Hydro-biogeochemical coupling beneath a large polythermal Arctic glacier: implications for subice sheet biogeochemistry

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Hydro-biogeochemical coupling beneath a large polythermal Arctic glacier: Implications for subice sheet biogeochemistry

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[1] We analyze the interannual chemical and isotopic composition of runoff from a large, high Arctic valley glacier over a 5 year period, during which drainage evolved from a long-residence-time drainage system feeding an artesian subglacial upwelling (SGU) at the glacier terminus to a shorter-residence-time drainage system feeding an ice-marginal channel (IMC). Increased icemelt inputs to the SGU are thought to have triggered this evolution. This sequence of events provides a unique opportunity to identify coupling between subglacial hydrology and biogeochemical processes within drainage systems of differing residence time. The biogeochemistry of the SGU is consistent with prolonged contact between meltwaters and subglacial sediments, in which silicate dissolution is enhanced, anoxic processes (e.g., sulphate reduction) prevail, and microbially generated CO₂ and sulphide oxidation drive mineral dissolution. Solute in the IMC was mainly derived from moraine pore waters which are added to the channel via extraglacial streams. These pore waters acquire solute predominantly via sulphide oxidation coupled to carbonate/silicate dissolution. We present the first evidence that microbially mediated processes may contribute a substantial proportion (80% in this case) of the total glacial solute flux, which includes coupling between microbial CO₂-generation and silicate/ carbonate dissolution. The latter suggests the presence of biofilms in subglacial/ice-marginal sediments, where local perturbation of the geochemical environment by release of protons, organic acids, and ligands stimulates mineral dissolution. These data enable inferences to be made regarding biogeochemical processes in longer-residence-time glacial systems, with implications for the future exploration of Antarctic subglacial lakes and other wet-based ice sheet environments.


1. Introduction

[2] There is mounting evidence that the beds of glaciers and ice sheets are host to diverse communities of microorganisms [Sharp et al., 1999; Skidmore et al., 2000; Gaidos et al., 2004; Mikucki et al., 2004; Skidmore et al., 2005; Foght et al., 2004]. These microorganisms are intimately involved in chemical weathering, mediating both oxic and anoxic oxidation of sulphide minerals, the latter using NO₃ and Fe(III) [Bottrell and Tranter, 2002; Wadham et al., 2004; Wynn et al., 2006], and adding CO₂ to meltwaters through respiration [Tranter et al., 2002]. In doing so, they perturb local pH/Eh conditions and create microniches for microbial colonization [Tranter et al., 2005]. Recent work has highlighted the potential importance of these deep subsurface microbial communities in regulating global biogeochemical cycles. For example, the release of bioavailable iron from sediments in melting icebergs is a potential stimulant of ocean primary productivity, with consequent negative feedback effects on climatic warming via CO₂ drawdown [Raiswell et al., 2006, 2008]. Others have suggested that anoxia generated beneath continental ice sheets may stimulate microorganisms to produce methane during periods of ice advance and carbon burial [Skidmore et al., 2000; Weitemeyer and Buffett, 2006; Wadham et al., 2008]. This methane may be stored as clathrate beneath the ice sheet until ice sheet decay stimulates its rapid release to the atmosphere [Weitemeyer and Buffett, 2006].

[3] While the activity of subglacial microbial communities in wet-based zones of glaciers and is well established
Table 1. Description of Changes in Drainage on the Western Finsterwalderbreen Margin

<table>
<thead>
<tr>
<th>Year</th>
<th>Drainage Description</th>
<th>Estimation of the % of Meltwater Routed by the SGU/IMC</th>
<th>Data Employed in This Study</th>
<th>Source Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>1994</td>
<td>SGU sustained for the 1994 melt season</td>
<td>&lt;10%/&gt;90%</td>
<td>SGU major ion chemistry</td>
<td>Wadham et al. [1998]</td>
</tr>
<tr>
<td>1995</td>
<td>SGU sustained for the 1995 season and routed snowmelt to the terminus</td>
<td>&lt;10%/&gt;90%</td>
<td>SGU major ion chemistry</td>
<td>Wadham et al. [1998]</td>
</tr>
<tr>
<td>1997</td>
<td>SGU captures drainage from IMC and conveys 90% of meltwater drainage on the western margin</td>
<td>&gt;90%/&lt;10%</td>
<td>SGU major ion chemistry plus δ18O and δ34S of sulphate</td>
<td>Wadham et al. [2004]</td>
</tr>
<tr>
<td>1999</td>
<td>SGU no longer present, and all meltwater emerged via the IMC</td>
<td>0%/100%</td>
<td>SGU major ion chemistry, plus δ18O and δ34S of sulphate</td>
<td>Wadham et al. [2007]; Cooper et al. [2002]</td>
</tr>
</tbody>
</table>

*Data presented in this paper and source references.*

[Tranter et al., 2005], there is little data on microbial processes beneath ice sheets and the sensitivity of subglacial biogeochemistry to temporal shifts in the configuration and characteristics of meltwater flow paths. Any reorganization of subglacial flow paths and associated changes in meltwater residence times and rock:water ratios will impact on the balance of biogeochemical processes and associated export of key nutrients to downstream environments. Valley glaciers possess dynamic subglacial drainage systems, which evolve seasonally from long-residence-time distributed drainage systems (residence times of days to months) to discrete networks of channels (residence times of hours) [Nienow et al., 1998]. Recent work has demonstrated the presence of subglacial meltwater routing at the base of the Greenland Ice Sheet, driven by rapid drainage of surface lakes [Das et al., 2008]. Beneath Antarctica subglacial lakes form via the accumulation of meltwater generated by geothermal heating of basal ice, and periodically drain when their volume attains a critical threshold [Wingham et al., 2006; Fricker et al., 2007]. This suggests that some of the basal regions of ice sheets may evolve their drainage system structure over time, with consequent perturbation of subglacial biogeochemistry and microbial community composition [Tranter et al., 2005; Hodson et al., 2008]. While hydro-biogeochemical data from ice sheets is sparse, insights into the coupling between hydrological drainage system evolution and subglacial biogeochemistry can be gained from large valley glacier systems. Here, a polythermal regime (similar to that on ice sheet margins) and longer flow paths for meltwaters (and hence, longer water residence times) provide case studies that are intermediate between small, temperate valley glaciers and continental-scale ice sheets.

We employ the interannual chemical and isotopic composition of runoff from a large, high Arctic valley glacier, Finsterwalderbreen (Svalbard), over a 5 year period (1994–1999) to infer changes in subglacial biogeochemistry associated with drainage reorganization. Previous work from individual years [Wadham et al., 1998, 2000, 2004, 2007] has indicated that two principal drainage systems contribute runoff to the western glacier margin: a predominantly long-residence-time drainage system feeding an artesian subglacial upwelling outflow (SGU) and a shorter residence time channelized drainage system, which culminates in an ice-marginal channel (IMC). We estimate that residence times in the SGU are >1 day, and a matter of hours in the IMC. Independent hydrological observations over the 5 year study period clearly indicate that the SGU routed an increasing proportion of runoff on the western margin between 1994 and 1997, accounting for 90% of runoff by 1997 and forming an artesian fountain >1 m high at the terminus. However, the SGU was no longer present during the summers of 1999 and 2000 and all runoff was routed entirely by the IMC, signifying a shift in meltwater routing. This sequence of events provides a unique opportunity to identify (1) the forcing mechanisms for this subglacial drainage evolution and (2) modes of coupling between subglacial biogeochemistry and hydrology on larger glacier systems during drainage reorganization. In doing so, it provides a framework for testing a series of hypotheses regarding biogeochemical processes beneath continental-scale ice sheets (e.g., Antarctic subglacial lakes and their interconnecting flow paths). It has particular relevance to the Greenland Ice Sheet under climate warming scenarios where an increased extent of surface melting may initiate surface to bed hydrological coupling for the first time in areas where previously all melt was supplied from basal melting and routed via a distributed drainage system [Steffen et al., 2004].

2. Field Study Area

[5] Finsterwalderbreen (77°28′N, 15°18′E) is located on the southern side of Van Keulenfjorden, Spitsbergen. The active glacier is 35 km² in area, 11 km in length and is polythermal [Wadham et al., 1998]. The lithology is mainly sedimentary, with the upper catchment and headwalls consisting of Precambrian carbonates, phyllite and quartzite, Permian sandstones, dolomite and limestones and the rest of the catchment comprising Triassic to Cretaceous siltstones, sandstones and shales [Dallmann et al., 1990].

