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Glacial Erosion Liberates Lithologic Energy Sources for Microbes and Acidity for Chemical Weathering Beneath Glaciers and Ice Sheets

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Wet-based regions of glaciers and ice sheets are now recognized to host unique and diverse microbial communities capable of influencing global biogeochemical cycles. However, the isolated nature of subglacial environments poses limitations upon the supply of protons for chemical weathering and energy sources (electron donors/acceptors) to support in situ microbial communities. A less well recognized source of these substrates is the release of gases from mineral structures, pore spaces or fluid inclusions and the generation of gases from the breakage of mineral bonds during the mechanical breakdown of rocks by moving ice. Here, we investigate the potential release of H2, CO2, CO, and short chain hydrocarbons, particularly CH4, by glacial erosion at rates relevant to chemical weathering and microbial activity beneath glaciers. A wide range of magmatic, metamorphic, and sedimentary rocks, and subglacial sediments from glaciated catchments in Greenland, Norway and Canada were ground in the laboratory to varying grain sizes and the release of gases was measured. The volume of gas released increased as the grain size of the ground sediments decreased. The results of these laboratory experiments were used to estimate rates of catchment-scale gas release based upon estimates of long term abrasion rates at each glacier. H2 generation was calculated to be sufficient to potentially support previously estimated rates of methanogenesis in the upper centimeters of subglacial sediment at a gneissic catchment in Greenland and a sedimentary catchment in Canada. Sufficient CO2 could be released by grinding to drive as much as 20% of subglacial chemical weathering at a metamorphic catchment in Svalbard, with potential implications for the inferred quantity of CO2 drawn-down from the atmosphere by glacial weathering. Rates of CH4 generation from grinding bedrock has the potential to be greater than subglacial microbial generation in a sedimentary catchment in Canada with carbon rich bedrock, suggesting a potentially important source of CH4 for methanotrophic microorganisms. We conclude that mechanical erosion beneath a range of glaciers generates significant quantities of gases which have the potential to enhance chemical weathering and/or support subglacial microbial communities in the deep icy biosphere.

Keywords: subglacial environment, rock grinding, chemical weathering, CO2 drawdown, hydrogen, methane, microbial energy source, gas generation
INTRODUCTION

Subglacial environments have become of increasing interest for understanding how microbial life survives in cold, dark, environments, and for investigating the impact of glaciation upon global biogeochemical cycles (e.g., Skidmore et al., 2000; Christner et al., 2012; Wadham et al., 2013). Despite darkness, high pressures, low temperatures, and assumed low nutrient input, the subglacial environment hosts active microbial ecosystems (e.g., Sharp et al., 1999; Christner et al., 2014; Dieser et al., 2014). The ubiquitous presence of geochemically reactive rock flour, coupled with isolation from surface inputs of organic matter and complete darkness, suggests that chemolithotrophic communities are important primary producers in subglacial systems (Sharp et al., 1999; Tranter et al., 2002; Boyd et al., 2014; Christner et al., 2014). Chemolithotrophic microbes exploit chemical disequilibria to derive their energy, often utilizing dissolved gases as electron acceptors or donors (Lovley and Goodwin, 1990; Sharp et al., 1999). For example, hydrogenotrophic methanogens can subsist upon dissolved gases (i.e., H₂/CO₂) as their sole energy source (Lovley and Goodwin, 1990). However, due to the isolation of areas of the subglacial environment from surface processes and the atmosphere, there is a limited supply of energy sources resulting in their depletion over time, particularly during long periods of glaciation (Wadham et al., 2004).

Similarly, the proton pool available to fuel chemical weathering reactions can become limited during long periods of isolation from the atmosphere and surface processes (Hallet et al., 1996). Important proton sources in subglacial environments include carbonic acid generated from the dissolution of atmospheric and microbially-respired CO₂ and the oxidation of sulphide minerals under both oxic and anoxic conditions (Tranter et al., 1996, 2002; Sharp et al., 1999; Montross et al., 2013). However, vast areas beneath ice sheets are isolated from the influx of surface waters carrying atmospheric CO₂. Varying rates of basal melt across the bed impacts the volume of palaeo-atmospheric CO₂ released from bubbles trapped in the ice, limiting this source in low melt areas. Proton supply via sulphide oxidation is only significant where there are sulphide-bearing rocks being physically eroded to comminuted rock flour. This source may become, at least locally, limited. Microbially-respired CO₂ has been shown to enhance chemical weathering rates (Montross et al., 2013), but would be limited in areas where there are limited growth substrates (e.g., low organic carbon) and subsequently lower rates of respiration.

We postulate that a previously unrecognized abiogenic source of protons (through the generation of carbonic acid) and microbial energy is the bedrock, which is eroded at rates of 0.01–100 mm yr⁻¹ by moving ice in wet-based areas via fracturing, plucking and abrasion (Hallet et al., 1996; Lee and Rutter, 2004; Cowton et al., 2012). Tentative evidence from the literature supports this notion. Gases trapped within rocks can be released by mechanical grinding (Zhang et al., 2014; Deeds et al., 2015; Wang et al., 2015). For example, crystalline rocks, such as granitoids, which constitute the major basement rocks beneath the Greenland Ice Sheet, contain gases (e.g., hydrocarbons) trapped within their minerals’ crystal structure or within fluid inclusions (Potter and Konnerup-Madsen, 2003). Sedimentary rocks can retain even larger volumes of hydrocarbons and CO₂ within their pore spaces and mineral structures (e.g., Martinelli and Plescia, 2005; Osborn and McIntosh, 2010; Wang et al., 2015). In addition, gases such as H₂ can be generated from mechanical grinding of rocks and minerals through fault action (e.g., Ito et al., 1999; Kameda et al., 2004). Despite the potential for glacial erosion to generate these gases, there has been only one experimental investigation, focusing solely on H₂ generation, using field-collected samples from glacial catchments to date (Telling et al., 2015).

This study aims to investigate the potential of subglacial mechanical erosion as an abiogenic source of gases, at generation rates that are relevant to chemical weathering and microbial activity. Rock and sediment samples from a range of lithologically contrasting (igneous, metamorphic, and sedimentary rock types) glaciated catchments across the Northern hemisphere were ground in a ball-mill to simulate the process of glacial abrasion. The release of H₂, CO₂, CO, and several short-chained hydrocarbons was quantified to determine their significance for providing protons for chemical weathering via carbonic acid generation and as electron acceptors and donors for microbial energy.

