Subglacial chemical erosion: seasonal variations in solute provenance, Haut Glacier d’Arolla, Valais, Switzerland

G. H. Brown,
Centre for Glaciology, Institute of Earth Studies, University of Wales, Aberystwyth, Dyfed SY23 3DB, Wales

M. Sharp,
Department of Earth and Atmospheric Sciences, University of Alberta, Edmonton, Alberta T6G 2E3, Canada

M. Tranter
Department of Geography, University of Bristol, Bristol BS8 1SS, England

ABSTRACT. This paper determines the provenance of solute in bulk meltwaters draining Haut Glacier d’Arolla, Valais, Switzerland, during the 1989 ablation season. Dissolved species are partitioned into components derived from sea salt, acid aerosol, dissolution of atmospheric CO₂, and lithogenic sources, namely carbonates, sulphides and aluminosilicates. A major conclusion is that trace geochemically reactive minerals in the bedrock contribute the bulk of the solute found in runoff. Seasonal changes in solute provenance and in the dominant chemical weathering process are observed. Whereas the chemical weathering of aluminosilicate minerals by carbonation reactions remains relatively constant during the ablation season, the chemical erosion of carbonates shows distinct seasonal variations, reflecting changes in the nature of the subglacial drainage system. Subglacial drainage structure and bedrock type are key controls on the extent of subglacial chemical weathering.

INTRODUCTION

The traditional view that chemical erosion is least intense in glacial environments, founded on the notion that low temperatures inhibit chemical weathering processes, has been overturned by numerous studies since 1970 (see Brown and others, 1994; Sharp and others, 1995). The traditional view failed to account for the importance of acid snowmelt, and underplayed the role of moisture availability, the freshly ground character of the rock flour, and geochemically reactive trace minerals. Regarding the extent to which water acts as a reagent, catalyst and carrier in chemical weathering reactions, its abundance should be crucial to the rate of chemical weathering reactions (Reynolds and Johnson, 1972; Lerman, 1979). Therefore, the seasonality of runoff in glaciated catchments is of especial significance, since the rate of dissolution is inversely related to time-dependent solute concentration, and hence the meltwater flushing rate (Lerman, 1979; Brown and others, in press).

A major focus of geochemical research in active glaciated regions has been to establish removal rates of major cations in meltwaters and to define chemical denudation rates. Despite temperature-dependence effects on the rate of dissolution (Drever and Zobrist, 1992; Velbel, 1993), such studies indicate rates of chemical erosion in glaciated catchments which are 1.2–2.6 times the continental average (Sharp and others, 1995). The efficacy of chemical weathering in glacial environments is attributable to high flushing rates, turbulent meltwaters, high suspended-sediment concentrations and the low buffering capacity of dilute meltwaters (Sharp and others, 1995). However, few studies have attempted to disaggregate the solute load of glacial runoff into crustal, atmospheric and snowpack sources (cf. Sharp and others, 1995), and to identify variations in solute sources and chemical weathering mechanisms on a seasonal time-scale (Tranter and others, 1993; Brown and others, 1994).

This paper analyzes the provenance of solute in the proglacial stream draining Haut Glacier d’Arolla during most of the 1989 ablation season, utilising meltwater-quality data derived from twice-daily sampling. Dissolved species are partitioned into components derived from sea salt, acid aerosol, dissolution of atmospheric CO₂, and lithogenic sources, namely carbonates, sulphides and aluminosilicates. Variations in solute provenance are then related to the seasonal evolution of the subglacial hydrological system.

FIELD SITE

Sampling was undertaken at Haut Glacier d’Arolla, the most southerly glacier in the Val d’Hérens, Valais, Switzerland. The glacier has an altitudinal range of ~2560–3500 m, and occupies approximately 6.3 km² of a
12 km² catchment. The maximum length of the glacier is approximately 4.2 km. A number of portals (2-5) contribute meltwater to the bulk runoff (Sharp and others, 1993). The bedrock geology is varied and consists of metamorphic and igneous rocks of the Arolla series of the Dent Blanche nappe, the highest tectonic unit of the Valais Alps (Dal Piaz and others, 1977; Mazurek, 1986). Geochemically reactive minerals, such as pyrite and calcite (Table 1), have been identified by microscopy and are present in trace amounts in many of the rocks throughout the catchment (Brown, 1991). In addition, XRD data from the main lithological units indicate that the main feldspars are albite, anorthite, microcline and sanidite, the main olivines are diopside, enstatite and spodumene. Actinolite, muscovite, corderite, hematite, hydrobasaluminate, quartz and talc are also present (personal communication from D. Webb, cited in Tranter and others, in press).