[6] Details of meltwater drainage are discussed previously by Wadham et al. [1998, 2000, 2004, 2007], and indicate that ~70% of the glacier’s meltwaters drain from the western margin via the SGU and IMC. The relative dominance of these two pathways altered markedly between 1994 and 1999, and a summary of this behavior is included in Table 1. We subdivide the 5 years of study into two periods based upon the principle route taken by subglacial meltwaters: the Artesian Drainage Period (AD period) (1994–1997), when the SGU contributed subglacial meltwaters to runoff on the western margin and the Channelized Drainage Period (CD period) (1999), when the SGU was no longer active and all
runoff was routed to the margin by channels, culminating in the IMC.

3. Methods

[7] Full details of sampling procedures for individual years have been described previously (see Table 1 for source). In brief, manual samples were collected daily from the SGU in 1994 (Days 222–231) and 1995 (Days 177–221) and twice daily in 1997 (Days 183–195). No runoff sampling was conducted in 1996 or 1998. The 1997 SGU samples can also be considered as bulk runoff samples, since the SGU dominated western margin drainage in this year. Daily bulk runoff samples were taken in 1999, which issued from the IMC following SGU evolution. Samples were analyzed for the major cations and anions by ion chromatography on a Dionex 4000i Ion Chromatograph, where precision and accuracy of analyses was ±2 and ±10%, respectively. Alkalinity was calculated from charge balance. Dissolved Si was determined colorimetrically following methods by Wadham et al. [2000] and precision and accuracy were ±3.4% and ±6.7%, respectively. Isotopic determinations on 1997 samples are described by Wadham et al. [2004]. The isotopic composition of sulphate in 1999 runoff was determined following these methods and those reported by Wadham et al. [2007]. In short, the $\delta^{18}\text{O}$-SO$_4^{2-}$ of the samples was determined by CO$_2$ equilibration (standard = SMOW). The sulphate in these samples was recovered as BaSO$_4$ for the analysis of $\delta^{34}\text{S}$ and $\delta^{18}\text{O}$-SO$_4^{2-}$ (standard = CDT and SMOW, respectively). Isotopic analyses were made on a VG SIRA10 isotope ratio mass spectrometer at the University of Leeds. Methods for bulk discharge monitoring in 1994, 1995 and 1999 can be found in the works by Wadham et al. [1998] and Cooper et al. [2002].

4. Results

4.1. Major Ion Chemistry

[8] Major ion chemical data are available in all years, with $\delta^{34}\text{S}$ and $\delta^{18}\text{O}$ isotopes on sulphate for 1997 and 1999 (Table 1). Cation and sulphate concentrations are corrected for the snowpack contribution using ion ratios to chloride as reported previously [Wadham et al., 1998, 2004, 2007]. Hence, all concentrations reported are derived from rock weathering alone.

[9] We employ the concentrations of a variety of chemical indices in order to infer forcing mechanisms for hydrological evolution and associated shifts in biogeochemical processes during the AD period and between the AD and CD periods. Specifically, we conduct three comparisons, described below, to facilitate this analysis: (1) hydrological forcing, (2) SGU biogeochemical evolution, and (3) AD to CD biogeochemical evolution.

[10] 1. We employ discharge data and Cl$^-$ concentrations in meltwaters to indicate hydrological forcing for the drainage reorganization observed during the 5 year study period. Cl$^-$ is derived from snowmelt, and is generally absent from ice-melt [Wadham et al., 1998]. Hence, interseasonal differences in Cl$^-$ are used to indicate contrasts in these two meltwater sources during and between the AD and CD periods. We also compare total bulk discharge variations and the relationship between Cl$^-$ and discharge for 1994 and 1995 (AD period) and 1999 (CD period) in order to infer the degree to which any interannual differences in meltwater sourcing might be a function of varying surface melt rates (and hence discharge).

[11] 2. The chemical composition of meltwaters issuing from the SGU between 1994 and 1997 is compared in order to identify any trends in subglacial biogeochemistry during the AD period. Specifically, we compare associations between the major crustal-derived ions and ion ratios.

[12] 3. We compare the chemistry of the western margin bulk runoff between 1997 (using data from the SGU, which dominated bulk discharge) and 1999 (when the SGU was absent and the IMC dominated discharge) in order to determine biogeochemical evolution between the CD and the AD periods. We do not present the full crustal chemical data for bulk runoff in 1994 and 1995, since chemical trends in these years mainly reflect mixing processes and the relative contributions of the SGU and IMC to total bulk discharge [Wadham et al., 1998]. Associations between bulk meltwater *SO$_4^{2-}$ concentrations and discharge in 1994–1995 and 1999 are employed purely to indicate changes in these mixing processes.

4.2. Hydrological Forcing

[13] Seasonal time series of bulk meltwater discharge from the western margin in 1994, 1995 (AD period) and 1999 (CD period) are displayed in Figure 1. The bulk discharge regime in 1999 differs fundamentally from that in 1994 and 1995 in two significant ways. First, there is a period of unusually high spring discharge in 1999 of up to 30 m$^3$ s$^{-1}$, compared with 1994 and 1995 when early season discharges were generally <5 m$^3$ s$^{-1}$. Second, the occurrence of a midseason outburst event in 1999 [Wadham et al., 2001] produces peak discharges double those in 1994 and 1995. These outburst events are commonly observed on the western margin of the glacier in summer and have been discussed previously [Wadham et al., 2001].

[14] Figure 2a displays the temporal variation in Cl$^-$ concentrations in SGU meltwaters during the AD period, and Figure 2b compares Cl$^-$ concentrations between bulk
runoff sampled during the AD period (1997) and CD period (1999), when the SGU and IMC, respectively, dominated discharge at the western margin. There is a very clear trend of decreasing Cl\(^{-}\) in all waters sampled from the beginning to the end of each monitoring period. This is matched by decreasing concentrations of Cl\(^{-}\) in SGU meltwaters from 1994 to 1997 and in bulk runoff from 1997 to 1999. The association between bulk meltwater Cl\(^{-}\) concentrations and bulk meltwater discharge for the AD period (1994, 1995) and CD period (1999) is displayed in Figures 2c and 2d. While seasonal decreases in Cl\(^{-}\) generally correlate with increasing discharge, interseasonal concentrations of Cl\(^{-}\) between the AD and CD periods cannot be explained by discharge alone. For example, bulk meltwater Cl\(^{-}\) concentrations in the CD period (1999) are lower than those from the AD period (1994–1997) for the same discharge. During the AD period, there is also a decrease in bulk meltwater Cl\(^{-}\) concentrations for the equivalent discharge from 1994 to 1995. There is no significant relationship between Cl\(^{-}\) and discharge in 1999. As reported elsewhere, some of the higher discharge points (>15 m\(^3\) s\(^{-1}\)) in this year refer to a subglacial outburst event [Wadham et al., 2001].

### 4.3. Biogeochemical Evolution

[15] The concentration of sulphate, derived from the oxidation of sulphide minerals [Wadham et al., 2004], displays similar trends to Cl\(^{-}\) both within and between individual years (Figures 3a and 3b). A notable exception is the low *SO\(_4^{2-}\) observed in the 1997 AD period meltwaters relative to all other samples. In general, the bulk meltwater SO\(_4^{2-}\) concentrations were higher and much more strongly correlated with bulk meltwater discharge during the AD period (1994 and 1995: R\(^2\) = 0.72 and 0.59, respectively) relative to the CD period (1999: R\(^2\) = 0.41). As for Cl\(^{-}\), relatively constant *SO\(_4^{2-}\) concentrations in 1999 data for some of the discharge points >15 m\(^3\) s\(^{-1}\) reflects a subglacial outburst event [Cooper et al., 2002].

[16] Associations between different ions can be used to infer the dominant chemical weathering processes in meltwaters during and between the AD and CD periods. Associations between the major ions and dissolved Si are presented in Figure 4, with the gradients, intercepts and R\(^2\) values for these associations described in Table 2. We remove the ion data associated with the subglacial outburst event in 1999, which briefly perturbed bulk meltwater solute chemistry. Plots presented in Figure 4 highlight distinct evolutionary trends both during the AD period and between the AD and CD periods. Relationships between *SO\(_4^{2-}\) and HCO\(_3^-\) are generally strong during the AD period (apart from 1994, where the small number of data points and the low gradient makes this relationship statistically insignificant) (Figure 4a). There is no relationship between HCO\(_3^-\) and *SO\(_4^{2-}\) during the CD period. This suggests that there was a change in the biogeochemical dissolution processes supplying sulphate...
from the AD period to the CD period. Intercepts in plots of HCO$_3^-$ against SO$_4^{2-}$ and Ca$^{2+}$+Mg$^{2+}$ against SO$_4^{2-}$ in all meltwaters sampled are high (Figures 4a and 4b) and exceed the theoretical value for carbonate hydrolysis ($= 220 \mu$eq L$^{-1}$) [Tranter et al., 2002].