MATERIALS AND METHODS

Rock and Sediment Sample Selection

Rock samples were collected from a diverse range of glaciated catchments in the Northern hemisphere. These glaciers were Mittivakkat (MG) and Leverett (LG) in Greenland, Engabreen (EG) and Midtvei Lovenbreen (ML) in Norway, and Robertson (RG) in Canada. The details of the catchment location, general lithology and rock sample type(s) are shown in Table 1. Glacial sediment samples were also obtained at two catchments (LG and ML). The LG sediment was melted out of a basal ice sample. The ML sediment was collected immediately in front of the glacier terminus and had thus been recently exposed. Approximate percentage areal composition of five major minerals (quartz, feldspar, biotite, muscovite, and pyroxene) was determined for the crystalline rock samples (Table 1). Identification was conducted using established physical properties of minerals that were visible with the naked-eye or with a hand-lens (x10 magnification). Mineral quantification was not conducted for the sediment samples or the RG rock samples because they were too fine-grained to identify by this method.

Experiments to Simulate Gas Release via Glacial Erosion

A sledgehammer on a metal plate was used to reduce the rock samples in size. To minimize contamination, samples were wrapped in durable polyethylene bags and the plate and hammer were cleaned between samples with ethanol (100%). The 250 µm–2 mm fraction was collected and dried at 105°C for 48 h to remove moisture prior to milling. The rock and sediment samples were ground to finer powders in a zirconium oxide ball-mill sealed with a PTFE O-ring. We advocate that this process is
The ball-mill was present).

of major rock forming minerals, quartz (Q), feldspar (F), muscovite mica (Mu), biotite mica (Bt) and pyroxene (Py) (NB totals may not be 100 % where other minerals were present).

<table>
<thead>
<tr>
<th>Glacial catchment</th>
<th>Location</th>
<th>Catchment lithology</th>
<th>Sample type(s)</th>
<th>Mineral composition (areal %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MG</td>
<td>Southeast Greenland</td>
<td>igneous &amp; metamorphic</td>
<td>porphyritic granite</td>
<td>Q 60  F 15  Mu 2  Bt 15  Py n.d.</td>
</tr>
<tr>
<td>LG</td>
<td>West Greenland</td>
<td>basement rock (metamorphic)</td>
<td>orthogneiss</td>
<td>Q 35  F 30  Mu n.d.  Bt 5  Py 30</td>
</tr>
<tr>
<td>EG</td>
<td>Norway</td>
<td>metamorphic</td>
<td>mica-schist</td>
<td>Q 10  F 5  Mu 55  Bt 15  Py n.d.</td>
</tr>
<tr>
<td>ML</td>
<td>Northwest Svalbard</td>
<td>metamorphic</td>
<td>schist</td>
<td>Q 30  F 50  Mu 50  Bt 10  Py n.d.</td>
</tr>
<tr>
<td>RG</td>
<td>Alberta, Canada</td>
<td>sedimentary</td>
<td>hard* muddy carb.</td>
<td>Q –  F –  Mu –  Bt –  Py –</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>soft* muddy carb.</td>
<td>Q –  F –  Mu –  Bt –  Py –</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>shale</td>
<td>Q –  F –  Mu –  Bt –  Py –</td>
</tr>
</tbody>
</table>

(-) sediment samples and the rock samples of RG were too fine grained for the method of identification (Rock and sediment sample selection). (n.d.) mineral not detected.

(*) “hard” and “soft” describes the “hardness” of the two samples relative to each other based on how quickly the starting sample clasts were powdered during grinding. “muddy carb.” is short for muddy carbonate.

analogous to glacial abrasion where clasts frozen into the glacier’s base grind the bedrock beneath, producing large quantities of fine-grained sediment (e.g., Hallet et al., 1996). The ball-mill was modified with two valves so it could be flushed with 5.0 grade Argon, which was dried with an in-line moisture trap (Agilent), prior to grinding. Approximately 10 g of sample was ground in triplicate for 1, 3, 5, 10, and 30 min using a ball mill (Fritsch Planetary Mono Mill Pulverisette 6) at 500 rpm. The same length of grinding produced different mean grain sizes, depending upon the rock type, and thus comparison between the samples was standardized to mean grain size measured by Mastersizer. Grinding beyond 30 min did not produce smaller grain sizes. Immediately after grinding, 5 mL of 5.0 grade argon was injected into the ball-mill (to maintain ambient pressure) using a gas-tight syringe and mixed with the headspace before removing 5 mL of sample for analysis. Quartz mineral specimens (purchased from thegeologysuperstore.com) were used as a procedural blank to ensure that no gases were generated by the ball-mill during grinding. The quartz minerals were pre-crushed and sieved, by the same method as the samples, before being furnace at 1000°C for 2 h, in order to dissociate gases from the mineral surface and release gases trapped within fluid inclusions. Grinding of the prepared quartz was conducted in duplicate for 1, 10, and 30 min. Methane was the only gas detected in the blank runs and averaged at 0.01 nmol CH₄ g⁻¹ (standard deviation of 0.002 nmol g⁻¹) across all the blank runs. Minor (0.3 nmol g⁻¹) carbon dioxide was present in the Argon gas used to flush the headspace. No other gases were detected in the procedural blank.

### Analysis of Gases Released During Grinding

Gas samples were injected into an Agilent 7890A gas chromatograph (GC) fitted with a methaniser (at 395°C) and two detectors: an FID (flame ionizing detector, at 300°C) and a TCD (thermal conductivity detector, at 250°C). Separation of methane (CH₄), carbon dioxide (CO₂), ethane (C₂H₆), ethylene (C₂H₄), and propane (C₃H₈) was achieved using a molecular sieve 5A柱, 80–100 mesh, 8 ft × 1/8-inch column. Separation of hydrogen (H₂) was achieved using a Haysep D 80–100 mesh, 2m × 1/8-inch SS column. The columns were held at 25°C for 4 min, before being ramped at 50°C per min to 200°C.

Gas samples obtained from grinding rock samples from two catchments with contrasting geologies (sedimentary RG and granitic MG) were also analyzed for carbon monoxide (CO) on an SRI 8610C GC fitted with a molecular sieve 5A column at 30°C and a reduction gas detector (mercuric oxide with UV lamp) held at 295°C.

