**Seasonal release of anoxic geothermal meltwater from the Katla volcanic system at Sólheimajökull, Iceland**

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**Abstract**

Understanding patterns of geothermal and volcanic activity at many of Iceland’s most active volcanic systems is hampered by thick overlying ice, which prevents direct observation and complicates interpretation of geophysical signals. Katla is a prime example, being a large and restless volcanic system covered by the 740 m thick Mýrdalsjökull ice cap, whose eruptions have triggered some of the most powerful known meltwater floods in historical times. To shed new light on geothermal and subglacial hydrological processes at Katla, we have determined the sulphate isotopic composition of a series of glacial meltwater samples discharged from Sólheimajökull, a valley glacier of Mýrdalsjökull, between 2009 and 2012. Dual isotopic analysis of δ34S and δ18O in dissolved sulphate allow identification of source mixing processes and chemical evolution during subglacial meltwater transport. Strikingly, meltwater δ18OSO4 signatures indicate redox conditions at the glacier bed, which are inverse to those normally encountered at Arctic and Alpine glaciers. Discharge of reduced, anoxic meltwater in summer, rather than winter, points towards seasonal release of geothermally derived volatile gases. We attribute this to headward expansion of the channelized subglacial drainage system during the summer melt season, accessing key areas of geothermal activity within the Katla caldera. Volatile release may be further enhanced by unloading of overburden pressure due to snowpack melting in the summer season. In winter, restriction of subglacial channels to lower elevations effectively seals geothermal fluids and dissolved gases beneath the ice cap, with only sporadic release permitted by periodic increases in subglacial water pressure. When the subglacial drainage configuration permits access to key geothermal areas, sulphate isotopic signatures thereby form sensitive indicators of geothermal activity occurring deep beneath the Mýrdalsjökull ice cap.

Key words: Iceland, Mýrdalsjökull, Sólheimajökull, Geothermal water, glacial hydrochemistry, sulphate isotopes

**Introduction**

Glacial meltwater hydrochemistry is typically used as a fingerprinting technique for deciphering physical, chemical and biological processes occurring deep beneath ice masses (e.g. Galeczka et al., 2014; Hodson et al., 2005; Tranter et al., 1993, 1997). Such information is particularly significant in the context of glaciated volcanoes, where ice cover complicates interpretation of geophysical signals, making eruptive events difficult to forecast or identify (Guðmundsson et al. 2008). Melting of ice in glaciated volcanic systems may stimulate eruptive and geothermal activity (e.g. Pagli and Sigmundsson 2008, Tuffen 2010), which in turn can destabilise ice bodies (Bell 2008), creating positive feedback that may have major environmental implications (Huybers and Langmuir 2009). Several studies have addressed the discharge and geochemical composition of episodic flood waters generated through subglacial volcanic eruptions (e.g. Elefsen et al., 2002; Galeczka et al., 2014; Gíslason et al., 2002;), although few have used sensitive geochemical indicators to monitor the continual evolution of ice-capped geothermal systems (Kristmannsdóttir et al., 2002). Here, we evaluate the incidence of subglacial geothermal activity at Sólheimajökull, Iceland, an outlet glacier from the Mýrdalsjökull ice cap, which conveys large quantities of volatiles dissolved in meltwater outlet streams. The Sólheimajökull subglacial drainage system conveys volatile species derived from the ice-covered Katla caldera on the Mýrdalsjökull-Katla system, with the frequent smell of hydrogen sulphide emanating from the meltwater streams providing a strong indication of persistent subglacial geothermal activity (Björnsson et al., 2000). Existing research has documented the background chemistry in bulk meltwater outputs from the catchment and tentatively attributed these to variations in the strength of the Katla geothermal field (Lawler et al., 1996; Sigvaldason, 1963). However, the drivers of geothermal activity, and whether any associated chemical signal could be used reliably to understand processes in the subglacial geothermal system at Katla, remain poorly understood. Stable isotopes allow fingerprinting of subglacial chemical sources, their mixing and chemical evolution during meltwater transport. The rapid oxidation of reduced sulphurous gases to sulphate, which is subsequently relatively stable in solution, make the dual isotopic analysis of sulphate an attractive contender as an indicator of subglacial geothermal activity. This paper aims to i) identify the sources and biogeochemical cycling of sulphur beneath the Mýrdalsjökull ice cap ii) explain seasonal variations in redox status at the glacier bed and iii) evaluate the use of sulphur isotopes as indicators of subglacial geothermal activity. We begin by briefly reviewing the nature of sulphate acquisition and isotopic signatures within glaciated catchments to provide context to the study.

**Sulphur acquisition and isotopic signatures within glaciated catchments**

Sources of sulphur within glaciated catchments are predominantly restricted to atmospheric deposition and release from catchment geology. Atmospheric sulphur deposition is composed of marine aerosol and industrial pollution, with minor components from volcanic outgassing and continental biogenic emissions. Oxidation in the atmosphere to sulphate and subsequent deposition as rain/snowfall creates a supraglacial sulphate input to the glacier which varies in concentration and isotopic composition depending upon the dominance of each of the above sources. Sulphur isotopic values of each of the sources range between marine aerosol, which varies little from +21 ‰ (Rees et al., 1978) and continental biogenic emissions that form a light isotopic end member ranging from approximately 0 to -30 ‰ (Nielson, 1974). Volcanic aerosol emissions approximate 0 ‰ (Nielsen et al., 1991) and the release of sulphur gases from combustion of fossil fuels depends upon the sulphur isotopic composition of the source materials which has the capability of stretching over the whole range of values given above (see inventory in Newman et al., 1991; Nielson, 1974). Global precipitation values thereby exhibit isotopic signatures that reflect this mixed composition. For δ34SSO4 found within European precipitation (including snowfall) the typical range is between -3 to +9 ‰ reflecting a high proportion of industrial pollution (Mayer, 1998); for precipitation over mid-latitude sites, the range resides between -2.5 to + 13.5 ‰ (Jenkins and Bao, 2006), whilst δ34SSO4 extracted from shallow ice cores in central Antarctica range between +9.3 to +18.1 ‰ (Patris et al., 2000) and +14.5 to +15.0 ‰ (Jonsell et al., 2005) reflecting a dominance of marine, and volcanic input in the absence of industrial pollution.