Slopes of associations between HCO$_3^-$ and SO$_4^{2-}$ (Figure 4a) and Ca$^{2+}$+Mg$^{2+}$ and SO$_4^{2-}$ (Figure 4b) are low during the AD period (0.28 and 0.66 for HCO$_3^-$/SO$_4^{2-}$ in 1994 and 1997, respectively, and 1.28 and 1.34 for Ca$^{2+}$+Mg$^{2+}$/SO$_4^{2-}$ in 1994 and 1997, respectively), while concentrations of Na$^+$ and K$^+$ are high (Figures 4d and 4e). Both these features are consistent with the dissolution of silicate minerals by protons generated during sulphide oxidation [Tranter et al., 2002]. The 1995 SGU waters represent an enigma here since they display the highest concentrations of Na$^+$ and K$^+$ of all waters sampled (Figure 4e), but have high slope values in both relationships of HCO$_3^-$ against SO$_4^{2-}$ (gradient = 1.4) and Ca$^{2+}$+Mg$^{2+}$ against SO$_4^{2-}$ (gradient = 1.82) (Table 2). The regression line for the association between SO$_4^{2-}$ and Ca$^{2+}$+Mg$^{2+}$ for the 1999 bulk meltwaters was lower (slope = 1 in 1999) (Table 2) than those for the SGU meltwaters in all years, as determined by ANCOVA one way tests (at the 99% confidence level).

Decreases in the concentrations of Na$^+$ and K$^+$ from 1994/1995 to 1997 during the AD period (Figure 4e) suggest a diminishing importance of silicate dissolution as the SGU evolves hydrologically. Na$^+$ and K$^+$ display slightly higher concentrations in bulk runoff from the SGU compared to the CD period (Figure 4e). The relationship between Na$^+$ + K$^+$ concentrations and HCO$_3^-$ (Figure 4e) is markedly stronger than that with SO$_4^{2-}$ for most waters sampled (Figure 4d), suggesting that Na$^+$ and K$^+$ are principally acquired with HCO$_3^-$, rather than SO$_4^{2-}$.

Associations between ratios of (Ca$^{2+}$+Mg$^{2+}$): (Na$^+$+K$^+$), K$^+$:Na$^+$ and Ca$^{2+}$:Mg$^{2+}$ (Figure 5) and ionic strength can be used to infer the changing dominance of chemical dissolution processes with changing residence time during and between the AD and CD periods. Whereas divalent ions are usually derived from carbonate weathering (carbonate and dolomite), Na$^+$ and K$^+$ are derived predominantly from the dissolution of silicate minerals. Hence, ratios of Ca$^{2+}$+Mg$^{2+}$:(Na$^+$+K$^+$) can be used to indicate the relative dominance of carbonate versus silicate dissolution. K$^+$ derives predominantly from the dissolution of biotite, released nonstoichiometrically from the bedrock during subglacial comminution [Anderson, 2007]. Thus, ratios of K$^+$:Na$^+$ can be used to indicate the contribution to total runoff by meltwaters draining areas of the glacier bed where there is active comminution (e.g., distributed system). Ratios of Ca$^{2+}$+Mg$^{2+}$ indicate the relative importance of dolomite verses carbonate dissolution, since a proportion of Mg$^{2+}$ released is greater by dolomite dissolution. Typically, the
proportion of dolomite dissolution increases with increased meltwater residence time [Sharp et al., 2002]. Figure 5a indicates that there is a progressive increase in the ratio of divalent to monovalent ions with decreasing ionic strength and moving from the AD to CD period. Comparison of bulk meltwaters from the SGU (1997) and IMC (1999) periods indicates higher ionic strength and lower ratios of \(*\text{Ca}^{2+} + *\text{Mg}^{2+} : *\text{Na}^+ + *\text{K}^+\) in the former case. Ratios of \(*\text{K}^+ : *\text{Na}^+\) and \(*\text{Ca}^{2+} + *\text{Mg}^{2+} : *\text{SO}_4^{2-}\) show similar trends with ionic strength for the different years and periods (Figures 5a and 5b).

[10] Data on the $\delta^{34}$S and $\delta^{18}$O composition of sulphate exist for bulk runoff from the AD (1997) and CD (1999) periods, when the SGU and IMC, respectively, dominated runoff. The 1997 data are discussed by Wadham et al. [2004]. The $\delta^{34}$S values during the AD period plot in a steep array, with the $\delta^{34}$S inverse to the sulphate mass fraction of \(\text{SO}_4^{2-} : (\text{SO}_4^{2-} + \text{HCO}_3^-)\) (hereafter “SMF”).
Figure 6a). The lower end of this range corresponds to values typical of bedrock sulphide at $-17.94$‰ [Wadham et al., 2007]. Although the $\delta^{34}$S of CD runoff plots across the same range as that for the AD period, there is no significant association with SMF in the former case. There is also no temporal trend in $\delta^{34}$S values from the IMC (time series not shown). The $\delta^{18}$O of sulphate in the SGU meltwaters displays consistent values of around $+25–27$‰ (Figure 6b), which are about 35‰ heavier than CD meltwaters. This signifies a major shift in the isotopic

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composition of sulphate in meltwaters between the two periods.

5. Discussion

5.1. Interannual Changes in Hydrological Forcing Regime

[20] Previous research based on 1994 and 1995 data has inferred that the SGU drains a snowmelt-fed subglacial distributed drainage system, involving meltwater flow via a basal debris layer [Wadhams et al., 1998]. Decreases in Cl concentrations (Figure 2a) in the SGU between 1994 and 1997 provide clear evidence for strengthened coupling between the SGU and glacier surface icemelt production over this time period, attaining full coupling in 1997 when virtually all meltwater on the western margin was routed by the SGU. We note that rainfall during the study seasons was minimal (27–29 mm) compared with the amount of runoff produced by ice melt [Wadhams et al., 1998; Cooper et al., 2002]. Hence, it is unlikely that rainfall was the driver of the hydrological evolution. We infer that relatively low, but steadily rising water fluxes (via increased icemelt) to the SGU between 1994 and 1997 (Figure 1) enabled the SGU to be maintained on the western margin over this time period. Gradual rising fluxes resulted in a concomitant enlargement of flow paths and decrease in SGU water pressures in the near terminal zone, resulting in drainage capture from the IMC. This sequence of events matches well to hydrological theory, which states that increased discharge is accompanied by decreasing water pressures in conduits, and that larger channels should capture drainage from smaller ones if hydrological connections exist [Röthlisberger, 1972]. Similar phenomena have been observed on the polythermal glacier, Erdmanbreen, Svalbard during the course of a single melt season, where rising discharges were accommodated increasingly by a subglacial upwelling, at the expense of drainage via ice-marginal channels [Hodson and Fergusson, 1999].

[21] If the progressive meltwater capture by the SGU from 1994 to 1997 is a result of falling water pressures in this environment compared to the IMC, the reversal of this process from 1997 to 1999 must reflect increasing pressures in the SGU relative to the IMC and the forcing of water into the latter drainage system. The high short-duration spring runoff event in 1999 (Figure 1) is held as a significant driver for this evolutionary behavior. This could have triggered drainage migration (e.g., by hydrofracture through cold ice), toward the more favorable (at this time) lower-pressure channelized system (IMC) [Röthlisberger, 1972], as is observed on temperate valley glaciers during discharge increases [Nienow et al., 1998]. Further increases in melt input during the 1999 melt season (maximum discharge = 46 m$^3$ s$^{-1}$) and strengthening of surface-to-bed hydrological connectivity, as indicated by lower Cl$^-$ concentrations, are inferred to have sustained this channelized routing of meltwater for the rest of the melt season. Whether the chemical weathering environment that supplied the SGU during the AD period contributed meltwaters to the IMC during the CD period can only be inferred from the crustal ion data, and will be conducted in subsequent discussion. We contend that the SGU was sustained as the dominant drainage feature on the western margin from 1994 to 1997 since the melt inputs in spring were small (Figure 1) and increased slowly, enabling the SGU to adjust its configuration to the rising water fluxes such that only moderate water pressures were attained. The frequent observation of pressurized subglacial upwellings on Svalbard glaciers, which shift their location as a response to frontal collapse in the glacier terminal region and changing melt rates is consistent with these assertions [Hodson and Fergusson, 1999; Hodson et al., 2000; Rippin et al., 2003; Wynn et al., 2006]. In particular, high spring melt rates and elevated pressures in subglacial upwellings were inferred by Rippin et al. [2003] to drive diversion of drainage from the upwellings to an ice-marginal/subglacial channel on another Svalbard glacier, Midtre Lovénbreen.

