on the distribution and activity of microbial iron reduction beneath glaciers and ice sheets, despite the potential for this process to contribute iron to downstream ecosystems through glacial runoff. The iron-reducing enrichment cultures initiated with subglacial sediments in the present study indicate that viable iron-reducing microorganisms are widespread beneath glaciers and, based on activity data, are likely to be cold-adapted. Members of the *Geobacter* genus dominate two of the cold temperature enrichments (FL and R) while *Desulfosporosinus* dominates three of the environments (E, F, LW; see Fig. 2). These two genera appear to be responsible for most of the iron reduction observed, though species of *Desulfitobacterium* and *Rhodoferax* may also contribute. Further work is warranted to demonstrate that microorganisms present in these sediments that belong to these genera are indeed capable of microbial iron reduction (MIR) and to attempt to quantify their contribution to iron and carbon cycling in situ.

The possibility that iron reduction detected in the enrichments was indirect cannot be ruled out. Given the prevalence of sulfate-reducing genera such as *Desulfosporosinus*...
in these positive enrichments, the most plausible indirect mechanism of reduction would be via the abiotic reduction of ferric iron by sulfide following microbial sulfate reduction. However no sulfide production (characterized by non-magnetic black precipitate or the recognizable rotten egg smell) was observed in these enrichments. Furthermore, the data presented in Fig. 1 are for second-generation enrichments initiated with a 10% (v/v) inoculum from initial enrichments. Since no sulfate was added to enrichment medium, and no sulfide was observed in either stage of enrichments, we believe our data provide evidence of direct iron reduction.

Compared to other, less comprehensive studies, our results suggest microbial iron reduction in subglacial environments is characterized by substantial metabolic and genetic diversity. The vast difference in phylogenetic diversity between two low-temperature enrichments initiated with sediments from different locations beneath the Leverett Glacier (Fig. 2) further highlights this diversity. The presence of iron-reducing microorganisms in subglacial systems has been identified using culture-independent methods in a number of studies (Skidmore et al., 2005; Lanoil et al., 2009; Yde et al., 2010; Marteinsson et al., 2013; Mitchell et al., 2013), yet only two studies investigated subglacial sediments proximal to those used here. Yde et al. (2010) identified 16S rRNA gene sequences affiliated with the genera *Rhodoferax* and *Geobacter* in basal ice from the margin of Russell Glacier. More recently, Cameron et al. (2016) also found evidence for the presence of *Rhodoferax*, *Geobacter* and *Desulfoferax* in 16S rRNA gene sequence libraries from subglacial sediments draining Leverett Glacier; of these *Rhodoferax* was particularly abundant, representing more than 20% of reads in some samples. Similarly, Mitchell et al. (2013) detected 16S rRNA gene sequences affiliated with *Rhodoferax* in sediments sampled from beneath Robertson Glacier, Alberta, Canada; sequences affiliated with *Geobacter* were not detected. In this study we detected sequences closely related to *Rhodoferax* as the dominate sequence type in enrichments from Leverett Glacier (Fig. 2), though sequences affiliated with *Geobacter* were also detected in the enrichment.

To date, only two studies have enriched for MIR in subglacial sediments. The first, by Foght et al. (2004), enriched for iron-reducing microorganisms in sediments from beneath two temperature glaciers in New Zealand using ferric citrate, yeast extract and tryptone in a semisolid medium. The enrichments were positive for microbial iron reduction, but the study found the microorganisms responsible to be few in number. This is consistent with the expectation that iron-reducing microorganisms should not be dominant in subglacial communities, since they depend on a supply of organic carbon from primary producing chemolithoautotrophs (e.g. Boyd et al., 2014). It is possible that organic carbon of pre-glacial origin (legacy organic carbon) could serve as a carbon source, but this has yet to be demonstrated. The second study to enrich for microbial iron reduction was conducted by Mikucki and Priscu (2007), using outflow sediments from Blood Falls in Antarctica as their inoculum and medium containing amorphous iron oxide and yeast extract, with a pressurized H<sub>2</sub> headspace. The 16S rRNA analysis of the sediment identified sequences closely related to the iron reducer *Geopsychrobacter electrophilus*, although an isolate from a positive microbial iron reduction enrichment was a close relative of *Shewanella frigidimarina*. In all other studies *Rhodoferax* has been reported and implicated in subglacial MIR (Skidmore et al., 2005; Lanoil et al., 2009; Mitchell et al., 2013).