Catchment geology contributes to the sulphur load in glacial meltwaters through release of sulphur from geothermal activity and weathering of bedrock. Release of sulphur species from geothermal areas are documented to have a downstream impact on the load and isotopic composition of sulphate leaving glaciated catchments, dependent upon the strength of the geothermal field and the dissolution of volatile sulphur species into ground waters (e.g. Gíslason et al., 2002; Robinson et al., 2009). Bedrocks contain sulphur usually in the form of pyrite, gypsum, or secondary mineral precipitates. Studies of bedrock sulphur isotope composition from volcanic rocks within Iceland range from -2 to +0.4 ‰ for basalts, and up to + 4.2 ‰ for acidic volcanic rocks (Torssander, 1989). δ34S values of pyrite from rhyolite clasts at Skeiðarársandur range between -1.2 to +2.6 ‰ (Robinson et al., 2009) and δ34S values of lava incrustations (sublimates deposited in solidifying lavas) reflect closely the isotopic signature of host basalts and precursor volcanic gas species (Torssander, 1988). Acquisition of sulphur by bedrock weathering beneath a glacier proceeds via oxidation and dissolution reactions. Both of these reactions are largely dependent upon rock-water contact time and availability of dissolved gases, governed according to the subglacial or proglacial environment of solute acquisition and associated hydrological characteristics (e.g. Bottrell and Tranter, 2002; Wadham et al., 2007; Wynn et al., 2006).

Different types of subglacial drainage system are widely held to exist beneath glaciers and ice sheets, their characteristics dependent on meltwater discharge, temperature distribution at the ice-bed interface and the topography, permeability and rigidity of the glacier bed. Within temperate glacial systems such as those considered as a part of this paper, research recognises two main types of subglacial drainage; firstly discrete systems in which water is confined to a few channels or conduits, and secondly, distributed systems in which water is stored and transported over all or large parts of the glacier bed (e.g. Kamb, 1987; Lliboutry, 1968, 1979; Walder, 1986;). Discrete systems are efficient transporters of meltwater exhibiting rapid flow through a well-connected conduit system; in contrast, distributed systems are inefficient, with meltwater flowing slowly through poorly-connected networks. Distributed systems often take the form of linked cavity drainage networks, in which cavities permit temporary water storage and narrow connections or orifices link the cavities together. As subglacial discharge increases, melting of ice from the roof of the cavities offsets some of the ice pressure that would cause cavity closure and large low-pressure cavities effectively capture drainage from small high-pressure cavities, enlarging as they do so. Eventually the linked cavity drainage network becomes unstable at high meltwater discharges and the system switches to a discrete, efficient system (Fountain and Walder, 1998).

Both discrete and distributed drainage systems (termed ‘quick flow’ and ‘delayed flow’ respectively according to the residence time of meltwater at the bed within each) are not mutually exclusive, but compete on an annual basis such that the quick flow channelized system demonstrates headward expansion during the melt season and the delayed flow linked cavity system becomes spatially restricted to areas where a deep snowpack minimises the flux of water to the glacier bed. During periods of minimal melting, channelized systems close due to ice deformation, although cavity networks remain open and demonstrate spatial expansion due to a reliance on ice dynamics for cavity formation (Fountain and Walder, 1998). On temperate Icelandic glacial systems, where year-round melting usually persists in the lower reaches of the glacier, discrete channelized systems remain a permanent feature. Changes to the extent of the channelized network are determined through seasonal water flux to the glacier bed and mirror changes in the extent of the distributed system (Björnsson, 1988; Flowers, et al., 2003). This switch in the hydrology is mimicked through a ‘switch’ in the chemistry, associated with the degree of rock-water contact and availability of atmospheric gases afforded by each system. Distributed systems with extensive rock-water contact drive enhanced solute acquisition. The limited supply of atmospheric gases to the glacier bed determines the suite of available reaction mechanisms with many reactions in distributed systems occurring under conditions of low redox status (Wynn et al., 2006). Quick flow systems transit the subglacial environment with such rapidity that the degree of rock-water contact is a limiting factor on solute acquisition from bedrock weathering (Tranter et al., 1993). Acquisition of solutes from bedrock weathering continues through the proglacial zone through reaction with suspended sediments (e.g. Brown et al., 1994; Fairchild et al., 1999), weathering of near surface moraine and from contributing groundwaters (Robinson et al., 2009; Wadham et al., 2007), and re-dissolution of gypsum and other secondary mineral precipitates (Cooper et al., 2002).

Where ambient environmental conditions are fully oxidizing and the supply of atmospheric gases do not represent a limiting factor, there is very limited fractionation of sulphur isotopes during oxidative weathering (Taylor et al., 1984). Dissolution of secondary mineral precipitates and volatile geothermal gases are also documented to incur minimal isotopic fractionation (Ohmoto and Rye, 1979). The dissolved sulphate load in glacial meltwaters thereby retains an isotopic signature that directly reflects the isotopic composition of the geologic parent materials. However, under strongly reducing conditions, dissolved sulphate can be microbially reduced to produce sulphide, leaving a residual pool of dissolved sulphate enriched in 34S and 18O (eg. Spence et al., 2001; Strebel et al., 1990). Under anoxic conditions beneath glaciers which are closed to the external environment, a positive relationship between δ34SSO4 and δ18OSO4  in the residual dissolved sulphate pool is indicative of microbial sulphate reduction (Wadham et al., 2004). At redox conditions of intermediate status, representing areas of the subglacial system that are poorly connected to the atmosphere, oxidizing agents other than atmospheric oxygen are used to drive oxidation weathering, mediated through microbiological activity (e.g. Bottrell and Tranter, 2002). During this process, the sulphur isotopic composition of the source bedrock is preserved in the product sulphate, although oxygen source mixing and isotopic exchange between water and sulphur intermediate species leave a distinctive δ18OSO4 signature serving as an important indicator of redox status.