Calibration standards were run twice daily on both GCs, apart from ethylene, which was calibrated with a standard obtained later in the experimental period as the peak was initially unidentified. The percentage variance, limit of quantification and detection for the standards are displayed in Table 2. Concentrations of the samples were calculated from a linear regression line of manual dilutions of certified (+/- 5 %) standards with 5.0 grade Argon. Gas concentrations were converted to moles using the Ideal Gas Law, corrected for dilution, and normalized to the mass of dry sediment.

### Grain Size and Total Carbon Analysis

The grain size after grinding was measured to standardize comparison of the samples. Mean grain size was measured using a Malvern Mastersizer 3000 which utilizes laser diffraction to estimate a mean grain size from 5 replicates per sample. The mean percentage variation for the 5 replicates was 3.5 % (n = 200). An Elemental Analyser (EA) 1110 was used to measure percentage weight of total carbon in an 8 to 19 mg, <250 µm, well-mixed aliquot of each sample which.

<table>
<thead>
<tr>
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<td></td>
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<td>soft* muddy carb.</td>
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</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>shale</td>
<td>Q –  F –  Mu –  Bt –  Py –</td>
</tr>
</tbody>
</table>
TABLE 2 | The limit of quantification (variance) and detection (LOD) for each gas analyzed.

<table>
<thead>
<tr>
<th>Gas</th>
<th>LOD, ppm</th>
<th>LOD, nmol g⁻¹ equivalent</th>
<th>Variance nmol g⁻¹ equivalent</th>
<th>% variance (n = 48)</th>
<th>Standard concentration, ppm (+/- 5 %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H₂</td>
<td>0.3</td>
<td>2.0</td>
<td>2.0</td>
<td>1.8</td>
<td>493</td>
</tr>
<tr>
<td>CO₂</td>
<td>0.2</td>
<td>1.5</td>
<td>1.5</td>
<td>1.7</td>
<td>406</td>
</tr>
<tr>
<td>CH₄</td>
<td>0.1</td>
<td>0.6</td>
<td>0.6</td>
<td>0.8</td>
<td>195</td>
</tr>
<tr>
<td>C₂H₆</td>
<td>0.1</td>
<td>0.2</td>
<td>0.2</td>
<td>0.1</td>
<td>19</td>
</tr>
<tr>
<td>C₃H₆</td>
<td>0.3</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>104</td>
</tr>
<tr>
<td>C₂H₄</td>
<td>0.02</td>
<td>0.04</td>
<td>&lt;0.04</td>
<td>0.03</td>
<td>2.2</td>
</tr>
<tr>
<td>CO</td>
<td>0.2</td>
<td>0.02</td>
<td>0.02</td>
<td>0.2</td>
<td>12</td>
</tr>
</tbody>
</table>

Serial dilution was not conducted for CO.
^ for ethylene, n = 7.
^^ for CO, n = 24.

was flash heated to 1000°C. The EA was calibrated with a certified Aspartic acid standard containing 36.14 % total carbon (n = 3, per 20 samples). Reproducibility of the total carbon content analyses was determined using a soil reference standard containing 2.36 % total carbon with the precision at 0.07 wt % total carbon (n = 4). The limit of detection was 0.01 wt % total carbon.

Suspended Sediment Flux and Discharge Analysis for ML (2016) and RG

To compare the potential gas generation by glacial erosion between catchments the rate of gas production was scaled up to a “catchment-rate”. To conduct this calculation, the flux of suspended sediment (SS) was used. We assumed that this is approximately equivalent to the rate of mechanical erosion; a common method used to calculate erosion rates (e.g., Hallet et al., 1996) and that which is likely to be representative of the average erosion rate over longer timescales despite the potential for errors on an annual basis. For the 2016 ML melt season and the 2014 RG melt season, this was calculated here using the product of the SS concentration and discharge measurements. The SS flux at ML was measured at the eastern subglacial upwelling (78.8956°N, 12.0730°E). There was no evidence of a western subglacial upwelling forming that year, as has occasionally been common method used to calculate erosion rates (e.g., Hallet et al., 1996). SS concentration was measured (n = 16) every 3–4 days from the 24th June to the 21st July 2016. Water was collected in a rinsed 1 L Nalgene bottle and 100–500 mL (depending on the perceived turbidity) was filtered using a hand-pump with a Nalgene filtration tower onto a 0.45 µm cellulose nitrate filter paper. The paper was dried at 50°C for approximately 24 h and the dry weight of sediment per liter of water calculated and averaged (Table 3). The average grain size of the SS samples was 18.4 µm (n = 7; section Grain size and total carbon analysis). The discharge (m³ s⁻¹) was calculated for 9 days over the melt season from water velocity (m s⁻¹) and water depth (m) measured at 5–7 points across the width (m) of the river (number of measurements varied with river width). This was extrapolated across the estimated number of days the subglacial upwelling was flowing (47 days) to give a total melt season discharge of approximately 6.1 × 10⁶ m³. The product of the SS concentration and discharge provided an estimated SS flux of 95 × 10⁸ g during the summer melt season, the same order of magnitude as that reported by Hodson et al.’s (2000) study.

Here, we present the first data for SS flux and bulk meltwater discharge for RG. An average SS concentration for RG of 0.19 g L⁻¹ was obtained by the same method as described for ML from measurements taken on 3 days (n = 5/day), spread over the 2014 melt season. The average SS grain size was measured as 4.0 µm (n = 7; section Grain size and total carbon analysis; Table 3). The SS concentration was multiplied by an estimated average discharge of 0.3 m³ s⁻¹ over a 62-day melt season. This discharge is the average for July during the 2010 RG melt season, and the assumed average for August as both had similar numbers of positive degree days (PDD; 310 cf. 320) (Doxsey-Whitfield, 2012; Scanlon, 2017). We assumed that the average discharges in 2010 and 2014 were comparable, as the July and August 2010 PDD values were close to the 2006–2012 average for July (350) and August (340), and thus appear to be broadly representative (Scanlon, 2017).

TABLE 3 | Suspended sediment (SS) flux, average concentration and average grain size in meltwater for each glacier catchment.