**METHODS AND TECHNIQUES**

Bulk meltwaters were sampled ~100 m from the glacier snout twice daily at 1000 and 1700 h local time (approximating to minimum and maximum diurnal discharge, respectively), from 1 June (Julian day 152) to 31 August 1989 (Julian day 243). The samples were immediately vacuum filtered through 0.45 μm cellulose nitrate membranes, and stored in pre-cleaned plastic bottles. Total alkalinity was determined colorimetrically in a field laboratory to an end-point of pH 4.5 using BDH mixed indicator solution and 1 mmol HCl. Precision was ±2%. The concentration of HCO₃⁻ was determined from the total alkalinity by correcting for acid needed to acidify a volume of deionised water equal to the volume of the aliquot and the titre. The major cations (Ca²⁺, Mg²⁺, Na⁺, K⁺) were determined by AAS, using an air-acetylene flame. Spectrochemical buffers, La(NO₃)₃ and CsCl, were added to overcome chemical interferences and ionisation, respectively. Accuracy was ±5%. Major anions (Cl⁻, NO₃⁻, SO₄²⁻) were determined by ion chromatography on a Dionex 4000i. Accuracy was ±3%.

Bulk meltwater dissolved-ion concentrations were separated into crustal, sea-salt and snowpack derived components, as detailed in Sharp and others (1995). All NO₃⁻ and Cl⁻ was assumed to be atmospherically derived, with sea-salt contributions of Ca²⁺, Mg²⁺, Na⁺, K⁺ and SO₄²⁻ derived from standard sea-water ratios with Cl⁻ (Holland, 1978). Atmospherically derived SO₄²⁻ associated with acid sulphate aerosols was calculated from the average Cl⁻:SO₄²⁻ ratio measured in the 1992/93 winter snowpack (0.4107). Since the early stages of aluminosilicate dissolution are non-stoichiometric (see Lerman, 1979), atmospheric HCO₃⁻ derived from the carbonation of aluminosilicates was estimated from the sum of *Na⁺ and *K⁺ (where * denotes non-sea-salt sodium and potassium) rather than from a ratio with silica (although carbonation of aluminosilicates probably also contributes to the fluxes of Ca²⁺ and Mg²⁺). Therefore, this represents the lower limit for CO₂ drawdown associated with the carbonation of aluminosilicates. Coupled sulphide oxidation and carbonate dissolution (SO₄/CD) generates all the crustally derived

---

**Table 1. Mineralogical composition (%) of fine material (<5 mm) from sampling locations A–I on the medial moraines of Haut Glacier d’Arolla (data from Brown, 1991)**