Under atmospheric conditions, the oxidation of sulphide to sulphate is dependent upon obtaining an oxygen source from both atmospheric O2 and H2O (Equation 1).

4FeS2(s) + 15O2(aq) + 14H2O(l) → 16H+(aq) + 4Fe(OH)3(s) + 8SO42-(aq) (1)

However, if atmospheric oxygen is not available, the reaction can still proceed, albeit through microbial mediation under reducing conditions, using water as the sole oxygen provider (Equation 2).

FeS2 + 14Fe3+ + 8H2O → 2SO42- + 15Fe2+ + 16H+ (2)

Due to the large isotopic difference between oxygen sources used during the process of oxidation (H2O = -9.66 ‰ (average value of Sólheimajökull melt water δ18OH2O); atmospheric O2 = +23.88 ‰ (Barkan and Lus, 2005)), the varying proportion in which they are incorporated into the sulphate molecule can inform about the environmental redox status at the point of formation.

The instability of intermediate reaction products during the formation of sulphate means up to three of the four constituent oxygen atoms undergo rapid equilibration with the surrounding water (Holt and Kumar, 1991). At low temperature and pH, the final oxygen incorporated into the sulphate molecule is stable and thereby indicative of source (Lloyd, 1968). Sulphate molecules, which contain at least 25 % oxygen obtained from an atmospheric source, are thereby indicative of a subglacial drainage system which is connected to the atmosphere. For those sulphate molecules with >75 % oxygen obtained from H2O, it can be assumed these were formed under conditions of low redox status through microbially-mediated reaction mechanisms utilising Fe3+ as an oxidising agent. A threshold isotopic value can be calculated to express this classification of sulphate molecules according to the ambient redox conditions in their environment of formation (Equation 3) (after Bottrell and Tranter, 2002).

δ18Othreshold = [(23.88 – 8.7) x 0.25] + (0.75 x δ18OH2O) (3)

Where +23.88 ‰ represents the isotopic value of atmospheric oxygen (Barkan and Lus, 2005), which incurs an experimentally determined fractionation of -8.7 ‰ during incorporation into the sulphate molecule (Lloyd, 1968). Where sulphur intermediate species are a product of this oxidation reaction (eg. elemental sulphur and thiosulphate), bacterial disproportionation reactions can lead to the formation of hydrogen sulphide and / or sulphate (Bottcher et al., 2005). The extent of fractionation associated with disproportionation depends on the presence of either iron or manganese as oxidizing agents (Bottcher et al., 2001a, b) but may enrich product sulphate in 34S by up to + 35 ‰ and in 18O by up to +21 ‰ compared to precursor sulphur compounds and water respectively. Repeated cycles of disproportionation and re-oxidation will deplete product sulphides in 34S beyond the envelope of fractionation expected through processes of sulphate reduction alone (Canfield and Thamdrup, 1994).

Glaciers have been observed to fluctuate in subglacial redox status largely according to seasonality. In High Arctic and temperate glacial systems, spring season and early summer subglacial meltwaters suggest low redox status at the bed of the glacier. Full oxygenation returns during times of high summer discharge (Bottrell and Tranter, 2002; Wynn et al., 2006; Wynn et al., 2007). Evidence for occasional recurrence of low redox status during the summer season has been attributed to persistence of the distributed drainage system under portions of the glacier and its variable mixing with oxygen saturated quick flow melt waters on a range of timescales (Bottrell and Tranter, 2002; Irvine-Fynn and Hodson, 2010).

Due to the limited isotopic fractionation of geological sulphur species under oxidizing conditions and the frequent overlap in isotopic signatures between sulphur released from geothermal sources and those released from mineral sources, the use of sulphate isotopes as indicators of subglacial geothermal processes is still poorly documented and understood. Attribution of any chemical fingerprint to understanding subglacial geothermal processes must rely on being able to deconvolve mixing of sulphur sources, as well as indicate mechanisms of biogeochemical cycling and chemical evolution during transport. Here, we use stable environmental isotopes of δ34SSO4 and δ18OSO4 in conjunction with conventional meltwater chemical assays to identify the sources and biogeochemical cycling of sulphur beneath the Mýrdalsjökull ice cap. We implicate the role of subglacial geothermal processes in contributing to the sulphur load and trace the redox evolution of the subglacial hydrological system, which appears to be driven by seasonality in the release of geothermally-produced fluids.

**Methods**

***Site description***

The Mýrdalsjökull ice cap covers the Katla volcanic system, which constitutes the central Katla volcano and fissure swarm (Duller et al., 2012; Sturkell et al., 2010). The Katla caldera is 600-750m deep, and covers an area of 100 km2. Its rim is breached in three places, which provides outflows of ice to feed the three major outlet glaciers of Sólheimajökull, Kötlujökull and Entujökull (Björnsson et al., 2000, Sturkell et al., 2010). Geothermal activity within the caldera is manifest as several 10-50m deep ice cauldrons within and at the caldera rim, with a total geothermal output of a few hundred megawatts (Guðmundsson et al., 2007). The drainage divide of Sólheimajökull continues into the south western rim of the Katla subglacial caldera, making it possible for volcanically-generated and geothermal ice melt to drain through Sólheimajökull as jökulhlaups (Björnsson et al., 2000, Larsen, 2010; Larsen et al., 2001). Most historic jökulhlaups generated by Katla eruptions have drained through neighbouring Kötlujökull onto Mýrdalssandur to the east; however, some floodwaters generated by the 1860 Katla eruption were routed via Sólheimajökull (Björnsson et al., 2000; Hákonarson, 1860). After this event, the eruption centre moved to the northeast, confining jökulhlaups to Kötlujökull and Mýrdalssandur until 1999, when a volcanically-generated jökulhlaup burst from Sólheimajökull (Larsen, 2010; Roberts et al., 2003; Russell et al., 2010). A number of smaller floods generated by geothermal activity are also known to periodically exit the glacier.