<table>
<thead>
<tr>
<th>Glacier</th>
<th>SS flux (10⁸ g a⁻¹)</th>
<th>SS flux year(s)</th>
<th>SS conc. (g L⁻¹)</th>
<th>SS grain size (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MG</td>
<td>176</td>
<td>2005¹</td>
<td>0.5¹</td>
<td>23.2²</td>
</tr>
<tr>
<td>LG</td>
<td>29,000</td>
<td>2009-2012²</td>
<td>1.1³</td>
<td>20.7⁴</td>
</tr>
<tr>
<td>EG</td>
<td>165</td>
<td>1987, 1989-1993⁵</td>
<td>0.1⁶</td>
<td>23.2²</td>
</tr>
<tr>
<td>ML</td>
<td>95</td>
<td>2009⁷, 2010⁸⁷</td>
<td>1.2⁸</td>
<td>25.7⁷,⁸</td>
</tr>
<tr>
<td>RG</td>
<td>3</td>
<td>2010, 2014⁸⁷</td>
<td>0.2⁸</td>
<td>4.0⁸</td>
</tr>
</tbody>
</table>

Where data for multiple melt seasons is available, SS flux is an average. SS grain size is the average measured in that catchment, (*) denotes data not available, therefore, the average of the non-sedimentary catchments (LG and ML) was used. Year of the season used to calculate SS flux is shown and this, SS concentration and SS grain size are from published values or obtained here.¹ Hoshizaki and Merfield, 2006;² Hawkins et al., 2015;³ Hawkins et al., 2016;⁴ Boge, 2014;⁵ Boge, 1996;⁶ Boge, 1996;⁷ Hodson et al., 2015;⁸ Hodson et al., 2004; this study (section Suspended sediment flux and discharge analysis for ML (2016) and RG).

Calculation of Catchment-Scale Gas Generation From Grinding Experiments

To compare potential gas generation by glacial erosion for catchments of contrasting size and erosion rates, the rate of gas production per m² of the catchment per day was calculated using...
Equation 1, after Telling et al. (2015):

\[ G_{\text{(catchment)}} = \frac{G \times E}{A \times 365} \]  

where, \( G_{\text{(catchment)}} \) is the catchment-wide rate of gas production (\( H_2, \ CH_4, \) or \( CO_2 \)) generated by mechanical grinding (nmol m\(^{-2}\) d\(^{-1}\)); \( G \) is the amount of gas generated per gram of sample (nmol g\(^{-1}\)); \( E \) is the flux of suspended sediment (SS) exported from the catchment per year (g a\(^{-1}\)); \( A \) is the area of the catchment (m\(^2\)) and 365 converts the rate from yearly to daily. Note that \( G \) is the amount of gas generated per gram of sample when ground to the average grain size of SS measured in glacial runoff from the catchment (Table 3). \( G \) is calculated using the regression line equation of gas production against grain size (Figure 1). Values used for \( E \) and the average grain size of SS in each catchment are shown in Table 3. Where \( G_{\text{(catchment)}} \) was calculated for \( H_2 \) generation by silica-water reactions at LG (section Hydrogen), the value for \( G \) was 11.4 nmol H\(_2\) g\(^{-1}\) as calculated by Telling et al. (2015).

These calculations assume that the measured suspended sediment flux in runoff from each catchment is in equilibrium with the rate of mechanical erosion. This is likely to be true over decadal and longer timescales, as otherwise sediment beneath glaciers would either run out or would accumulate. However, there is the potential for the erosion rate to be out of equilibrium with the suspended sediment flux in individual years, particularly at small, slowly eroding glaciers which are underlain by subglacial till (e.g., RG and ML). Here, in the absence of other data we have made the assumption that the present day rate is equivalent to a longer term abrasion rate measured over a number of years. These calculations also assume that the average suspended sediment grain size has been achieved from grinding an original grain size of between 250 \( \mu \)m and 2 mm. This is unlikely to be true, therefore, \( G_{\text{(catchment)}} \) is likely a lower estimate as the reduction of bedrock to 2 mm sized particles will likely generate additional gas. Where multiple rock types were present in the catchment, we selected the dominant rock type to calculate values of \( G \). The only exception to this was for \( CO_2 \) at LG which was calculated from experiments in which basal sediment was ground, rather than the bedrock (gneiss) which did not produce quantifiable \( CO_2 \). Previous studies have shown that basal ice sediment at LG has an organic carbon content of 0.44 \% (dry weight) (Lawson et al., 2014) which was presumably the origin of the \( CO_2 \) released during grinding of the basal sediment.

Calculation of Supported Rates of Hydrogenotrophic Methanogenesis

To determine the significance of the \( H_2 \) generated by glacial erosion to subglacial microbial processes, the degree of hydrogenotrophic methanogenesis that could be sustained by grinding-generated \( H_2 \) and \( CO_2 \) was determined for two catchments with contrasting geologies; gneissic LG and sedimentary RG. To calculate the depth of sediment throughout which hydrogenotrophic methanogenesis could be sustained by \( H_2 \) generated from grinding, we used Equation 2:

\[ D = \frac{H_2_{\text{(catchment)}}}{CH_4_{\text{(methanogenesis)}} \times 4 \times S} \]  

Where, \( D \) is the depth of sediment (m) throughout which methanogenesis could be supported by grinding-generated \( H_2 \). \( H_2_{\text{(catchment)}} \) is the \( H_2 \) generated by mechanical grinding from Equation 1 (nmol m\(^{-2}\) d\(^{-1}\)), \( CH_4_{\text{(methanogenesis)}} \) is the measured rate of \( CH_4 \) production by methanogenesis at the catchment derived from incubation studies (nmol g\(^{-1}\) d\(^{-1}\)), 4 is the stoichiometric relationship between \( H_2 \) and \( CH_4 \) during methanogenesis (Equation 3), S is the grams of sediment per meter cubed where the wet sediment has a density of 2 \( \times \) 10\(^8\) g m\(^{-3}\) (as measured by Stibal et al., 2012) and therefore converts the rate of methanogenesis from nmol CH\(_4\) g\(^{-1}\) d\(^{-1}\) to nmol CH\(_4\) m\(^{-2}\) d\(^{-1}\). For RG, values used for \( CH_4_{\text{(methanogenesis)}} \) were 0.0002–0.0012 nmol CH\(_4\) g\(^{-1}\) d\(^{-1}\), as measured during incubations at 4°C of basal sediment from RG (Boyd et al., 2010). For LG, 0.00018 nmol CH\(_4\) g\(^{-1}\) d\(^{-1}\) was used for \( CH_4_{\text{(methanogenesis)}} \) (Stibal et al., 2012). This rate was measured during incubation at 1°C of sediment from Russell Glacier, which is adjacent to, and has the same bedrock geology as LG (Stibal et al., 2012).