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
<th>G</th>
<th>H</th>
<th>I</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbonates:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcite</td>
<td>5.67</td>
<td>11.45</td>
<td>0.00</td>
<td>0.00</td>
<td>0.42</td>
<td>3.55</td>
<td>1.35</td>
<td>0.00</td>
<td>0.41</td>
</tr>
<tr>
<td>Siliicates:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Quartz</td>
<td>27.53</td>
<td>39.02</td>
<td>58.48</td>
<td>66.40</td>
<td>64.35</td>
<td>42.55</td>
<td>55.61</td>
<td>66.75</td>
<td>60.94</td>
</tr>
<tr>
<td>Orthoclase</td>
<td>0.00</td>
<td>0.23</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.25</td>
<td>0.00</td>
</tr>
<tr>
<td>Microcline</td>
<td>0.61</td>
<td>0.00</td>
<td>0.00</td>
<td>0.60</td>
<td>0.63</td>
<td>0.00</td>
<td>0.90</td>
<td>2.25</td>
<td>0.00</td>
</tr>
<tr>
<td>Plagioclase</td>
<td>0.40</td>
<td>0.00</td>
<td>0.39</td>
<td>0.20</td>
<td>0.00</td>
<td>1.18</td>
<td>0.22</td>
<td>2.00</td>
<td>2.25</td>
</tr>
<tr>
<td>Perthite</td>
<td>0.61</td>
<td>0.00</td>
<td>0.78</td>
<td>0.40</td>
<td>0.63</td>
<td>0.95</td>
<td>1.35</td>
<td>0.00</td>
<td>0.82</td>
</tr>
<tr>
<td>Tremolite-actinolite</td>
<td>8.70</td>
<td>2.57</td>
<td>0.78</td>
<td>3.02</td>
<td>2.53</td>
<td>0.71</td>
<td>0.00</td>
<td>0.50</td>
<td>1.23</td>
</tr>
<tr>
<td>Hornblende</td>
<td>0.20</td>
<td>0.00</td>
<td>0.00</td>
<td>0.80</td>
<td>0.42</td>
<td>0.24</td>
<td>0.00</td>
<td>0.00</td>
<td>1.43</td>
</tr>
<tr>
<td>Amphibole (misc.)</td>
<td>4.86</td>
<td>2.10</td>
<td>1.56</td>
<td>3.84</td>
<td>5.49</td>
<td>3.31</td>
<td>1.57</td>
<td>0.50</td>
<td>2.66</td>
</tr>
<tr>
<td>Sphene</td>
<td>0.40</td>
<td>3.04</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Zoisite</td>
<td>4.25</td>
<td>7.24</td>
<td>2.92</td>
<td>0.20</td>
<td>0.21</td>
<td>0.71</td>
<td>1.79</td>
<td>1.50</td>
<td>0.81</td>
</tr>
<tr>
<td>Epidote</td>
<td>1.21</td>
<td>0.93</td>
<td>0.39</td>
<td>2.82</td>
<td>0.42</td>
<td>0.00</td>
<td>0.30</td>
<td>4.50</td>
<td>5.50</td>
</tr>
<tr>
<td>Biotite</td>
<td>10.93</td>
<td>3.97</td>
<td>1.75</td>
<td>0.20</td>
<td>0.42</td>
<td>5.44</td>
<td>0.45</td>
<td>1.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Mica (misc.)</td>
<td>10.53</td>
<td>4.67</td>
<td>3.51</td>
<td>2.41</td>
<td>1.68</td>
<td>8.27</td>
<td>9.87</td>
<td>7.00</td>
<td>2.25</td>
</tr>
<tr>
<td>Chlorite</td>
<td>7.29</td>
<td>5.61</td>
<td>1.17</td>
<td>3.62</td>
<td>6.96</td>
<td>9.22</td>
<td>4.48</td>
<td>3.25</td>
<td>6.75</td>
</tr>
<tr>
<td>Opaques:</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mainly pyrite</td>
<td>3.24</td>
<td>2.10</td>
<td>1.56</td>
<td>1.21</td>
<td>1.05</td>
<td>4.96</td>
<td>1.35</td>
<td>2.75</td>
<td>0.82</td>
</tr>
<tr>
<td>Others</td>
<td>0.00</td>
<td>0.47</td>
<td>0.19</td>
<td>0.20</td>
<td>2.95</td>
<td>0.24</td>
<td>0.00</td>
<td>2.75</td>
<td>0.00</td>
</tr>
<tr>
<td>Total</td>
<td>100.40</td>
<td>99.99</td>
<td>99.99</td>
<td>99.99</td>
<td>99.56</td>
<td>100.01</td>
<td>100.02</td>
<td>100.00</td>
<td>100.00</td>
</tr>
</tbody>
</table>
SO$_4^{2-}$, yielding one equivalent of HCO$_3^-$ and two equivalents of Ca$^{2+}$ + Mg$^{2+}$. Snowpack acidity from acid sulphate and nitrate aerosols was assumed to weather carbonate minerals to produce crustally derived HCO$_3^-$. The residual HCO$_3^-$ was assumed to arise equally from the dissolution and dissociation of atmospheric CO$_2$ and the carbonation of carbonate minerals, which produced all the crustally derived non-SO/CD Ca$^{2+}$ and Mg$^{2+}$.

RESULTS AND DISCUSSION

Figure 1 shows the discharge and suspended-sediment concentration records for much of the 1989 ablation season (Brown and others, 1994). Discharge and discharge amplitude increased as the season progressed, accompanied by an increase in suspended-sediment concentration, reflecting the seasonal evolution of the drainage system (Brown and others, 1994).