Sólheimajökull itself is 8km long, non-surging and characterised as a temperate glacier. It supports a maximum ice thickness of 433m (Kruger et al., 2010; Mackintosh et al., 2002; Russell et al., 2010; Sigurdsson, 2010) and ranges in elevation from 1500 m.a.s.l. down to an over-deepened snout with bed elevation 50m below sea level (Lawler et al., 1996; Le Heron and Etienne, 2005; Russell et al., 2010; Scharrer et al., 2008). The total area of the glacier is approximately 78 km2 with the accumulation zone comprising approximately 56 % of the total glacier area (Lawler et al., 1996; Le Heron and Etienne, 2005; Roberts et al., 2000). Dynamic advance and retreat cycles are exhibited by the glacier (Schomacker et al., 2012), which are frequently asynchronous with other glaciers along the south coast of Iceland (Dugmore and Sugden, 1991). The proglacial geomorphology reflects this dynamic activity, leaving behind a vivid sequence of moraine assemblages and glaciofluvial outwash features (Staines et al., In Press). The supraglacial drainage across the ice-free area of Sólheimajökull is poorly developed due to extensive crevassing and moulin connectivity to englacial and subglacial hydrological pathways. Proglacial channel morphology has evolved from 2009 to 2012, from a configuration of two melt water channels conveying water away from the glacier on the western and eastern side of the catchment respectively, to a well-established proglacial lake which has gradually increased in size from 2010 to the present day (Figure 1). Jökulsá á Sólheimasandi is the bulk meltwater stream leaving the catchment; it has long been known by nearby inhabitants as Fulilaekur or ‘stinky river’ as the smell of hydrogen sulphide is locally prevalent (Thorarinsson, 1939). Jökulsá á Sólheimasandi carries meltwaters from Sólheimajökull as well as two additional stream inputs originating from outside the catchment. The river Jökulsárgil drains Jökulsárgilsjökull, a small valley glacier flowing from Mýrdalsjökull, and enters the Jökulsá á Sólheimasandi proglacial system (Figure 1). Fjallgilsá is a small river of non-glacial origin that drains an area of grassland to the south east of Mýrdalsjökull and joins Jökulsá á Sólheimasandi around 2km down-sandur from the glacier snout. Jökulsá á Sólheimasandi continues to flow in an anastomosing channel, passing under the N1 road bridge 5km from the glacier front where river stage, EC and water / air temperature are automatically logged by the Icelandic Meteorological Office. The geology of the catchment comprises distinct volcanic bedrock units, each thought to predominantly represent a single eruptive episode. Between each unit, striated erosion surfaces support tillites, fluvioglacial deposits and occasional sub-aerial weathering defining periods of landscape denudation and glacial erosion (Carswell, 1983). A detailed chemostratigraphy of each unit is provided in Carswell, 1983, highlighting the dominant basalt lava type found within the majority of units to be classified as a transitional alkalic type. The sulphur content of the rocks within the whole succession has not previously been documented, although the sulphur composition of mixed subglacial till and pyrite-bearing lithologies recently revealed by the receding ice margin are discussed below.

***Field sampling***

Field samples were collected over a period of 4 years from 2009 to 2012, comprising two summer and one spring season sampling campaign. A limited number of samples were also collected in the September of each year. The precise time intervals during which sampling was undertaken, the number of samples collected in each sampling interval, and the temporal comparability of sample collection between seasons are detailed within Tables 1 and 2. Sample collection sites remained at consistent locations throughout the four years, as far as the evolving nature of the proglacial system allowed (Figure 1) and included meltwaters sampled from supraglacial locations, at sites flanking the edges of the proglacial lagoon, subglacial upwellings located at the frontal ice margin, bulk meltwaters located at the outlet of the lagoon and 5 km downstream beneath the N1 road bridge, and streams of external catchment origin. Repeat samples were taken at consistent intervals at each collection site throughout the period of monitoring. At each site, measurements of EC, pH and DO were taken using a handheld multi-parameter WTW 340i combination meter (Wissenschaftlich-Technische Werkstätten GmbH). Water samples were collected and filtered through 0.45 µm cellulose nitrate membrane filters using a handheld Nalgene filter pump. Aliquots were bottled (LDPE Nalgene bottles) without headspace and determination of major ions, cations and trace metals was undertaken within one month of return from the field (only select ions are reported here). Hydrogen sulphide was extracted for isotopic analysis as zinc sulphide, following addition of zinc acetate and collection by filtration (0.45 µm cellulose nitrate membrane filters). For sulphate isotope pre-concentration, 0.5 litres of filtered water was loaded onto a 1 ml volume of ion exchange resin immediately after collection (AG2-X8 and 50W-X8 resin for anion and cation removal respectively). Water samples collected for determination of D/O isotopes were bottled unfiltered and without headspace to minimise the fractionating effects of evaporation.

***Laboratory analysis***

Laboratory analyses of major anions were determined by ion chromatography (Dionex ICS4000 system). Precision based on repeat analysis of reference standard material of comparable concentration to the samples being analysed was calculated as +/-5 %. Limits of detection were calculated based on blank determinations subjected to the same collection and storage procedure as field samples.

In preparation for analysis of sulphate isotopes, anions were removed from the AG2-X8 resin using aliquots of 1M ultrapure HCl to a total volume of 1.5 ml and stored in micro-centrifuge tubes. Addition of 0.2 ml 1M BaCl2 solution to resin eluents enabled precipitation of sulphate as barium sulphate. Samples were allowed to crystallise for approximately 72 hours under refrigeration prior to centrifugation and repeated washing. Product pellets of barium sulphate were oven dried before weighing into tin or silver capsules for δ34S and δ18O analysis respectively.