[22] The fact that Cl$^-$ concentrations decrease only slightly between 1997 (AD period) and 1999 (CD period) (Figures 2a and 2b) on Finsterwalderbreen, but ion associations are dramatically different (Figure 4) also supports the notion that changes observed in runoff biogeochemistry from the AD to the CD period were not simply a consequence of dilution but
reflect fundamental changes in drainage system configuration (i.e., predominantly distributed drainage to predominantly channelized drainage). This sequence of events on Finsterwalderbreen provides a unique case study for identifying modes of coupling between subglacial drainage reorganization and biogeochemistry, as is discussed in section 5.2.

5.2. Identification of Coupling Between Subglacial Hydrology and Bed Biogeochemistry During Drainage Reorganization

[23] The reorganization of the drainage system on Finsterwalderbreen associated with the growth and disappearance of the SGU between 1994 and 1999 has a direct impact on biogeochemical processes in the Finsterwalderbreen catchment. Most strongly evident are the decreases in the concentrations of crustal ions in SGU meltwaters between 1994 and 1997, and particularly \( \text{SO}_4^{2-} \) (Figures 3b and 3d), signifying a reduction in residence times and rock:water ratios during SGU growth and meltwater capture from the IMC. The much smaller crustal ion decreases in bulk meltwaters from 1997 (AD period) to 1999 (CD period) (Figures 3 and 4) suggest similar residence times during these two time periods. However, dramatic shifts in ion associations (Figure 4) indicate different drainage flow path characteristics between the two periods, and hence chemical weathering processes. The details of this biogeochemical evolution are most clearly identified from associations between ions and from isotopic data as identified in the following discussion.

[24] Sulphate is commonly employed as a diagnostic indicator of distributed type drainage beneath glaciers, where reactive sulphide surfaces are continually produced in subglacial environments by comminution [Tranter et al., 1993]. The generally strong associations between \( \text{SO}_4^{2-} \) and \( \text{HCO}_3^- \) (Figure 4a) in the SGU during the AD period are consistent with strong coupling between carbonate dissolution and sulphide oxidation in this subglacial chemical weathering environment, where the following reaction applies:

\[
4\text{FeS}_2 + 16\text{CaCO}_3 + 15\text{O}_2 + 14\text{H}_2\text{O} \rightleftharpoons 16\text{Ca}^{2+} + 16\text{HCO}_3^- + 8\text{SO}_4^{2-} + 4\text{Fe(OH)}_3
\]  

(1)

The relatively high concentrations of \( \text{SO}_4^{2-} \) in the 1999 (CD period) meltwaters and lack of any strong relationship between \( \text{SO}_4^{2-} \) and \( \text{HCO}_3^- \) (Figure 4a) implies that, if \( \text{SO}_4^{2-} \) derives from sulphide oxidation, the protons generated by this process do not dissolve carbonates. However, the lack of any correlation between \( \text{SO}_4^{2-} \) ("Na" + K") and \( \text{SO}_4^{2-} \) (Figure 4d) suggests that neither do protons from sulphide oxidation dissolve silicates. This leads us to infer that the provenance of \( \text{SO}_4^{2-} \) in 1999 runoff is different to that of the SGU during the AD period. We contend that \( \text{SO}_4^{2-} \) in 1999 runoff derives from ice-marginal moraine pore waters, which are either entrained directly by ice-marginal channel waters and/or added to the IMC via the numerous extraglacial streams that can be observed along the western glacier margin. Previous work shows that these pore waters are rich in \( \text{SO}_4^{2-} \), which is acquired initially by sulphide oxidation in moraines, but subsequently precipitates from solution as \( \text{Ca}^{2+}-\text{Mg}^{2+}-\text{SO}_4^{2-} \) salts due to freeze and evaporative concentration in summer [Cooper et al., 2002]. The salts redissolve during summer rain events, producing high \( \text{Ca}^{2+}-\text{SO}_4^{2-} \) concentrations in moraine pore waters [Cooper et al., 2002]. These pore waters are typically saturated with respect to carbonate, and hence contain low concentrations of \( \text{HCO}_3^- \) [Cooper et al., 2002]. The hypothesis that \( \text{SO}_4^{2-} \) in 1999 runoff is predominantly derived from the addition of \( \text{SO}_4^{2-} \)-rich moraine pore waters is supported by the poor relationships between \( \text{HCO}_3^- \) and both \( \text{SO}_4^{2-} \) and \( \text{Ca}^{2+}-\text{Mg}^{2+} \) in 1999 runoff, but a strong relationship between \( \text{Ca}^{2+}-\text{Mg}^{2+} \) and \( \text{SO}_4^{2-} \) (Figure 4). The much lower gradient (1.1) in plots of \( \text{SO}_4^{2-} \) versus \( \text{SO}_4^{2-} \) (Figure 4b) compared with SGU meltwaters is consistent with the known composition of moraine pore waters, where \( \text{Ca}^{2+} \) and \( \text{SO}_4^{2-} \) occur in ratios of \( 1:1 \) [Cooper et al., 2002; Wadham et al., 2007]. The strong ice-marginal influence in the CD period meltwaters matches with observations on cold-based glaciers, where contact of runoff with ice-marginal moraines contributes significant solute, and particularly \( \text{SO}_4^{2-} \), to bulk runoff [Tranter et al., 2002]. It suggests that during the CD period continuously draining, long-residence-time subglacial meltwaters on the western margin were either absent or very limited.

[25] Coincident with a shift in the dominant sulphate source between the AD and CD periods are changes in the redox chemistry of runoff, as displayed by the \( \delta^{18}\text{S} \) and \( \delta^{18}\text{O} \) composition of sulphate in 1997 and 1999 bulk meltwaters (Figure 6). These isotopes are well known to provide information upon oxic versus anoxic sulphide oxidation, in addition to sulphate reduction in glaciers [Bottrell and Tranter, 2002; Wadham et al., 2004, 2007]. The degree of anoxia determines which of these three processes control the isotopic composition of the sulphate. Under oxic conditions, sulphides are oxidized by molecular oxygen (equation 2)), reverting to Fe(III) as conditions go suboxic to anoxic (equation 3)). As anoxia increases still further, and electron acceptors such as oxygen, nitrate and Fe(III) become depleted, sulphate reduction becomes viable, provided an organic carbon source exists (equation 4)). Oxidation of sulphate in an oxic environment should produce \( \delta^{18}\text{S}-\text{SO}_4^{2-} \) values close to that of the bedrock sulphide (mean = \(-17\%o \) [Wadham et al., 2007] and an \( \delta^{18}\text{O}-\text{SO}_4^{2-} \) of \(+11\%o \), derived from a mixture of 3.5 of 4 oxygen atoms from \( \text{O}_2 \) \(+23.7\%o \) and 0.5 from water molecules \( +12 \) to \(-15\%o \), according to equation (2) [Dole et al., 1954; Kroopnick and Craig, 1972; Lloyd, 1967; Taylor et al., 1984; van Everdingen and Krouse, 1985; Holt and Karm, 1991]):

\[
\begin{align*}
\text{FeS}_2 + 3.5\text{SO}_2 + 16\text{H}_2\text{O} & \rightleftharpoons 2\text{SO}_4^{2-} + 2\text{Fe}^{2+} + 2\text{H}^+ \\
\text{FeS}_2 + \text{Fe}^{3+} + 8\text{H}_2\text{O} & \rightleftharpoons 2\text{SO}_4^{2-} + 2\text{Fe}^{2+} + 16\text{H}^+ \\
\text{SO}_4^{2-} + 2\text{CH}_2\text{O} & \rightleftharpoons \text{H}_2\text{S} + 2\text{HCO}_3^- 
\end{align*}
\]

(2) (3) (4)

[26] Sulphate in neither the 1997 (AD period) nor the 1999 (CD period) bulk meltwaters displays an isotopic composition in line with that for oxic sulphide oxidation (Figure 6). The isotopic composition of sulphate during the AD period is consistent with the occurrence of bacterial sulphate reduction, where the consistency of \( \delta^{18}\text{O} \) at around \(+25\%o \) is evidence that bacterial sulphate reduction has attained isotopic equilibrium [Wadham et al., 2004]. Here,
the residual sulphate is enriched in $\delta^{34}$S and $\delta^{18}$O, since the lighter isotope is preferentially incorporated into the $\mathrm{H}_2\mathrm{S}$ reaction product (equation (4)). Sulphate reduction in the SGU in 1997 provides one explanation for the low $\delta^{34}$SO$_4^-$ concentrations observed in this year, and their similarity to 1999 despite the higher concentrations of all other ions (Figure 4).