The subglacial hydrological system of Alpine glaciers appears to be adequately described by two principal flow components (Collins, 1978; Oerter and others, 1980; Tranter and Raiswell, 1991). Quick-flow waters are derived largely from icemelt, and pass rapidly through a channelled hydrological system of ice-walled channels and major arterial conduits (Sharp, 1991; Brown and others, 1994). Conversely, delayed-flow waters are derived largely from snowmelt, and pass more slowly through a distributed hydrological system (e.g., linked, water-filled cavities [Walder, 1986]). These two components mix to form bulk meltwaters draining the glacier, with the channelised system expanding headwards at the expense of the distributed system as the melt season progresses (Brown and others, 1994).

Sources of aqueous protons

Chemical erosion by Alpine glacial meltwaters is predominantly by the broad class of reactions known as acid hydrolysis (Tranter and others, 1993). Raiswell (1984) has suggested that the anionic content of glacial meltwaters demonstrates the acid source used in the chemical weathering of rock minerals. HCO$_3^-$ is the dominant anion where the dissolution and dissociation of atmospheric CO$_2$ provides protons to fuel chemical erosion (Equations (1) and (2)).

\[
\text{CaCO}_3(s) + CO_2(aq) + H_2O(aq) \rightarrow \text{Ca}^{2+}(aq) + 2HCO_3^-(aq) \quad (1)
\]

\[
\text{CaAl}_2\text{Si}_2\text{O}_8(s) + 2CO_2(aq) + 2H_2O(aq) \rightarrow \text{Ca}^{2+}(aq) + 2HCO_3^-(aq) + H_2\text{Al}_2\text{Si}_2\text{O}_8(s) \quad \text{partially weathered feldspar} (2)
\]

Conversely, SO$_4^{2-}$ is the dominant anion where protons are derived from the oxidation of sulphide minerals (Equation (3)).

\[
4\text{FeS}_2(s) + 15\text{O}_2(aq) + 14\text{H}_2\text{O}(aq) = 16\text{H}^+(aq) + 4\text{Fe(OH)}_3(s) + 8\text{SO}_4^{2-}(aq) \quad \text{ferric oxyhydroxides} (3)
\]

Typically, chemical reactions are coupled, such that solute acquisition in glacial meltwaters involves species in solid, liquid and aqueous phases (Tranter and others, 1993), e.g.,

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

Therefore, the proportion of HCO$_3^-$ and SO$_4^{2-}$ in the bulk meltwaters will reflect the relative dominance of the two major sources of protons driving subglacial chemical erosion (Fig. 2). Expressed as the ratio of HCO$_3^-$ to (HCO$_3^-$ + SO$_4^{2-}$), a ratio of 1.0 would signify carbonate reactions involving pure dissolution and acid hydrolysis, consuming protons derived from atmospheric CO$_2$ (Equations (1) and (2)). Conversely, a ratio of 0.5 suggests coupled reactions involving the weathering of carbonates by protons derived from sulphide oxidation (Equation (4)).

It is clear from Figure 2 that when flow from the distributed system, which underlies the whole glacier in early June (Brown and others, 1994), dominates bulk discharge early in the ablation season, the ratio of HCO$_3^-$/HCO$_3^- +$ SO$_4^{2-}$ (hereafter the C-ratio) (~0.65) suggests that coupled reactions, involving carbonate dissolution and protons derived primarily (though not exclusively) from the oxidation of sulphide minerals, largely control bulk meltwater composition. It
Brown and others: Subglacial chemical erosion

is unlikely that a system of linked cavities (Walder, 1986) is in intimate contact with a large source of atmospheric CO₂ to enhance the C-ratio to 0.5 (Tranter and others, 1993). Other possible sources of CO₂(aq) in the distributed system may include HCO₃⁻ in meltwaters feeding the distributed system, the release of bubbles trapped in basal ice facies (Tison and others, 1993; Tranter and others, in press), microbial oxidation of organic carbon (Tranter and others, in press) and the oxidation of elemental free carbon. Carbon occurs both as a combined form (mainly as carbonates of Ca²⁺ and Mg²⁺) and as a free element (e.g. diamond and graphite) in addition to that associated with CO₂ (Greenwood and Earnshaw, 1984). Graphite is widely distributed, associated with quartz and metamorphosed sedimentary silicate rocks such as mica-schists and gneisses all of which are present in the Haut Glacier d’Arolla catchment (Brown, 1991). Indeed, this recycling of carbon from sediments to fresh waters via rock weathering forms an integral part of the global carbon cycle (e.g. Holland, 1978; Stumm and Morgan, 1981). The importance of these coupled reactions in controlling meltwater composition in the delayed-flow component is also evident to a lesser extent in the C-ratio (<0.75) during the major recession flow event at the end of July (Julian day 212) (when snow fell on the glacier surface, reducing surficial meltwater inputs to the hydroglacial system, resulting in delayed flow waters forming an increasing proportion of the bulk runoff), and in the diurnal variability from mid-July onwards when the C-ratio is lower at daily minimum discharge. Conversely, when the quick-flow component dominates maximum diurnal bulk meltwater discharge from mid-July onwards (Brown and others, 1994), the C-ratio (0.75–0.9) suggests protons are derived primarily from the dissolution and dissociation of atmospheric CO₂.