***Mass spectrometric analysis***

 34S/32S and 18O/16O ratios of product barium sulphate and zinc sulphide were determined by elemental analyser linked to a continuous flow isotope ratio mass spectrometer. Sulphur isotopes from samples collected during the 2009 field season, were analysed using a Thermo Flash EA linked to a Delta XP mass spectrometer at the NERC Isotope Geosciences Facility (NIGL). All other samples were analysed using an Elementar Pyrocube elemental analyser linked to an Isoprime 100 continuous flow mass spectrometer at the University of Lancaster. Combustion of samples within tin capsules in the presence of vanadium pentoxide at 1050 °C yielded SO2 for determination of δ34SSO4 and pyrolysis within silver capsules in the presence of carbon black catalyst at 1450 °C yielded CO for determination of δ18OSO4. δ34S values were corrected against VCDT using within run analyses of international standard NBS-127 and SO5 (assuming δ34Svalues of +21.1 ‰ and +0.5 ‰ respectively and δ18O was corrected to VSMOW using NBS-127 and SO6 (assuming δ18O values of +9.3 ‰ and -11.3 ‰ respectively). Within-run standard replication (1 SD) was better than +/-0.3 ‰ for both sulphur and oxygen isotope values.

Water sample D/H and 18O/16O ratios were determined by continuous flow isotope ratio mass spectrometry at both the University of Lancaster (Elementar pyrocube elemental analyser linked to an Isoprime 100 mass spectrometer; datasets 2010 – 2012) and the University of Birmingham (Eurovector elemental analyser and multiflow linked to a GV Isoprime mass spectrometer; dataset 2009). For D/H determination, both analytical laboratories used sample injection aliquots of 0.3 μl followed by reduction to hydrogen at 1050°C over chromium metal catalyst. δ18O analyses at the university of Lancaster were undertaken in pyrolysis mode using sample injection of 0.4 μl over glassy carbon chips at 1450°C. At the University of Birmingham, analysis was undertaken by equilibration with mixed headspace gas (95% He, 5% CO2). δD and δ18O values were corrected against laboratory calibration standards relative to V-SMOW and presented within the manuscript as non-normalised values. Within-run standard replication (1 SD) was better than +/-1 ‰ for δD in both laboratories. Standard replication for δ18O was better than +/-0.4 ‰ and +/-0.1 ‰ for pyrolysis and equilibration techniques respectively.

**Results**

Sulphate concentrations, isotope values (δ34SSO4 and δ18OSO4, δ18O/DH2O) and physico-chemical data are compiled and presented within Table 1 and 2. A plot of water isotope data (Fig. 2a) differentiates between supraglacial and subglacial meltwater sources during the summer season. However, this source distinction does not prevail during the spring season of 2012, where all water sources cluster tightly within the range dominated by water of supraglacial origin during summer (Fig. 2b). Isotope values presented in Table 1 demonstrate a range of δ34SSO4 between -0.8 ‰ to +7.7 ‰ with these end member values attributed to supraglacial meltwaters and stream waters originating from outside the catchment respectively. Dissolved hydrogen sulphide supports an isotopic composition ranging between +5.8 to +6.2 ‰ (*n* = 5) (Table 1). Bedrock from within the catchment does not contain measurable sulphur concentrations except for a pyrite bearing microcrystalline basalt (approximately 10 % volume of rounded vesicles that are 1-4 mm across and partly filled with cubic pyrite crystals <0.5 mm in length) found within a few key locations at the recently exposed ice margin and within samples of mixed subglacial till. The isotopic signature of this microcrystalline basalt (and mixed subglacial till) ranges between +0.1 to +1.6 ‰ (*n* = 5) and is capable of directly influencing subglacial waters which have undergone rock-water contact. A clear sample site distinction using both sulphur isotope values and concentrations is expressed as a mixing between supraglacial meltwaters, bedrock weathering and stream input of external catchment origin dependent upon seasonality (Fig. 3a,b). Oxygen isotopes within sulphate (δ18OSO4) range between -8 ‰ to +5.5 ‰, and lie either side of the threshold for depicting reducing / oxidising environments (Fig. 4a,b). A pronounced seasonality to the δ18OSO4 signatures depicts summer subglacial conditions, which are of a sub-oxic nature, and spring conditions which are fully oxidised.

**Discussion**

**Seasonal evolution of the glacier drainage system**

The hydrological evolution of Sólheimajökull operates in the fashion generally expected of any temperate glacier system. Discharge is at its lowest in the winter season due to a reduction in surface melting, although the volume of melt produced should be enough to retain open channel flow conditions in the lower portions of the ice mass. Above the winter snowline however, the snowpack thickens with altitude. Such conditions of snowpack storage limit the release of meltwaters to the glacier bed and subglacial hydrology existing beneath an ice mass with a deep snowpack can be expected to remain confined to linked cavity drainage, poorly connected to the channelized system existing at lower elevation (Fountain and Walder, 1998). Deuterium and oxygen isotopes of water would appear to support this hydrological configuration. Altitude has a fractionating effect upon isotopic signatures of δ18O/DH2O such that values become depleted in 18O and D as an air mass ascends to the upper reaches of the glacier (Arnason, 1977). This generates an offset from the Local Meteoric water Line (LMWL) and an altitudinal range of isotopic values in the glacier surface snowpack. The presence of distinctive isotopic values for the supraglacial and subglacial water sources in Fig. 2a would suggest the source waters originate from different altitudinal ranges. Supraglacial waters constitute some of the heaviest isotopic values on the local meteoric water line. Due to the crevassed nature of the glacier surface, supraglacial streams sampled in the lower reaches of the glacier are likely to be locally sourced and thereby originating from low elevation. The light isotopic end member on Fig. 2a is composed of upwelling waters of subglacial origin, likely representing snow / ice melt originating from higher elevation (cf. Griselin et al., 1994). During the spring season of 2012, this light isotopic end member is not apparent due to the absence of any subglacial upwelling waters and the glacier drainage system still being spatially restricted to the low elevations (Fig. 2b, Table 1). Enhanced summer season melting and headward expansion of drainage channels will subsequently re-establish a high volume subglacial drainage network penetrating beneath the accumulation zone and transporting meltwaters of high elevation origin.