[27] There is no evidence for bacterial sulphate reduction in the CD period (1999) bulk runoff. Here, the $\delta^{18}$O-SO$_4^-$ is consistent with predominantly anoxic sulphide oxidation using Fe(II) (equation (3)), where the $\delta^{18}$O-SO$_4^-$ adopts a value close to that of environmental water [Bottrell and Tranter, 2002], −12 to −12.9‰ in this case.

[28] During the CD period, $\delta^{18}$O in SO$_4^-$ (mean = −8.9‰ in this case) is slightly higher than environmental water (Figure 6b), suggesting that a small proportion of sulphides are being oxidized under oxic conditions. Isotope mass balance can be used to calculate the mean percentage contribution of oxic versus anoxic sulphide oxidation to sulphate in the 1999 meltwaters in the following manner,

$$
\langle \delta^{18}\text{O-SO}_4^- \times \text{SO}_4^- \rangle_{\text{bulk}} = \langle \delta^{18}\text{O-SO}_4^- \times \text{SO}_4^- \rangle_{\text{oxic}} + \langle \delta^{18}\text{O-SO}_4^- \times \text{SO}_4^- \rangle_{\text{anoxic-Fe(II)}}
$$

where SO$_4^-$ is sulphate concentration and $\delta^{18}$O-SO$_4^-$ is the isotopic composition of the sulphate. Hence, $\langle \delta^{18}$O-SO$_4^-$ \times \text{SO}_4^- \rangle_{\text{bulk}}$ refers to the sample composition as measured. We set the $\langle \delta^{18}$O-SO$_4^-$ \times \text{SO}_4^- \rangle_{\text{anoxic-Fe(II)}}$ to the mean $\delta^{18}$O-H$_2$O recorded in proglacial meltwaters (−12.5‰). We calculate the $\langle \delta^{18}$O-SO$_4^-$ \times \text{SO}_4^- \rangle_{\text{oxic}}$ as described above, taking −12.5‰ as the mean $\delta^{18}$O-H$_2$O and assuming that 3.5 of 4 oxygen atoms in the sulphate produced derive from the atmosphere ($\delta^{18}$O = 23.7‰) and 0.5 from water (gives +11.6‰ for $\delta^{18}$O-SO$_4^-$). Equation (5) is then solved to give the mean concentrations of SO$_4^-$ produced oxicly and anoxically, since (SO$_4^-$)$_{\text{bulk}}$ = (SO$_4^-$)$_{\text{oxic-Fe(II)}}$ + (SO$_4^-$)$_{\text{oxic}}$. Calculations give values of 15% and 85%, respectively, and are consistent with a predominantly anoxic source for most sulphate in these meltwaters.

[29] The $\delta^{18}$O-SO$_4^-$ and $\delta^{34}$S-SO$_4^-$ composition of CD period (1999) meltwaters plot within the range reported for proglacial moraine groundwaters [Wadhams et al., 2007], with a scatter in $\delta^{34}$S-SO$_4^-$ that arises from variation in the initial $\delta^{34}$S bedrock composition [Wadhams et al., 2007]. This is consistent with the hypothesis that most sulphate in 1999 runoff derives from the addition of moraine pore waters to the IMC. Typically 88% of sulphate is produced anoxically using Fe(II), and 12% is acquired under oxic conditions in moraine pore waters [Wadhams et al., 2007], which is in broad agreement with the values calculated above.

[30] Isotope data are not available prior to 1997 and so it is impossible to infer anoxic versus oxic conditions in the SGU chemical weathering environment in 1994 and 1995. The data for 1997, however, strongly suggest that the SGU drained an anoxic environment where sulphate reduction was prevalent. We infer that the contribution to bulk runoff by long-residence-time subglacial drainage was small or absent following drainage reorganization between 1997 and 1999 (apart from during outburst events). Hence, during the CD period sulphate was acquired initially via anoxic sulphide oxidation in ice-marginal moraines, with temporary storage as sulphate salts that subsequently became mobilized by melt and input to the ice marginal channelized environment. This shift from microbially mediated sulphate reduction to sulphide oxidation under anoxic conditions implies a major change in the biogeochemical weathering environment draining the glacier western margin.

[31] Further information on microbial processes is encapsulated in the intercept values in plots of both HCO$_3^-$ against *SO$_4^-$ and *Ca$^{2+}$+*Mg$^{2+}$ against *SO$_4^-$ (Figures 4a and 4b and Table 2). Intercepts of ~220 µeq L$^{-1}$ are commonly reported for subglacial and other meltwaters, and reflect the acquisition of Ca$^{2+}$+Mg$^{2+}$ and HCO$_3^-$ in the absence of SO$_4^-$, by carbonate hydrolysis (equation (6)) [Tranter et al., 2002]:

$$
\text{CaCO}_3 + \text{H}_2\text{O} \Leftrightarrow \text{Ca}^{2+} + \text{HCO}_3^- + \text{OH}^-
$$

[32] Increases above the 220 µeq L$^{-1}$ intercept value in AD and CD period meltwaters point to an additional source of HCO$_3^-$. In addition, the increases in the gradient of the association between HCO$_3^-$ and *SO$_4^-$ in 1995 meltwaters to above the maximum theoretical value for sulphide oxidation/ carbonate dissolution (gradient = 1) suggest that some of this additional HCO$_3^-$ is linked to processes involving SO$_4^-$ (Table 2). The SGU is out of contact from the atmosphere, precluding an atmospheric source for excess HCO$_3^-$ between 1994 and 1997. The most plausible scenario in this case is that the HCO$_3^-$ is derived from CO$_2$ produced via microbial respiration, which subsequently fuels chemical dissolution in a subglacial debris layer. Similar hypotheses have been used to explain excess HCO$_3^-$ in borehole waters from temperate valley glaciers [Tranter et al., 1996]. In 1995, the gradient of >1 for the association between HCO$_3^-$ and SO$_4^-$ (Table 2) may reflect bacterial sulphate reduction (and HCO$_3^-$ generation via the oxidation of organic carbon) in the very long residence time, low-discharge waters. A decrease in sulphate reduction as discharges rises in the SGU would reduce the excess HCO$_3^-$ added to meltwaters by this process, steepening the overall gradient of the HCO$_3^-$/SO$_4^-$ relationship. Essentially this is a drainage evolution trend.

[33] Since meltwaters follow a channelized/ice-marginal routing to the glacier terminus during the CD period (1999), an atmospheric source for the excess HCO$_3^-$ is plausible. One feature of the 1999 chemical data set, however, is inconsistent with this hypothesis. The strong relationships between *Na$^+$+*K$^+$ and HCO$_3^-$ in AD and CD period meltwaters (and weak associations between these cations and *SO$_4^-$ (Table 2)) suggest that some of the excess HCO$_3^-$ is balanced by *Na$^+$ and *K$^+$. The significant associations between dissolved Si and HCO$_3^-$ in 1995 and 1999 (Table 2), suggest that this is not simply an ion exchange phenomenon where Ca$^{2+}$ and Mg$^{2+}$ (acquired by CO$_2$, and balanced by HCO$_3^-$ in solution) are exchanged for K$^+$ and Na$^+$ on sediment exchange sites. The inverse association between the ratio of (Ca$^{2+}$+Mg$^{2+}$):(*Na$^+$+*K$^+$) and isotonic strength (Figure 5a) is also consistent with this assertion. This relationship is opposite to what might be anticipated for ion exchange, where divalent ions are exchanged for monovalent ions as exchange surfaces become more selective at lower ionic strength [Tranter et al., 2002]. We note that this is contrary to
the original interpretation for the high excess \( *\text{Na}^{+} \) observed in SGU waters, when a smaller amount of data were available [Wadhams et al., 1998]. If our current interpretation of the data is correct, some of the excess \( \text{CO}_2 \) observed in meltwaters must directly dissolve silicates (up to 42% in 1995 as indicated by gradients of \( *\text{Na}^{+} + *\text{K}^{+} \) versus \( \text{HCO}_3^{-} \) (Table 2)). \( \text{CO}_2 \) derived from an atmospheric source is very unlikely to fuel the dissolution of silicate minerals. Unless the \( \text{CO}_2 \) is generated close to silicate mineral weathering sites, the unfavorable kinetics of silicate dissolution [Tranter et al., 1993] will favor the carbonation of carbonates rather than silicates, according the following equations:

\[
\text{Ca}_{1-x} (\text{Mg}_x) \text{CO}_3 + \text{CO}_2 + \text{H}_2 \text{O} \Leftrightarrow (1-x) \text{Ca}^{2+} + x \text{Mg}^{2+} + 2\text{HCO}_3^{-} (\text{Mg})
\]