Protons to fuel chemical erosion may also be derived from the dissolution of acid sulphate and nitrate aerosols in the seasonal snowpack (Tranter and others, 1993). Significant contributions from this source of acidity are limited to the early melt season (Fig. 3), as a result of fractionation processes within the snowpack (Johannessen and Henriksen, 1978) and the up-glacier retreat of the seasonal snow cover as the ablation season proceeds (Tranter and others, 1993).

**Lithogenic solute provenance**

Contributions of solute from various rock types are often calculated by assuming that their contribution is proportional to their mass (Garrels and Mackenzie, 1971). However, minerals weather at different rates, and examples of such weathering sequences are common in the geochemical literature (e.g. Goldich, 1938; Stumm and Morgan, 1981). Therefore, the relative abundance and reactivity of available minerals within a catchment will affect the degree and nature of chemical erosion. Using HCO₃⁻ as an indicator of the relative proportions of solute contributed by lithogenic (coupled SO₂/CD erosion, dissolution of carbonates by acidity derived from the snowpack (SP/CD) and carbonation of carbonates (C/CD)) and atmospheric (carbonation of aluminosilicates and carbonates) sources (Fig. 4), the calculations detailed above suggest that the weathering of carbonates supplies the vastly greater part of the HCO₃⁻ in solution. This is in direct contrast to the actual percentage of carbonate within the bedrock, which from samples of fines (<3 mm diameter) from supraglacial moraines is <12%, and generally <5% (Table 1; Brown, 1991). While the total contribution from carbonate minerals is in excess of 85%, the proportion of solute acquired from carbonation of carbonates and SO₂/CD reactions, and the weathering mechanisms producing the solute, change as the ablation season progresses. This is clearly illustrated in Figure 4, and will be discussed in more detail in the following paragraphs, where we discuss solute provenance from the major lithogenic sources.

(i) Carbonation of carbonates

The stoichiometry of Equation (1) suggests that the proportion of the HCO₃⁻ load derived from lithogenic and atmospheric sources is equal in equivalence units. If it is assumed that acidity derived from the seasonal snowpack (Fig. 3) is predominantly neutralised by the weathering of carbonates during the early melt season (based on the idea that most solute appears to come from the weathering of lithogenic carbonates; Sharp and others, 1995), this augments the levels of early-season HCO₃⁻ derived from the carbonation of carbonates, resulting in a relatively constant contribution of litho-
suspended sediment concentrations, spherical stems are dominated by coupled sulphide oxidation and derived from the dissolution and dissociation of atmospherically derived HCO$_3^-$ (b) is derived from the dissolution and dissociation of atmospheric CO$_2$ associated with the hydrolytic weathering of aluminosilicate and carbonate minerals.

**Genetic HCO$_3^-$ throughout the entire ablation season (Fig. 5a).** This is in direct contrast to the proportion of HCO$_3^-$ derived from the dissolution and dissociation of atmospheric CO$_2$ (Fig. 5b), which shows a steady increase as the ablation season proceeds, reflecting the headwards expansion of the channelised system, the increasing proportion of quick-flow waters in runoff, and increasing suspended-sediment concentrations.

**(ii) Coupled sulphide oxidation and carbonate dissolution**

Recent models of solute acquisition beneath Alpine glaciers suggest that chemical erosion in the distributed system is dominated by coupled sulphide oxidation and carbonate dissolution (Tranter and others, 1993, in press; Brown and others, 1994; Equation (4)). This is confirmed in the provenance calculations, which illustrate a peak in HCO$_3^-$ (SO/CD), SO$_4^{2-}$ (crystal) and Ca$^{2+}$ + Mg$^{2+}$ (SO/CD) during the early part of the ablation season (Julian days 152-165), when delayed flow waters dominate runoff (Fig. 6). The role of SO$_4/CD$ reactions in the distributed system is also evident to a lesser extent during the recession flow event at the end of July (Julian days 212-220).