**Sources of sulphur in the Sólheimajökull system**

Key sources of sulphur in the Sólheimajökull system during summer 2010 are depicted through Fig. 3a. Key inputs to the system consist of supraglacial meltwater, sulphur obtained from bedrock weathering, dissolved hydrogen sulphide and sulphur within streams that enter from outside the catchment. Mixing between these sources in variable proportion dominates the sulphur isotopic composition in the main meltwater streams.

Summer season Supraglacial runoff from the glacier is necessarily low in sulphate concentration, reflecting the dilute nature of ice melt. The isotopic values are very similar to those found in the catchment bedrock (catchment bedrock = +0.1 to +1.6 ‰) and probably reflect either bedrock dust deposition and subsequent release through surface weathering, or dissolution of volcanic aerosol (~ 0‰ (Nielson, 1974)) attached to ash particulates deposited on the surface of the glacier (e.g. Flaathen and Gíslason, 2007; Olsen et al., 2013). Despite the close proximity of the ocean, there is little isotopic evidence for any marine input (~ +21 ‰ (Rees et al., 1978)). This contrasts markedly with the work of and Gíslason and Torssander (2006) in which the main component of Icelandic winter precipitation is seen to be marine derived (δ34SSO4 = +13 to +19 ‰). This contrast is likely due to volcanic and bedrock sources detailed above being trapped during snowpack deposition, dominating the meltwater composition when released during melt. Streams originating from outside the catchment form another end-member to the mixing plot, forming some of the heaviest sulphur isotopic signatures (up to +7.7 ‰). These are likely obtained through contribution from soil and groundwater leachates during passage through high elevation pastures prior to entry into the Sólheimajökull catchment. The deposition of ash from the eruption of Eyjafjallajökull in May 2010 and its entrapment in surface vegetation is likely the cause of lighter isotopic values in streams of external catchment origin in summer 2010 compared to spring 2012. Further biogeochemical controls on the sulphur isotopic composition of waters emanating from vegetated soils include processes of assimilation (uptake) of sulphur into vegetation, re-mineralisation of organic matter, and long term storage of sulphur in both vegetation and soil. During processes of assimilation and mineralisation, a slight preference for incorporation of 32S into the reaction products creates a fractionation of 1-2 ‰ (Krouse et al., 1991; Mitchell et al., 2001; Thode et al., 1991). Net uptake (assimilation) of sulphate by vegetation typically leaves soil sulphate solutions slightly enriched in 34S (Zhang et al., 1998) and hence explains why the streams of external catchment origin form a sulphur isotopic end member enriched in 34S relative to that released during snow and ice melt. Sulphur isotopic signatures obtained from the catchment bedrock match closely with those obtained from meltwaters of subglacial origin (sampled directly from upwelling zones), suggesting bedrock weathering forms a third end-member to the system (Table 1 and Figure 3a). Sulphur isotopic signatures collected from the bulk meltwater output (denoted ‘mixed meltwater outlet’ in Figure 3) mainly plot between these three end member signatures, representing input from only three of the four possible sources. The fourth source (oxidation of dissolved hydrogen sulphide, δ34S = +5.8 to +6.2 ‰) seems to hold only a weak influence over meltwater sulphate isotopic composition. Despite glacial meltwaters previously recorded to contain concentrations of H2S up to 3 ppm during the summer season (Lawler et al., 1996; Sigvaldason, 1963), this sulphur source does not seem to dominate the isotopic composition of the sulphate molecule. Optimal conditions promoting the oxidation of aqueous H2S require pH 6.8-7.5. However, the meltwater pH of subglacial and bulk discharge is frequently below this value and a large proportion of residual oxygen is required to enable efficient oxidation (dissolved oxygen concentrations are typically below 100 % during the summer season, Table 1). Rates of hydrogen sulphide oxidation are also limited by low temperatures. Environmental conditions are therefore not conducive to rapid oxidation of H2S. Whilst conditions of < 100 % DO may be limiting to the oxidation of aqueous H2S, oxidation of pyrite can proceed under sub-oxic conditions through the use of alternative oxidising agent FeIII under microbial catalysis (Equation 2), hence sulphate concentrations within the catchment meltwaters remain dominated by those obtained through mineral weathering and atmospheric deposition.

The chemical composition of subglacial water does not remain constant throughout the summer melt season but changes with subglacial hydrological evolution and source mixing. Subglacial waters periodically upwell along lines of structural weakness where ice is thin close to the margin (e.g. along remnants of thrust planes, and at the head of debris stripes), initially under pressure with high concentrations of sulphate and an isotopic composition similar to the surrounding bedrock. Sulphate content becomes more dilute with time (days) and isotopically similar to bulk meltwaters leaving the catchment (Fig. 3a). This evolution of δ34SSO4 likely represents initial drainage of isolated parts of the glacier bed, associated with increasing water pressures in distributed cavities. Subsequent chemical and isotopic dilution demonstrates mixing with other water sources (e.g. supraglacial recharge, stream waters originating from outside the catchment mixing beneath the floating dead ice area at the glacier snout) as the drainage system evolves. This is typical of the chemical switch depicted within Bottrell and Tranter, (2002) and Wynn et al. (2006).

Samples collected for sulphur isotopic analysis at the road bridge, representing the outer limits of the sampled catchment approximately 5 km from the glacier snout, reside partly outside the mixing plot of Fig. 3a. This could be due to downstream oxidation of remaining hydrogen sulphide to sulphate, driven by stream turbulence and an increase in water temperature. On-going reactivity of suspended sediment and contributions from varying lithology throughout the catchment may also drive this subtle shift in isotopic composition. Alternatively, extra-glacial stream Fjallgilsá enters the main meltwater channel 3km upstream from the bridge sampling site and has been shown in subsequent years to convey sulphate at lower concentrations, but enriched in 34S compared to the main stream (Table 1). Either way, it is apparent that solute loads and isotopic determinations of sulphate undertaken at the edge of the catchment are not truly representative of processes occurring beneath the glacier and ice cap, but are associated with downstream mixing of water sources and geochemical evolution away from point of emergence from the subglacial environment.