Although members of the genera *Desulfoferax* are conventionally thought of as sulfate-reducing bacteria, results from this study serve as compelling evidence that they are capable of low-temperature iron reduction, though visually undetectable levels of sulfate reduction in the enrichment cultures cannot be ruled out. Only one characterized strain belonging to this genus is known to carry out dissimilatory iron reduction: *Desulfoferax lacus* (Ramamoorthy et al., 2006). This species was identified in two of the enrichments in which *Desulfoferax* was abundant, namely Engabreen and Russell. This genus has only been reported in one other study of cold glacial ecosystems (Marteinsson et al., 2013), in which a number of bacterial 16S rRNA gene clones from DNA extracted from subglacial lake sediments from beneath Vatnajökull ice cap in Iceland were closely associated with *Desulfoferax*. The authors hypothesized that *Desulfoferax* is carrying out the reduction of sulfate and other oxidized sulfur species in this environment, whereas *Geobacter*, also detected in their bacterial clone library, carries out ferric iron reduction. The results from
our study suggest that Desulfosporosinus may be able to reduce iron in this subglacial lake. Indeed a number of sulfate-reducing taxa are capable of conserving energy from iron reduction (e.g. Desulfospororum polyspora, Sass et al., 2004; Desulfotalea psychrophila, Knoblauch et al., 1999), which is consistent with the higher energy yield associated with iron reduction when compared to sulfate reduction (Neal et al., 2003).

Whilst a true determination of psychrophily requires metabolic and growth data, on the basis of activity data alone, the iron-reducing microorganisms enriched in our current study appear to be cold-adapted, and at least cold-tolerant (Fig. 1). While numerous studies have provided evidence for cold-adapted microorganisms (e.g. Thiothriases sp. RG5; Harrold et al., 2016) and their activities (e.g. nitrification and nitrate reduction (Boyd et al., 2011) and methanogenesis (Telling et al., 2015)), only a handful of iron-reducing microorganisms have been characterized as psychrophilic to date, namely Desulfogrificus oceanense (optimal $T = 10^\circ$C, min. $-1.8^\circ$C; Knoblauch et al., 1999), Desulfuromonas valbardenis and D. ferrireducens (both optimal $14^\circ$C, min. $-2^\circ$C; Vandeike et al., 2006). A larger number of iron-reducing microorganisms can tolerate temperatures near or below freezing (e.g. Pelobacter propionicus, Schink, 1984; Lonergan et al., 1996), and the sediment samples used in this study had been frozen to $-20^\circ$C for months to years prior to use in enrichments. However, the vast majority of characterized iron-reducing strains are psychrotolerant, capable of growth at low temperatures but have much higher optimal growth temperatures, such as strains of Geobacter (Nevin et al., 2005; Sung et al., 2006; Shelobolina et al., 2007, 2008), Desulfuromusa (Liesack and Finster, 1994) and Pelobacter (Schink, 1984). The results from our study suggest that iron-reducing microorganisms in a variety of subglacial ecosystems are adapted to the cold conditions that characterize these environments.

4.2 Potential controls on activity of iron-reducing microorganisms beneath glaciers

A key determinant of the in situ activity of iron-reducing microorganisms is the availability of electron donors and ferric iron, as well as the absence of oxidizers with higher reduction potentials such as oxygen ($O_2$) or manganese oxide ($MnO_2$), which should be preferentially utilized for energetic reasons. In anoxic environments depleted in $MnO_2$, understanding the supply of both electron donors and ferric iron is important in constraining the activity of iron-reducing microorganisms in situ. We note that the type and availability of organic matter will vary greatly between subglacial catchments, depending on factors such as hydrological isolation from the surface, timings of advance and retreat over soils and vegetation, and underlying lithology. Although some glaciers have overridden soils, lacustrine or marine deposits rich in organic carbon of different bioavailabilities (e.g. Russell and Leverett

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dissolved ferrous iron from anoxic subglacial environments. The source of this Fe(II) remains unknown, though microbial iron reduction is a plausible, albeit underdemonstrated, explanation (Hawkings et al., 2014). Results from our study indicate that microbial Fe(III) reduction is a possible source of the Fe(II) emanating from glaciers worldwide and may be responsible for a portion of the Fe(III) flux to marine ecosystems from glacial systems. Since Fe(III) availability often limits the primary production of marine phototrophs, the activity of iron-reducing bacteria in subglacial environments could have an indirect but significant effect on global carbon budgets (Statham et al., 2008; Death et al., 2013). Iron cycling can also have an important impact on the availability of phosphorus to microorganisms, due to the adsorption of phosphorus to solid phase Fe oxyhydroxide particles (Gunnars et al., 2002). Further studies are required to characterize in situ Fe(III) reducing metabolic activity and to quantify its impact on the export of bioavailable iron to past and present marine ecosystems. Subglacial environments are considered analogous to potential habitats for life on Mars (Skidmore et al., 2000; Christner et al., 2008; Fisher and Schulze-Makuch, 2013), and microbial iron reduction has been identified as a plausible metabolism to fuel microbial life (Nixon et al., 2012). This study lends weight to this hypothesis and suggests similar past or present perennially cold and dark environments are within the bounds of habitability.

5 Data availability

The raw sequencing data have been submitted to NCBI Sequence Read Archive (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA374495, Nixon et al., 2017) with the project accession number of SRP099404 (samples E, L, FL and R are under accession no. SRR5332312 and samples LW and F under accession no. SRR5332311).

The Supplement related to this article is available online at doi:10.5194/bg-14-1445-2017-supplement.

Competing interests. The authors declare that they have no conflict of interest.

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