(7)

\[
\text{CaAl}_2\text{Si}_2\text{O}_8 + 2\text{CO}_2 + 2\text{H}_2 \text{O} \Leftrightarrow \text{Ca}^{2+} + 2\text{HCO}_3^{-} + \text{H}_2\text{Al}_2\text{Si}_2\text{O}_8
\]

(8)

where \( \text{Ca}_{1-x} (\text{Mg}_x) \text{CO}_3 \) is magnesium calcite, \( \text{CaAl}_2\text{Si}_2\text{O}_8 \) is anorthite (calcium feldspar), and \( \text{H}_2\text{Al}_2\text{Si}_2\text{O}_8 \) is weathered feldspar surfaces.

[34] This suggests that the \( \text{CO}_2 \) was produced close to the sites of mineral dissolution, and most probably by microbial respiration. There are numerous reports in the literature where microorganisms accelerate the dissolution of a variety of silicates by the localized production of excess protons and organic ligands [Bennett et al., 2001; Liermann et al., 2000; Drever and Stilings, 1997; Brantley and Stilings, 1996]. In doing so, they derive macronutrients (e.g., N and P) and trace nutrient metals such as Fe, V and Mn [Valsami-Jones et al., 1998; Brantley et al., 2001]. Recent laboratory studies have shown that subsurface microorganisms preferentially colonize nutrient-bearing silicate surfaces and subsequently dissolve the silicate matrix in order to extract and use limiting inorganic nutrients [Rogers et al., 1998]. The \( *\text{Na}^{+} + *\text{K}^{+} \) and \( \text{HCO}_3^{-} \) data presented here provide evidence that similar processes may operate in a subglacial and ice-marginal environment, and in doing so accelerate chemical weathering rates. These assertions are consistent with microscopy studies which have detected high concentrations of microbial cells attached to clay grains in basal ice from the Greenland Ice Sheet, coincident with high concentrations of carbon dioxide and methane gas [Tung et al., 2006].

[35] The relative dominance of silicate versus carbonate dissolution in smaller valley glaciers, has been linked previously to meltwater residence times, such that silicate dissolution is enhanced in long-residence-time meltwaters as carbonates become exhausted or saturation is attained with respect to calcite [Sharp et al., 2002]. Given the inferred decrease in residence times and rock:water ratios for western margin meltwaters between the AD and CD periods, similar shifts in the balance between silicate and carbonate dissolution might be expected on our larger-scale system. Decreasing concentrations of \( *\text{Na}^{+} \) and \( *\text{K}^{+} \) between 1994 and 1997 to the lowest measured concentrations in 1999 (Figures 4d and 4e) suggest a reduction in the extent of silicate weathering as the upwelling captures an increasing volume of meltwater during the AD period and then evolves to a channelized drainage system during the CD period. Gradients of associations between \( \text{HCO}_3^{-} \) and \( *\text{SO}_4^{2-} \) in SGU meltwaters from 1994 and 1997 fall below the theoretical value of 1 for coupled sulphide oxidation/carbonate dissolution, indicating coupling between sulphide oxidation and silicate dissolution (equation (9)),

\[
4\text{FeS}_2 + 16\text{Na}^{+} + 5\text{AlSi}_3\text{O}_8 + 15\text{O}_2 + 86\text{H}_2\text{O} \\
\Leftrightarrow 16(1-x)\text{Na}^{+} + 16\text{K}^{+} + 8\text{SO}_4^{2-} + 4\text{Al}_2\text{Si}_4\text{O}_{10} + 32\text{H}_2\text{SiO}_4 + \text{Fe(OH)}_3
\]

(9)

where \( \text{Na}_{1-x} \text{K}_x \text{AlSi}_3\text{O}_8 \) is alkali feldspar and \( \text{Al}_2\text{Si}_4\text{O}_{10} \) is kaolinite.

[36] The high slope values in 1995 are an exception here and may reflect drainage evolution trends with increased microbial \( \text{CO}_2 \) generation as discharges fall, as previously discussed. The lack of any significant relationship between \( \text{HCO}_3^{-} \) and \( *\text{SO}_4^{2-} \) or between \( *\text{Na}^{+} + *\text{K}^{+} \) and \( *\text{SO}_4^{2-} \) concentrations (Figures 4a and 4d) during the CD period suggests coupling between sulphide oxidation and silicate dissolution is very limited, if not absent. This is consistent with a decrease in residence times, as the SGU drainage system becomes more efficient and subsequently evolves to a channelized/ice-marginal system in 1999. Correspondingly, the ratio of divalent (carbonate derived) to monovalent (silicate derived) ions relates inversely to ionic strength, and clearly indicates a shift from silicate dissolution (e.g., 1995) to increasing amounts of carbonate dissolution as the SGU develops during the AD period and subsequently evolves to channels between 1997 and 1999 (Figure 4a). This may also reflect the access of meltwaters to new sediment sources as drainage evolution takes place, and new sources of carbonate minerals are tapped by meltwaters and subsequently dissolved. Declining \( *\text{K}^{+}:*\text{Na}^{+} \) ratios with increased ionic strength (Figure 5b) are difficult to link to silicate dissolution, but may reflect loss of \( *\text{K}^{+} \) during the formation of clay minerals in the SGU in 1995 and 1994 [Holland, 1978] when residence times were probably at their peak. Decreasing \( *\text{Ca}^{2+}:*\text{Mg}^{2+} \) ratios with increased ionic strength (Figure 5c) are consistent with previous work which has suggested that dolomite dissolution is enhanced in long-residence-time systems [Sharp et al., 2002]. Hence, as SGU residence times and rock:water ratios decrease with increasing icemelt input, carbonate (and limestone relative to dolomite) dissolution becomes increasingly important over silicate dissolution and clay mineral formation decreases. This situation was most strongly developed during the CD period, when meltwaters took a channelized and subsequently ice-marginal routing to the terminus.

5.3. Evidence for and Quantification of Microbial Mediation of Chemical Weathering

[37] Drainage reorganization between 1994 and 1999 is associated with clear changes in runoff biogeochemistry, with implications for microbial processes and microbial diversity in the Finsterwalderbreen catchment. Sulphate reduction in 1997 (AD period) is superseded by oxic and anoxic sulphide oxidation in 1999 (CD period), signifying a shift from anoxic to oxic/suboxic conditions. All three of these processes are mediated by different types of microorganisms [Wadhams et al., 2004]. In addition, the change from anoxic to suboxic/oxic conditions in the subglacial environment has implications for the composition of microbial (e.g., anaerobes versus...
The role of microbes in mediating chemical weathering is clear during both the AD and CD periods, the role of microbes is apparent in dissolving silicate minerals through the generation of CO$_2$ close to mineral weathering sites. This probably took place within a subglacial debris layer during the AD period and within ice-marginal moraines during the CD period. Concentrations of Na$^+$ and K$^+$ generated by this process decrease as the SGU becomes more efficient and subsequently evolves to a channelized drainage system. This suggests that sectors of the SGU hydrological environment became progressively less favorable for microbial habitation during drainage reorganization.