**(iii) Carbonation of aluminosilicates**

Relative to the carbonation of carbonate minerals, the weathering of aluminosilicates by carbonation reactions remains relatively constant during the ablation season (Fig. 4b), exhibiting only a slight increase as the melt season progresses. This is illustrated (Fig. 7a) by variations in atmospherically derived HCO$_3^-$ associated with the carbonation of aluminosilicates. Na$^+$ shows a monotonic increase as the ablation season progresses, associated with increased weathering of plagioclase- and sodic-feldspars (Fig. 7b). Following an early-season peak, K$^+$ also increases, especially during August, associated with increased weathering of biotite, muscovite and potash-feldspar (Fig. 7c). This suggests that part of the increasing proportion of solute derived from carbonation reactions between dilute quick-flow waters and suspended sediment is derived from aluminosilicate minerals in suspension as the ablation season progresses (Fig. 4; Brown and others, 1994).

**CONCLUSIONS**

To date, few studies have attempted to disaggregate the solute load of glacial runoff into crustal, atmospheric and snowpack sources, and to identify variations in solute sources and weathering mechanisms on a seasonal timescale. Therefore, snowpack-derived solute is usually...
incorporated in estimates of chemical denudation rates in
glacial catchments (Sharp and others, 1995). When
dissolved species are partitioned into components derived
from sea salt, acid aerosol, dissolution of atmospheric CO$_2$
and lithogenic sources, seasonal changes in solute
provenance and in the dominant chemical weathering
process are observed.

Trace geochemically reactive minerals in the bedrock
(e.g., carbonate and sulphide minerals) appear to
contribute the major proportion of solute found in bulk
runoff. The major anions are bicarbonate and sulphate,
suggesting that two primary sources of protons are utilised
to fuel the chemical erosion of glacial sediments: (i)
sulphide oxidation, and (ii) the dissolution and dissociation
of atmospheric CO$_2$. Solute contributions from the
seasonal snowpack are limited to the early melt season,
accompanied by a transient weathering of crustal
material associated with snowpack acidity.

Whereas the chemical weathering of aluminosilicate
minerals by carbonation reactions supplies a relatively
constant proportion of bulk meltwater solutes during the
ablation season, the chemical erosion of carbonates shows
distinct seasonal variations in the proportion supplied,
reflecting changes in the nature of the subglacial drainage
system. There are two subglacial environments in which
carbonates are chemically eroded, the distributed and
channelised drainage systems. Chemical weathering of carbonates in the distributed system is driven by protons derived from the oxidation of sulphide minerals. This peaks early in the ablation season and is relatively constant thereafter, reflecting the reduction in areal extent of the distributed system, and hence the proportion of delayed flow waters in the bulk discharge. In contrast, carbonation of carbonates is the dominant weathering process in quick-flow waters where there is usually free access to atmospheric CO₂. The magnitude of carbonation increases as the channelised drainage system grows throughout the ablation season, accompanied by an increase in the proportion of quick-flow waters and suspended-sediment concentrations in bulk runoff. In addition, carbonation of the aluminosilicate fraction of the suspended-sediment load also increases, reflected in the proportion of atmospheric HCO₃ derived from the carbonation of aluminosilicates, "Na" and "K". Therefore, the seasonal evolution of the subglacial drainage system plays an important role in determining both the source of solute and the weathering mechanism driving chemical erosion.

The chemical weathering of carbonates is enhanced during the early melt season by acidic snowmelt. The decrease in this mechanism of chemical weathering through the ablation season is offset by the increase in chemical weathering in the channelised drainage system driven by carbonation reactions. We conclude that subglacial drainage structure and the relative abundance and reactivity of available minerals within the catchment are key controls on the extent of subglacial chemical weathering.

ACKNOWLEDGEMENTS

This work was supported by a U.K. Natural Environment Research Council Studentship (GT4/88/AAPS/56) and Fellowship (GT5/F91/AAPS/3) to G.H.B. and by NERC Grants GR3/7004a and GR3/8114. The Royal Society and University of Wales Learned Societies Fund provided financial support to attend the conference in Reykjavik. Dr A. Barker is gratefully acknowledged for his contributions to the derivation of the lithological data in the Department of Geology, University of Southampton. C. Hill provided invaluable assistance in the field.

REFERENCES