In the absence of a subglacial water source in Spring 2012, bulk meltwater outputs are dominated by mixing between supraglacial meltwater and streams of external catchment origin (Fig 3b). The Jökulsárgil water source of external catchment origin dominates the chemical composition of the mixed outlet sampling site, demonstrating the importance of this input as an end member of the Jökulsá á Sólheimasandi system in the spring season. A discernable evolution of the water composition to isotopic values slightly enriched in 34S and of lesser sulphate concentration between the mixed outlet and bridge sampling site may be attributed to inputs from the Fjallgilsá stream, also of external catchment origin. It is possible that oxidation of small quantities of dissolved H2S may also form an end member of the mixing plot, thereby accounting for enrichment in 34S and increases in sulphate concentration. However, no detectable presence of H2S, would preclude this possibility. Furthermore, the lack of a subglacial water source due to the restricted nature of the subglacial drainage system at this time of the year, is demonstrated through the restricted range in δ18OH2O (Fig. 2b) and as such would prevent access to any subglacial source of H2S.

**Hydrological controls on subglacial geothermal volatile release**

During the summer season, large portions of the glacier bed are drained by channels operating under atmospheric pressure. During times of minimal surface ablation (winter-spring), reduced meltwater discharge should dictate that large channels are squeezed closed through ice deformation and drainage becomes confined to linked cavities, which are comparatively isolated from any atmospheric source (Fountain and Walder, 1998). Commensurate with a switch in hydrological configuration, a change in the redox status of drainage pathways would also be expected. Under such a model, winter season subglacial conditions should be driven towards a reducing status as weathering reactions consume all available atmospheric gases, whilst summer season conditions are closer to fully oxidising.

However, subglacial redox status at Sólheimajökull appears inverse to this expected pattern, supported through signatures of sulphate-oxygen isotopes. During the summer season, the δ18OSO4 signature of meltwaters which are at least partially sourced from the glacier bed, indicate the attainment of low redox status, calculated through equation 3 following Bottrell and Tranter, (2002). Full reducing, closed system conditions (eg, Hartmann and Nielsen, 2012) and the activity of microbially mediated sulphate reduction are not in evidence as δ18OSO4  values are far from exchange equilibrium with water (Fritz et al., 1989). Figure 4b demonstrates oxidizing conditions during the winter season, indicative of a dominating influence from channelized subglacial flow conditions which are well-connected to the atmosphere and draining meltwaters from below the winter snowline. The presence of low redox conditions during the summer season is counter-intuitive to expectations of the hydrological system in operation and demands a mechanism of either removing oxygen from the system, or for a source of anoxic water to contribute to the meltwater discharge.

The conceptual model shown in Figure 5 proposes a suite of controls determining the seasonal presence of meltwaters of low redox status. We propose that a combination of periodic increases in basal water pressure and summer season headward expansion of channelized drainage, enables connectivity to be established with a zone of subglacial geothermal activity beneath Mýrdalsjökull which discharges anoxic waters. The strength of this geothermal field is likely controlled through seasonal changes to the ice overburden pressure. Numerous cauldrons across the surface of Mýrdalsjökull, some of which reside within the Sólheimajökull catchment, provide surface expression of subglacial geothermal activity (Björnsson et al., 2000). High ice over-burden pressures and the inward-dipping ice surface topography at cauldrons would likely conspire to prevent continuous meltwater escape from the locale (Björnsson 1988, Guðmundsson et al. 2004). Only when subglacial water pressures become great enough will ice over-burden pressures be exceeded and / or hydraulic fracturing enable the escape of anoxic water. The onset of the summer melt season has the effect of establishing seasonal hydraulic connectivity with the anoxic geothermal zone, thereby encouraging the release of confined geothermal meltwaters charged with reduced gases (including hydrogen sulphide). These waters travel through both subglacial cavities and channels and upwell at the glacier snout throughout the summer season (Fig. 5a). The reduction in water pressure during transition from high pressure linked cavities to low (atmospheric) pressure discrete channelized drainage will also encourage the degassing of H2S initially emplaced under elevated overburden pressures. The limited oxygen content of the water drives bedrock sulphide oxidation utilising FeIII as an oxidising agent and dominates the chemical composition of the bulk meltwaters leaving the glacier catchment. The effects of bacterial disproportionation of sulphur intermediate compounds on the meltwater sulphate isotope content of meltwaters cannot be discounted (Bottcher et al., 2005), although the enrichment of δ18OSO4 compared to precursor water values is not great enough to unequivocally invoke this process. Low temperatures, low pH and below-saturated dissolved oxygen content (measured at the point of upwelling) all serve to minimise any oxidation of H2S gas. A further possible effect of summer reductions in overburden pressure, due to the combination of summer snowmelt and the switch to a low-pressure subglacial drainage system, is to stimulate geothermal activity. It has been proposed that seasonal variations in snowpack thickness cause stress readjustments that favour eruptive activity in the summer / late summer months (Albino et al., 2010), consistent with preference for summer / late summer eruptions during historical activity at Katla (Thordarsson and Larsen, 2007). The Katla geothermal-volcanic system therefore appears highly sensitive to modest pressure changes (Albino et al., 2010). However, minimal change in the size of the surface cauldrons on a seasonal basis would suggest that the strength of geothermal activity is predominantly seasonally invariant, as increased summer geothermal activity would be expected to drive deepening of surface depressions (e.g. Guðmundsson et al., 2004). Evidence for anoxia is present in the samples collected in September of each year (Table 2), sometime after which the commencement of high elevation snowfall and the restriction of the subglacial hydrological system to lower elevations will cut off the supply of anoxic geothermal water and fully oxic conditions will return (Fig. 5b). An alternative process capable of generating δ18OSO4 values close to those of δ18OH2O, involves high rates of oxygen isotopic exchange between sulphate molecules and surrounding water under conditions of high temperature and low pH (Lloyd, 1968). However, geothermal sulphur sources hold only a weak influence over isotopic signatures of sulphate (Fig 3a) and it is likely that temperatures of melt waters at the ice-bed interface will not be high enough to promote oxygen isotopic exchange between water and sulphate obtained through bedrock weathering. Therefore, in our conceptual model (Fig. 5) the geochemical signature of meltwater is predominantly driven by hydrological reconfiguration. Only when the subglacial drainage system expands to connect with geothermal areas, do sulphate isotopes form sensitive indicators of subglacial geothermal activity through redox status imprinted in the sulphate molecule. Seasonal reconfiguration of meltwater drainage above the Katla geothermal field, may create shifting abundance and pressure of meltwater at the glacier base. This could also drive seasonal patterns of shallow low-frequency seismicity and pressure that encourage seismogenic slip at the ice-bedrock boundary in summer months (Jonsdottir et al. 2007, 2009).