The process of microbial mediation of sulphide oxidation is well established [Taylor et al., 1984]. We know from the isotopic composition of sulphate in the 1999 runoff that 85% of all sulphide was oxidized anaerobically in the SGU, mediated by microbial respiration (A). This assumes that 386 $\mu$eq L$^{-1}$ of sulphate can be produced by anoxic sulphide oxidation using oxygen supplied in oxygen saturated snowmelt [Tranter et al., 2002], with any excess above this quantity added by anoxic sulphide oxidation. Hence, we subtract this “oxic value” from the total mean sulphate concentration in each water type to give a minimum estimate (C) of the amount of sulphate produced microbially. In the maximum estimate (B), we assume that 100% of the sulphate in the SGU is produced anoxically, mediated by microbes. This assumes that the amount of oxygen delivered to the subglacial environment by regelation (5–6 mm melt per year) is also minimal, which is consistent with previous findings [Wadham et al., 2008]. In the minimum case, we assume that 386 $\mu$eq L$^{-1}$ of sulphate can be produced by anoxic sulphide oxidation using oxygen supplied in oxygen saturated snowmelt [Tranter et al., 2002], with any excess above this quantity added by anoxic sulphide oxidation. Hence, we subtract this “oxic value” from the total mean sulphate concentration in each water type to give a minimum estimate (C) of the amount of sulphate that must be produced anoxically. For both the SGU and 1999 data, we take the concentration of microbially derived sulphate and assume that each molar equivalent of sulphate is balanced by an equivalent quantity of cation charge. Hence values for microbially derived sulphate are multiplied by two to give minimum and maximum estimates of the solute produced microbially from this process.

The first process concerns solute acquired due to microbial CO$_2$ generation. We assume that the CO$_2$ generated by microbial respiration (A) is calculated from the intercepts in plots of HCO$_3^−$ versus $\text{SO}_4^{2−}$ (Table 2) minus the theoretical value for carbonate hydrolysis (220 $\mu$eq L$^{-1}$) [Tranter et al., 2002]. This assumes that all excess CO$_2$ is derived from microbial respiration, rather than the atmosphere; an assumption that is dealt with in section 5.2. We note that the standard error on the 1994 $\text{SO}_4^{2−}$/HCO$_3^−$ association is high (110 $\mu$eq L$^{-1}$) due to the small number of samples. We assume that the excess HCO$_3^−$ concentration is balanced by an equivalent amount of cation charge, which has to be the case for stoichiometric reasons. Hence the excess HCO$_3^−$ is multiplied by two to give the total solute attributable to microbial CO$_2$ production. This method gives a minimum estimate of the excess “microbial” CO$_2$ since it excludes Na$^+$ and K$^+$ which may be acquired by carbonation using microbial CO$_2$. The same calculation is performed for our maximum and minimum estimates.

The second process concerns solute acquired due to microbial mediation of sulphide oxidation. Since we do not know the degree to which microbes are involved in oxic sulphide oxidation within the catchment, we assume for simplicity that only anoxic sulphide oxidation is microbially mediated, which is well established [Taylor et al., 1984]. We quantify the average proportion of solute in meltwaters from 1994 to 1999 which is “microbially generated” by these two processes to give maximum and minimum estimates of microbial solute acquisition (see Table 3, where letter labels in the following text refer to parameter identifiers in the table).

### Table 3. Calculations of Microbially Derived Solute Present During the SGU Meltwaters During the AD period (1994–1997) and Bulk Runoff During the CD Period (1999) at the Western Margin

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Excess Ca$^{2+}$</th>
<th>$\text{Mg}^{2+}$</th>
<th>Maximum Sulphate by Sulphide Oxidation</th>
<th>Minimum Sulphate by Sulphide Oxidation</th>
<th>Maximum Microbial Solute Concentration</th>
<th>Minimum Microbial Solute Concentration</th>
<th>Mean Ion Concentration</th>
<th>%MAXSolute$_{\text{mic}}$</th>
<th>%MINSolute$_{\text{mic}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>SGU 1994</td>
<td>468</td>
<td>475</td>
<td>89.0</td>
<td>1110</td>
<td>2900</td>
<td>65.0</td>
<td>85.0</td>
<td>38.0</td>
<td></td>
</tr>
<tr>
<td>SGU 1995</td>
<td>58.0</td>
<td>604</td>
<td>218</td>
<td>1320</td>
<td>552</td>
<td>33.0</td>
<td>14.0</td>
<td>12.0</td>
<td></td>
</tr>
<tr>
<td>SGU 1997</td>
<td>120</td>
<td>267</td>
<td>0.00</td>
<td>774</td>
<td>240</td>
<td>41.0</td>
<td>13.0</td>
<td>10.0</td>
<td></td>
</tr>
<tr>
<td>1999 Runoff</td>
<td>139</td>
<td>263</td>
<td>0.00</td>
<td>804</td>
<td>278</td>
<td>46.0</td>
<td>16.0</td>
<td>12.0</td>
<td></td>
</tr>
<tr>
<td>Mean SGU</td>
<td></td>
<td></td>
<td></td>
<td>1330</td>
<td>635</td>
<td>47.0</td>
<td>23.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean CDS</td>
<td></td>
<td>804</td>
<td></td>
<td></td>
<td></td>
<td>46.0</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Units are in microequivalents per liter ($\mu$eq L$^{-1}$).*
These calculations show that the amount of solute that is microbially generated in subglacial and ice-marginal environments could be as high as 50–65% of the total solute flux. This supports previous work that has demonstrated a correlation between the metabolic pathways employed by the major phylogenetic groups of microorganisms and runoff chemistry, suggesting that microbes are directly involved in mineral weathering [Skidmore et al., 2005]. It also suggests that microbial processes may play some role in enhancing chemical weathering rates beneath glaciers, which have been shown to exceed the continental average for the same specific discharge [Reynolds and Johnson, 1972; Anderson et al., 1997; Wadham et al., 1997; Hodson et al., 2000].

The impact of SGU hydrological evolution on microbially generated solute is evident in the gradual decrease in the calculated microbially generated solute concentrations between 1994 and 1997, and the decrease in the maximum percentage contribution to total solute from 1994 (65%) to 1995/1997 (39–46%). This suggests that the increased efficiency of the SGU associated with rising icemelt inputs impacted on microbial activity. This may have taken place either by loss of microbial habitats in the subglacial debris layer, or by increasing isolation of these habitats relative to the dominant flow paths as the drainage system became more efficient and concentrated into a smaller number of pathways. Contrary to what one might expect, the mean percentage of solute attributable to microbial activity was similar between the AD period (1997: 47%) and the CD (1999: 46%), suggesting that although the shift in drainage between the two periods lowered overall concentrations of microbially derived solute (Table 3, column D), the proportion of solute that can be said to be microbially derived did not change substantially (Table 3, column G). This suggests that debris rich glacial environments (ice-marginal or subglacial) both constitute favorable environments for microbial activity, as has been hypothesized elsewhere [Tranter et al., 2005].

5.4. Implications for Biogeochemical Processes Beneath Ice Sheets

The AD and CD data sets presented here provide a unique opportunity to infer biogeochemical processes beneath continental-scale ice sheets in two specific ways. First, the growth and evolution of the SGU between 1994 and 1997 enable inferences to be made regarding the balance of biogeochemical processes that prevail in subglacial environments as residence times and rock:water ratios increase, with 1994 and 1995 representing the longest-residence-time waters and 1997 the shortest. This provides a predictive framework for the biogeochemistry of large continental-scale ice sheets, where residence times are likely to be prolonged (days to months on Greenland and years beneath Antarctica), rock:water ratios are high and there is very limited contact with atmospheric gases (e.g., deformable sediments beneath ice streams). Second, the AD to CD transition at Finsterwalderbreen can be used to predict changes in bed biogeochemistry that might occur on contemporary ice sheets as the zone of surface melt expands with increased air temperatures [Steffen et al., 2004] and new areas of the bed begin to receive inputs from surface melt. We examine these two scenarios in turn in the following discussion by testing a series of hypotheses with regard to chemical dissolution under ice sheets.

1. Hypothesis 1 states: The significance of microbes in mediating subglacial chemical weathering is greater in longer-residence-time drainage systems, as might be found beneath ice sheets. We previously identified two processes that are attributable to microbial activity in chemical data sets from Finsterwalderbreen. First is the generation of microbial CO₂ that subsequently fuels the dissolution of carbonate and silicate minerals. Second is the anoxic oxidation of sulphide minerals using Fe(III), which is a microbially mediated process. In section 5.3, we calculated the potential solute fluxes from Finsterwalderbreen that are linked to these two types of microbial activity. If Hypothesis 1 is correct, we would expect to find a decrease in solute loads attributable to microbial activity, both in absolute terms and as a proportion of total solute loads, during the AD period as water residence times decrease. We might also expect to see progressively weakening relationships between HCO₃⁻ and (*Na⁺+*K) during the AD period, inferred to reflect decreased coupling between the microbial generation of CO₂ and silicate dissolution. It is clear from data during the AD period at Finsterwalderbreen that changes in residence times and rock:water ratios from 1994 to 1997 impacted on the biogeochemistry of meltwaters issuing from the SGU in these years. Beyond the decrease in ion concentrations observed due to rising icemelt input to the SGU chemical weathering environment, a shift in the balance of biogeochemical processes is identified. First, the role of microbes in mediating chemical dissolution decreased, both in absolute terms and as a proportional contribution to total solute loads during the AD period (Table 3, columns E and G and H), with the strongest microbial influence observed in 1994. Second, the role of microbial CO₂ in dissolving silicates decreased from 1995/1997 to 1999 (Figures 4 and 5) as residence times decreased. These observations support Hypothesis 1, indicating that microbes have the greatest impact on subglacial chemical weathering when meltwater residence times are prolonged, by the generation of protons via microbial respiration and microbially mediated sulphide oxidation. We would expect these processes to be further developed still on ice sheets, where flow paths may be >1000 km long and water residence times are, therefore, prolonged.