Since the \( H_2/CO_2 \) pathway for methanogenesis requires a source of \( CO_2 \) in addition to \( H_2 \), we also calculated whether sufficient \( CO_2 \) is generated by grinding to allow microbes to utilize mechanically generated \( H_2 \). This was calculated from the stoichiometric relationship of \( H_2 \) and \( CO_2 \) in the hydrogenotrophic methanogenesis pathway:

\[ 4H_2 + CO_2 \rightarrow CH_4 + 2H_2O \]  

Calculation of Theoretical Total Flux of CO\(_2\) and CH\(_4\) in Subglacial Discharge

For ML and LG, it was necessary to calculate an estimated flux of gas from grinding per year to allow direct comparison with fluxes in the literature. At ML, \( CO_2 \) fluxes can be compared with estimates of \( CO_2 \) “used” in a chemical weathering study at the catchment (Hodson et al., 2000). \( CH_4 \) fluxes can be compared with measured \( CH_4 \) export at LG. The latter has been inferred to be microbial in origin (Lamarche-Gagnon et al., accepted), thus providing a comparison between grinding-generated abiotic \( CH_4 \) and suspected subglacial microbial generation rates. The theoretical flux of \( CH_4 \) and \( CO_2 \) was calculated as shown:

\[ G_{\text{flux}} = G_{\text{(generated)}} \times E \]  

where, \( G_{\text{flux}} \) is the grinding-generated \( CH_4 \) exported from the catchment per year (t); \( G_{\text{(generated)}} \) is the gas generated per gram of sample (t [g sample]\(^{-1}\)); \( E \) is the flux of suspended sediment exported from the catchment per year (g a\(^{-1}\)). Values used for \( E \) are shown in Table 3 for LG. For ML, we used only data for the year closest (2000) to when the comparison study was conducted, where \( E \) was 113 \( \times \) 108 g a\(^{-1}\) and the mean SS grain size used to determine \( G_{\text{(generated)}} \) was 33.1 \( \mu \)m (Hodson et al., 2004). \( G_{\text{(generated)}} \) is calculated the same way as \( G_{\text{(catchment)}} \) in...
Equation 1 but converted to tons of gas per gram of sediment using the molecular weight of the gas.

RESULTS

Variations in rock hardness of the samples analyzed resulted in widely differing grain sizes when ground for the same time. Therefore, we used the average grain size of the ground sample rather than the grinding time to compare the amount of gas produced by grinding different samples. Decreasing average grain size equated to an increase in the amount of \( \text{H}_2 \), \( \text{CO}_2 \), \( \text{CO} \), \( \text{CH}_4 \), \( \text{C}_2\text{H}_6 \), \( \text{C}_3\text{H}_8 \), and \( \text{C}_2\text{H}_4 \) emitted during grinding (Figure 1). The only exception to this pattern was a decrease in the \( \text{CO}_2 \) emitted with decreasing average grain size for LG basal sediment. It is likely that the amount of gas emitted would continue to increase with further reduction in grain size, however, using this method we were not able to further decrease the grain size by grinding for longer than 30 minutes.

The amount of \( \text{H}_2 \) measured during the grinding of rock and sediment samples varied by two orders of magnitude depending on the rock type investigated (Figure 1A). Mica-rich samples, EG schist, ML schist, and ML sediment, produced up to one order of magnitude more \( \text{H}_2 \) than the non-mica-rich samples when ground to 17 \( \mu \)m (the average grain size of suspended sediment...
exported from LG, RG and ML, for which this data was available; Table 3).

CO$_2$ was detected during the grinding of all rock and sediment samples but was below the limit of quantification ($<1.5$ nmol $g^{-1}$) for the MG granite, LG gneiss, and EG schist. However, large amounts of CO$_2$ (µmol g$^{-1}$) were produced during grinding of RG and ML rock samples (Figure 1B). Correlations between the amounts of gases measured were investigated in order to identify potential release mechanisms. For example, the correlation between CO$_2$ and CH$_4$ was highly linear within individual rock samples ($R^2 > 0.8$), but not within the entire population of rock samples ($R^2 0.32$) due to the higher CO$_2$:CH$_4$ ratios of the ML samples than of the RG rocks (Figure 2A).

CO was analyzed during grinding of rocks from catchments with contrasting geologies; RG sedimentary rocks and MG granite. Small amounts of CO (1–4 nmol $g^{-1}$) were measured during grinding of MG granite. The three RG rocks produced similar amounts of CO (2–20 nmol $g^{-1}$) as each other when ground to the same average grain size (Figure 1C). There was a significant linear correlation between CO$_2$ and CO produced during grinding of the RG rocks ($R^2$ of 0.93). The strength of these correlations increased when considering only single rock types ($R^2$ of 0.96 to 0.99) (Figure 2B).

A wide range of hydrocarbon species were produced during grinding, including CH$_4$, C$_2$H$_6$, C$_2$H$_4$, C$_3$H$_2$, and C$_3$H$_8$ (Table 4). CH$_4$ and C$_2$H$_6$ were produced during grinding of all rock and sediment samples, except for LG granite, where C$_2$H$_6$ was not detected. CH$_4$ (Figure 1D) production was notably higher for the sedimentary RG samples than for the crystalline samples of EG, MG, and LG. LG sediment samples produced similar amounts of CH$_4$ at all grain sizes as LG gneiss, whereas ML sediment produced three times more CH$_4$ than the crushed ML schist, one of the dominant rock types in the catchment. This suggests that ML schist is not the dominant composition of the basal till. ML sediment behaved as an outlier in several correlations with elevated C$_2$H$_6$:CH$_4$
We therefore propose that the majority of which involve reactions with surface radicals which form on the fractured surfaces of minerals where bonds have broken. The first generation mechanism results from the reaction of water with silica radicals (Si) and is the most commonly attributed source of H2 released from the grinding of rocks, e.g., in fault zones (e.g., Sugisaki et al., 1983; Ito et al., 1999):

\[
\text{Si} + \text{H}_2\text{O} \rightarrow \text{Si} - \text{OH} + \text{H} \quad (5)
\]

\[
2\text{H} \rightarrow \text{H}_2 \quad (6)
\]

The second potential H2 generation mechanism involves the reaction of hydroxyl functional groups (–OH) from within the crystal structure with Si radicals which form on the surface of a fracture plane (Kameda et al., 2004; Hirose et al., 2011):

\[
\text{Si} + \text{ALOH} \rightarrow \text{Si} - \text{O} - \text{Al} + \text{H} \quad (6)
\]

This mechanism has been used to explain why the grinding of –OH rich minerals, such as phyllosilicates (e.g., muscovite and biotite mica) and the clays, generate more H2 than grinding other silicate minerals to similar surface areas, and therefore, producing similar amounts of silica radicals (Kameda et al., 2003).