**Conclusions**

Stable isotopes of sulphate and sulphide have been used with the ultimate aim of establishing a geochemical fingerprint of subglacial geothermal activity beneath the Mýrdalsjökull ice cap. The ubiquitous nature of sulphate in most glacial meltwaters and the expectation for rapid oxidation of reduced sulphurous gases, make the sulphate molecule an attractive contender as an indicator of subglacial geothermal processes. However, sulphur isotopes demonstrate that sulphate in the bulk meltwaters leaving the Sólheimajökull catchment can be accounted for through a three way mixing of sources, including oxidation of bedrock sulphides, supraglacial melt and stream water inputs of external catchment origin. Aqueous hydrogen sulphide, suspected to be of geothermal provenance does not appear to be readily oxidised to sulphate due to sub-optimal environmental conditions of pH, temperature and dissolved oxygen content. Sulphate and aqueous sulphide thereby remain two separate components of the sulphur biogeochemical cycle. However, oxygen isotopes within the sulphate molecule provide a strong sense of redox status, which demonstrate seasonal variation opposite to that expected in alpine and arctic glaciers which do not reside over a geothermal field. At Sólheimajökull, anoxic conditions prevail during the summer when drainage systems are fully developed and would normally be expected to convey waters under fully oxygenated conditions, suggesting a substantial input of reduced gases of geothermal origin. Under winter conditions, the waters are fully oxygenated suggesting limited geothermal contribution and an open hydrological system resulting from year round melting on the lower portions of the glacier. Seasonality of the δ18OSO4 geochemical fingerprint is considered the result of a strong summer geothermal contribution associated with subglacial drainage reconfiguration, and unloading due to surface melting. This is also likely closely linked with the seasonal patterns of shallow seismicity detected at Katla, and the tendency for geothermal events and eruptions to occur during the summer months. Oxygen isotopes of sulphate are thereby deemed to provide a sensitive indicator of subglacial processes, recording shifts in hydrological connectivity at a large geothermal system that is entirely obscured by ice. Further development of this conceptual model is now required in order to build a better understanding of the role of the overlying ice mass upon mediating subglacial geothermal activity and the controls exerted upon the release of dissolved volatiles through meltwater streams.

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**Figure captions:**

**Figure 1:** Location map showing the Mýrdalsjökull ice cap and outlet glacier Sólheimajökull (Fig. 1a). The main sampling sites and proglacial drainage configuration during 2009-2010 are depicted in Fig. 1b and during 2012 in Fig 1c.

**Figure 2:** Plot of δ18O/DH2O for meltwater samples collected during summer 2010 (Fig. 2a) and Spring 2012 (Fig. 2b). The Global meteoric Water Line (GMWL) is plotted as δ2H = 8.17 δ18O+11.27 (after Rozanski et al., 1993). The Local Meteoric Water Line (LMWL) is sourced from GNIP station Reykjavik (64**°** 07’ 48’’, 21**°** 55’ 48’’; 14 m.a.s.l).

**Figure 3:** δ34S Vs [SO4] for sulphur sources and bulk meltwaters within the Sólheimajökull catchment. Data displayed are from summer 2010 (Fig. 3a) and Spring 2012 (Fig 3b).

**Figure 4:** δ18OSO4 Vs calculated threshold for oxidation of sulphide under reducing conditions. Threshold calculations are depicted through equation 3 (after Bottrell and Tranter, 2002). Data displayed are from summer 2010 (Fig. 4a) and spring 2012 (Fig. 4b).

**Figure 5:** Conceptual diagram showing the relationship between hydrological evolution and overburden pressure release leading to seasonality in meltwater geothermal indicators. Fig. 5a) Spring season configuration showing discrete channelized subglacial hydrology limited to the lower portions of the glacier, a large ice and snow overburden pressure and limited connectivity to the zones of geothermal activity. Fig. 5b). Summer season configuration depicting a well-developed channelized subglacial hydrological system which accesses the geothermal zone. Reduced overburden pressure enhances geothermal activity, heat exchange and volatile release.

**Table 1:** Sulphate concentrations, isotope values (δ34SSO4 and δ18OSO4, δ18O / δ2HH2O) and physico-chemical data for Sólheimajökull, during spring and summer field campaigns 2009-2012.

--- = data unavailable. Values represent averages with *n* value provided. Data range provided in parentheses.

**Table 2:** Sulphate concentrations, isotope values (δ34SSO4 and δ18OSO4, δ18O / δ2HH2O) and physico-chemical data for Sólheimajökull, during September 2009-2012.

\*δ18OSO4 threshold has been calculated using δ18OH2O values calculated as an average of the summer season mixed outlet.

\*\*δ18OSO4 threshold has been calculated using δ18OH2O values calculated as an average of the summer season mixed outlet from 2009 and 2010.

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