2. Hypothesis 2 states: Silicate dissolution becomes dominant over carbonate dissolution as meltwaters become saturated with respect to calcite in long-residence-time drainage systems. This is likely to be enhanced beneath ice sheets. Since silicate dissolution generates predominantly Na⁺ and K⁺ ions and carbonate dissolution generates Mg²⁺ and Ca²⁺ ions, the ratio of the monovalent to divalent cations largely indicates the relative dominance of silicate versus carbonate dissolution between the different study periods. Under Hypotheses 2, we would expect to see low ratios of (*Ca²⁺+*Mg²⁺):(*Na⁺+*K⁺) in longer-residence-time waters (e.g., AD period) compared with low-residence-time waters (CD period). We would also expect to see an increase in this ratio as meltwater residence times in the SGU decreased between 1995 and 1997. Our data set shows that silicate (including biotite) and dolomite dissolution were enhanced relative to carbonate and limestone dissolution, respectively, in the longer-residence-time SGU meltwaters (1994 and 1995), as evidenced by low ratios of (*Ca²⁺+*Mg²⁺):(*Na⁺+*K⁺) (Figure 5a). Correspondingly, an increase in the importance of carbonate dissolution was
observed as residence times and rock:water ratios decreased from 1995 to 1999, as evident from the inverse association between \( \text{Ca}^{2+} + \text{Mg}^{2+} + \text{Na}^{+} + \text{K}^{+} \) and ionic strength (Figure 5a). These observations support Hypothesis 2. One might expect the features of the longer-residence-time chemical data sets (1994 and 1995) to be even more developed beneath continental-scale ice sheets. Within a deformable sediment layer beneath an ice stream one might expect to find high rates of microbial activity (including CO₂ generation via respiration and SO₄²⁻ produced by microbiologically mediated sulphate oxidation), which in turn fuels silicate and carbonate dissolution via released protons, and drives the bed anoxic as the oxidation of sulphides and organic carbon consumes available oxygen.

Hypothesis 3 states: Anoxia increases as water residence times increase within sub-ice sheet environments that are remote from the atmosphere, resulting in the dominance of anaerobic microbial processes, such as sulphate reduction. Anoxic conditions may be inferred from the δ¹⁸O-SO₄²⁻ and δ³⁴S-SO₄²⁻ of runoff, as discussed in section 5.2. If the concept that anoxia increases with increased residence times holds, the δ¹⁸O-SO₄²⁻ and δ³⁴S-SO₄²⁻ of runoff from Finsterwalderbreen should show evidence for decreasing anoxia as water residence times decrease during the AD period and between the AD and CD periods. Isotope data presented in this paper provide strong evidence that meltwater residence times have a strong impact on the degree of anoxia attained in the subglacial environment. During the AD period, when high solute concentrations imply prolonged water:rock contact times, sulphate reduction was apparent where the residual sulphate was enriched in δ³⁴S and δ¹⁸O. Sulphate reduction no longer occurred during the CD period and sulphate was produced by a combination of anoxic and oxic sulphide oxidation. This is consistent with Hypothesis 3. The degree of anoxia would be expected to be high beneath ice sheets, where flow paths may be >1000 km long. Here, one might expect all sulphide oxidation to proceed anoxically using Fe(III) and sulphate reduction to occur, provided an organic carbon substrate exists. For ice sheets where there is an organic carbon supply (e.g., overridden soils, vegetation and/or marine/lake sediments) one might predict methanogenesis to occur, as has been suggested elsewhere [Weitmeyer and Buffett, 2006; Wadham et al., 2008].

Hypothesis 3 states: The input of surface meltwater to new areas of the ice sheet bed and the subsequent evolution of a distributed drainage system to channels will be accompanied by evolution from a silicate-dominated to carbonate-dominated chemical weathering regime. Under this hypothesis, one might expect to see an increase in the ratio of Ca²⁺ + Mg²⁺ + Na⁺ + K⁺ as meltwater residence times decrease, indicating a greater proportional contribution of carbonate versus silicate dissolution to solute loads. The transition between the AD and CD periods on Finsterwalderbreen is accompanied by a progressive decrease in the significance of silicate and dolomite (versus calcite) dissolution, as indicated by increasing ratios of Ca²⁺ + Mg²⁺ : *Na⁺ + *K⁺ and *Ca²⁺ : *Mg²⁺. We would expect similar trends to occur upon initiation of channelized flow at the bed of an ice sheet, driven by a reduction in meltwater residence times. This would produce a decrease in saturation indices with respect to calcite, and carbonate dissolution would be favored above the kinetically less favorable silicate dissolution. The same argument follows for the dissolution of dolomite, where present. This may have implications for resident microbes, which depend upon silicate dissolution to
release trace quantities of essential life elements such as N and P [Valsamis-Jones et al., 1998; Bratley et al., 2001].

6. Conclusions

[51] The above synthesis provides a unique insight to the relationship between subglacial hydrology and biogeochemical processes beneath large, long-residence-time glaciers. During its growth period (1994–1997), the SGU at Finsterwalderbreen constitutes an anoxic subglacial weathering environment where favorable conditions for sulphate reduction are apparent, silicate dissolution is enhanced and microbes are important in driving chemical dissolution (including that of silicates). The subsequent evolution of this environment to a channelized/ice-marginal system in which residence times and rock:water ratios are reduced witnesses a shift in runoff biogeochemistry. During this CD period, most solute derives from chemical weathering in ice-marginal moraines, and particularly the dissolution of Ca\(^{2+}\)-Mg\(^{2+}\)-SO\(_4\)^{2-} salts, with a reduced importance of silicate versus carbonate dissolution. Both the AD and CD period data sets provide first time compelling evidence for coupling between CO\(_2\) generated by microbial respiration and the dissolution of silicate mineral surfaces; suggesting that biofilms may be important in accelerating chemical dissolution in subglacial and ice-marginal sedimentary environments. In both cases, microbial activity (via sulphide oxidation and CO\(_2\) generation) accounts for up to ~50% of the solute flux from the SGU and ice-marginal moraine environment. This is consistent with work which has suggested that there are strong linkages between microbial activity and chemical dissolution processes beneath glaciers. The reorganization of drainage between 1994 and 1999 provides insight into the likely balance of biogeochemical processes under ice sheets. This has relevance given the current scientific interest in Antarctic subglacial lakes [Priscu et al., 2005; Siegert et al., 2007], and the potential shift in subglacial drainage that may occur beneath the Greenland Ice Sheet with an increased extent of surface melting and input to previously untapped basal environments [Steffen et al., 2004]. The interior regions of ice sheets are likely to be characterized by even longer residence times and a greater degree of isolation from the atmosphere than the SGU. In this type of chemical weathering environment one might expect; enhanced silicate dissolution, high anoxia and the prevalence of sulphate reduction and methanogenesis, diverse microbial communities which map onto specific Ph/Eh microniches at the bed and a chemical weathering regime that is essentially driven by microbial CO\(_2\) and microbially mediated sulphide oxidation using Fe(III). Under conditions of distributed system drainage evolution to channels triggered by enhanced melt inputs such as might be observed on the Greenland Ice Sheet in the coming decades, one might expect the opposite set of biogeochemical processes, namely, oxic/suboxic conditions (with an absence of sulphate reduction/methanogenesis), a reduced significance of silicate versus carbonate dissolution, reduced microbial involvement in chemical dissolution and a decrease in microbial diversity due to substrate limitations. These inferences provide a predictive framework for subice sheet biogeochemistry, which may become important in the coming decade as deep, subice sheet environments in Greenland and Antarctica are sampled for the first time.

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References