It seems unlikely that the first mechanism (Equation 5) was the primary production mechanism in our experiments, given the significant quantities of H2 released under dry conditions. Although it is possible that some H2O, either as liquid or vapor, was released from fluid inclusions in our samples during grinding, we propose that an additional mechanism was required for the high amounts of H2 emitted during grinding of, in particular, EG schist, ML schist, ML sediment, and MG granite (Figure 1A). Therefore, to further investigate if the reaction of –OH groups with Si (Equation 6) better explains the high production of H2 measured, we compared the estimated content of the micas, biotite and muscovite, in hand specimen with the H2 generated when the sample was ground to 17 µm (the average grain size of suspended sediment exported from LG, RG, and ML, for which this data was available; Table 3). Of the samples here, excluding the RG rocks which were too fine-grained for hand-specimen mineral quantification, ML and EG schist had the most muscovite and biotite at an estimated 60 and 70 % area, respectively, (Table 1) and generated the most H2 when ground to 17 µm at 164 and 407 nmol g\(^{-1}\), respectively. Comparatively, ML quartzite, which had no detectable mica by this method, and LG granite, which had approximately 5 % mica, produced the lowest amounts of H2 when ground to 17 µm at 31 and 21 nmol g\(^{-1}\), respectively. The three sedimentary rocks from RG were too fine-grained for mica identification in hand-specimen. However, based upon known characteristics of the rock types (shale and muddy-carbonates), we can broadly infer that they contained high quantities of “mud,” i.e., clay minerals (hydrous aluminum phyllosilicates). Reaction of Si with –OH groups present in clays could supply H2 in addition to that released from pore spaces, thus helping to explain the high amounts of H2 released when grinding the RG rocks (Figure 1A). Our results broadly support that increased mica or clay content and, therefore, increased presence of –OH groups stimulates greater

![Figure 3A](https://example.com/figure3a.png) and H2:CO2 (Figure 3B) ratios compared with the wider population of rocks. The H2:CO2 and C2H6:CH4 relationships were highly correlated for all other rocks (R\(^2\) of 0.86 and 0.93, respectively) when ML sediment was not included.

**DISCUSSION**

**Sources and Generation Mechanisms for Gases During Glacial Grinding**

The potential sources of the different gases measured during grinding of rock and glacial sediment samples in this study likely reflect release from two primary mechanisms. First, gases may be released from pore spaces, fluid inclusions or crystal structures. Second, they may be formed as a consequence of bond breakages during grinding of the rock. The dominant source for each sample will be dependent upon the rock type (i.e., sedimentary vs. crystalline). We use our experimental data to infer the most likely source of each gas for the different rock types in the following sections.

**Hydrogen**

H2 was measured during grinding of all of the samples. There is some evidence that H2, generated by thermal degradation of organic matter during a rock’s formation, can be trapped within the pore spaces of sedimentary rocks (Suzuki et al., 2017). The breaking open of pore spaces during grinding may be an important source of H2 for the sedimentary RG rocks. For the crystalline samples (LG, MG, EG and ML) release of free H2 trapped within the rock is a less likely source because, although H2 can be contained within fluid inclusions, this is rare (Burke, 2001). We therefore propose that the majority of H2 emitted by the grinding of the crystalline samples and at least some of the H2 emitted by the sedimentary samples is likely formed during the grinding process, rather than released. There are two likely formation mechanisms for H2, both of which involve reactions with surface radicals which form on the

**TABLE 4** The hydrocarbons detected during grinding.

<table>
<thead>
<tr>
<th>Glacier</th>
<th>Sample</th>
<th>CH4</th>
<th>C2H6</th>
<th>C2H4</th>
<th>C2H2</th>
<th>C3H8</th>
</tr>
</thead>
<tbody>
<tr>
<td>MG</td>
<td>granite</td>
<td>6.6</td>
<td>&lt;</td>
<td>&lt;</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>LG</td>
<td>gneiss</td>
<td>3.0</td>
<td>&lt;</td>
<td>&lt;</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>EG</td>
<td>schist</td>
<td>3.8</td>
<td>&lt;</td>
<td>&lt;</td>
<td>-</td>
<td>&lt;</td>
</tr>
<tr>
<td>ML</td>
<td>schist</td>
<td>10.4</td>
<td>0.7</td>
<td>n.d.</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>ML</td>
<td>quartzite</td>
<td>16.0</td>
<td>0.5</td>
<td>&lt;</td>
<td>-</td>
<td>&lt;</td>
</tr>
<tr>
<td>RG</td>
<td>hard MC</td>
<td>6.9</td>
<td>0.3</td>
<td>1.1</td>
<td>-</td>
<td>&lt;</td>
</tr>
<tr>
<td>RG</td>
<td>soft MC</td>
<td>222</td>
<td>1.3</td>
<td>0.7</td>
<td>-</td>
<td>&lt;</td>
</tr>
<tr>
<td>RG</td>
<td>shale</td>
<td>196</td>
<td>1.4</td>
<td>0.5</td>
<td>-</td>
<td>&lt;</td>
</tr>
</tbody>
</table>

Where quantifiable, the amount (nmol g\(^{-1}\)) released when ground to an average grain size of 20 µm is shown (calculated from linear regression equation). (-) detected but below limit of quantification. (-) detected but not quantified as no standard available for acetylene (C2H2). MC is muddy carbone with “hard” and “soft” describing the relative “hardness” of the samples to each other based on how quickly the entire starting sample was powdered during grinding.

![Figure 1A](https://example.com/figure1a.png)
H₂ generation during grinding. However, more detailed analysis of mineralogical composition would be required to definitively test this hypothesis.

Several experiments have shown that the loss of –OH groups from a phyllosilicate mineral (e.g., by Equation 6) leads to an amorphous structure (e.g., Takahashi, 1959; Kristof et al., 1993; Martinelli and Plescia, 2005). Interestingly, a recent study by Hawkings et al. (2017) described significant quantities of amorphous material, particularly silica, on suspended sediments in LG's subglacial meltwaters. The authors hypothesized one potential source of this amorphous material as the mechanical grinding of bedrock beneath the ice sheet. The amorphous material on the suspended sediment grains was largely found on the edges of platy minerals (e.g., micas or clays) and contained varying amounts of auxiliary elements, including, aluminum (Hawkings et al., 2017). This tentatively supports the reaction of Si with –OH groups in phyllosilicates (Equation 6) being an important driver of H₂ generation by mechanical erosion.

We propose that the main mechanism for generation of H₂ by grinding of crystalline rocks was by reaction of Si-with –OH due to the presence of phyllosilicate minerals in all rocks, the occurrence of higher estimated mica contents with higher H₂ generation, and the presence of amorphous material containing Al on platy minerals. Even for rocks containing minor quantities of –OH-rich minerals (e.g., LG gneiss and ML quartzite), studies have shown that more than half the H₂ generated from grinding can arise from the fracturing of minor minerals (Kameda et al., 2003). This, in combination with the possible release of H₂O from fluid inclusions allowing some formation by Equation 5, explains the production of H₂ from samples that are poor in phyllosilicates.

A recent study by Telling et al. (2015) measured H₂ generated by rock-water reactions (Equation 5) during incubation of crushed rocks from glaciated catchments with water at subglacially-relevant temperatures (0°C). At LG, data was available to directly compare estimated long-term catchment-scaled rates of H₂ generation by rock-water reactions from the Telling et al. (2015) study with H₂ generation by dry grinding from this study. Generation of H₂ by dry grinding at LG (244 nmol m⁻² d⁻¹; Table 5) was the same order of magnitude as the generation previously inferred by rock-water reactions (151 nmol m⁻² d⁻¹).

**Carbon Dioxide and Carbon Monoxide**

Significant generation of CO₂ was measured during grinding of sedimentary rocks from RG and metamorphic rock and sediment from ML. Possible sources include release from fluid inclusions, release from pore spaces, dissociation from calcite/dolomite minerals and oxidation of the released CH₄. The latter process seems unlikely to be occurring in these experiments because grinding occurred in an oxygen free atmosphere, temperatures were unlikely to be high enough for CH₄ combustion (the mill was never hot to the touch), and 1–4 orders of magnitude more CO₂ was measured than CH₄. Therefore, the remaining possible sources of CO₂ (fluid inclusions, pore spaces, and crystal dissociation) are more likely and vary by rock type. For ML samples, CO₂ was probably released from CO₂-rich fluid inclusions which are common in many minerals and can form during or post-metamorphism (Burke, 2001; Diamond, 2001). For RG shale, the majority of CO₂ (as with the hydrocarbons, section Hydrocarbons) was likely released from pore spaces between the particles of clays and other minerals which were broken open upon grinding (Wang et al., 2015). For the RG muddy carbonates, we speculate that CO₂ was both released from pore spaces and released from the breakdown of calcite (CaCO₃) and/or dolomite (CaMg(CO₃)₂) crystals that make up the carbonate component of the rock. The latter process has been suggested to occur during the mechanical breakdown of marly carbonates, a similar rock type to the RG muddy carbonates, by the following mechanisms (Martinelli and Plescia, 2005):

\[
\text{CaCO}_3 \rightarrow \text{Ca} - \text{O} + \text{CO}_2 \quad (7)
\]

\[
\text{CaMg(CO}_3)_2 \rightarrow \text{CaMgO}_2 + 2\text{CO}_2 \quad (8)
\]

Emission of CO during grinding was analyzed for rocks from two catchments with contrasting geologies; high total carbon sedimentary RG and low total carbon igneous MG (Figure 1C). There is little in the literature about the release or production of CO during the mechanical breakdown of rocks/minerals. CO can be found in fluid inclusions in crystalline rocks (Burke, 2001) making this the likely source of CO from grinding MG granite. The Martinelli and Plescia (2005) study on the mechanical breakdown of marly-carbonaceous rocks (broadly similar in composition to the RG muddy carbonates) notes that CO is produced in “measurable quantities”. However, this was not quantified and no mechanism for release was described. The amount of CO and CO₂ released was highly linearly correlated for the RG rocks ($R^2 > 0.92$) (Figure 2B). We therefore tentatively suggest that the source of CO was similar to CO₂ for RG rocks, i.e., primarily pore spaces for the shale and both pore spaces and release from the breakdown of calcite and dolomite crystals for the muddy carbonates.

**Hydrocarbons**

A range of short chain hydrocarbon species were detected during grinding: CH₄ (methane), C₂H₆ (ethane), C₂H₄ (ethylene), C₂H₂ (acetylene), and C₃H₆ (propane). Most significantly, CH₂ was produced during grinding of all samples, with significantly higher amounts produced by grinding sedimentary rocks from RG than the other samples (Figure 1D). Three possible sources of hydrocarbons are fluid inclusions in the crystalline samples, pore spaces in the sedimentary samples and breakdown of carbon compounds in all samples. Fluid inclusions in igneous and metamorphic rocks commonly contain CH₄ with C₂H₆ as a more minor component and the higher hydrocarbons comparatively rare (Burke, 2001). This follows with the results found here: CH₄ was the most abundant hydrocarbon measured; C₂H₆, where quantifiable, was roughly 1–2 orders of magnitude lower; and lower or trace quantities of C₃H₈, C₂H₂ and C₂H₄ were found, if detected at all (Table 4). The higher CH₄ (1–2 orders of magnitude greater) and C₂H₆ (up to 1 order of magnitude greater) released when grinding the sedimentary RG rocks compared to all others (Figures 1D,E) is likely related to the presence of pore spaces which can trap gaseous hydrocarbons.
These hydrocarbons were likely generated from the breakdown of organic matter during the rock’s formation (Zhang et al., 2014; Wang et al., 2015). The generation of hydrocarbons from the mechanical breakdown of carbon compounds within the rock samples during grinding is a possible source for all samples. To investigate this, the total carbon content (weight percent) of the rock was measured and compared to the amount of hydrocarbons, and other carbon-based gases, measured during grinding. Increasing total carbon content broadly correlated with higher emission of CH<sub>4</sub> (R<sup>2</sup> of 0.72, p < 0.001; Figure 4). There was a weaker correlation with C<sub>2</sub>H<sub>6</sub> (R<sup>2</sup> of 0.46, p < 0.001) perhaps because C<sub>2</sub>H<sub>6</sub> was a comparatively minor component of gas released. CO<sub>2</sub> was poorly correlated (R<sup>2</sup> of 0.28, p < 0.001), possibly because of the different suggested primary generation mechanism (dissociation from the crystal structure of calcite and dolomite; section Carbon dioxide and carbon monoxide). CH<sub>4</sub> and C<sub>2</sub>H<sub>6</sub> were correlated across all samples (Figure 3A) which could indicate that the hydrocarbons trapped within the rocks (in fluid inclusions or pore spaces) originally formed in similar proportions. However, the formation mechanisms are vastly different, i.e., post-magmatic processes influencing fluid inclusion composition in igneous rocks (Potter and Konnerup-Madsen, 2003), such as those from LG and MG, or biogenic and/or thermogenic processes influencing pore space composition during the formation of shale (Wang et al., 2015). It is therefore possible that the linear relationship between CH<sub>4</sub> and C<sub>2</sub>H<sub>6</sub> could be explained by a common generation mechanism during grinding, i.e., from the mechanical breakdown of carbon compounds. Accurately distinguishing between the discussed sources of the measured hydrocarbons (fluid inclusions, pore spaces, and generation by breakdown of carbon compounds) would require in situ detection of gases in fluid inclusions, e.g., by raman microscopy. Interestingly, ML sediment was an outlier to the high collinearity of CH<sub>4</sub> and C<sub>2</sub>H<sub>6</sub> between all samples. If ML sediment was included within the analysis the R<sup>2</sup> of the correlation was reduced from 0.93 to 0.79. A similar pattern was seen for H<sub>2</sub>/CO<sub>2</sub> ratios (Figure 3B) which were elevated for ML sediment. This sample was collected from in front of the glacier terminus and consequently was exposed to the atmosphere. We therefore suggest that the elevated ratios for ML sediment reflected post-depositional processes, such as oxidation of CH<sub>4</sub> and dissolution, causing loss of CO<sub>2</sub> trapped within the minerals.

### TABLE 5 | Comparison of estimated long-term catchment production rates of H<sub>2</sub>, CO<sub>2</sub>, and CH<sub>4</sub> (nmol gas m<sup>−2</sup> day<sup>−1</sup>) by subglacial grinding.

<table>
<thead>
<tr>
<th>Glacier</th>
<th>Catchment area (km&lt;sup&gt;2&lt;/sup&gt;)</th>
<th>Sediment export (t km&lt;sup&gt;−2&lt;/sup&gt; a&lt;sup&gt;−1&lt;/sup&gt;)</th>
<th>H&lt;sub&gt;2&lt;/sub&gt; production (nmol m&lt;sup&gt;−2&lt;/sup&gt; d&lt;sup&gt;−1&lt;/sup&gt;)</th>
<th>CO&lt;sub&gt;2&lt;/sub&gt; production (nmol m&lt;sup&gt;−2&lt;/sup&gt; d&lt;sup&gt;−1&lt;/sup&gt;)</th>
<th>CH&lt;sub&gt;4&lt;/sub&gt; production (nmol m&lt;sup&gt;−2&lt;/sup&gt; d&lt;sup&gt;−1&lt;/sup&gt;)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MG</td>
<td>14.4</td>
<td>1,219</td>
<td>155</td>
<td>&lt;</td>
<td>21</td>
</tr>
<tr>
<td>LG</td>
<td>600</td>
<td>4,833</td>
<td>244</td>
<td>479</td>
<td>39</td>
</tr>
<tr>
<td>EG</td>
<td>36.2</td>
<td>456</td>
<td>311</td>
<td>&lt;</td>
<td>12</td>
</tr>
<tr>
<td>ML</td>
<td>5.4</td>
<td>1,751</td>
<td>369</td>
<td>26,965</td>
<td>25</td>
</tr>
<tr>
<td>RG</td>
<td>1.4</td>
<td>237</td>
<td>37</td>
<td>13,519</td>
<td>650</td>
</tr>
</tbody>
</table>

Rates were calculated here (Calculation of catchment-scale gas generation from grinding experiments). Sediment export rates were adapted from Table 3. (<) concentrations too low to calculate catchment rates.

![FIGURE 4](Image 314x350 to 541x555) The amount of CO<sub>2</sub>, CH<sub>4</sub>, and C<sub>2</sub>H<sub>6</sub> produced (nmol g<sup>−1</sup>) compared to the percentage weight of total carbon present in the sample.

### Glacial Erosion Liberates Potential Energy Sources for Subglacial Microbes

The generation of gases by mechanical grinding beneath glaciers and ice sheets could be significant for various microbiological processes in subglacial environments. A wide variety of microbes use different gases as their energy and/or carbon source (Conrad, 1996). Of the gases measured after grinding in these experiments, H<sub>2</sub>, CO<sub>2</sub>, CH<sub>4</sub>, and CO are the most relevant to microbial processes. In the subglacial environment, an abiogenic source of these gases from bedrock grinding has the potential to sustain microbial populations during long periods of isolation from the atmosphere and surface processes or where other sources are scarce, e.g., where subglacial sediments are carbon-poor. Notably, H<sub>2</sub> is utilized by many methanogens as a growth substrate, with CO<sub>2</sub> as the preferred electron acceptor (Equation 3). These hydrogenotrophic methanogens are an important species in isolated environments due to their ability to subsist upon abiogenic H<sub>2</sub> and CO<sub>2</sub> as their sole energy source (Sleep and Zoback, 2007; Hirose et al., 2011). Several
studies have identified the presence of methanogens, including hydrogenotrophic methanogens, in subglacial sediments and waters, and long-term incubations of subglacial sediments have shown active CH₄ generation (Boyd et al., 2010; Stibal et al., 2012).

In order for grinding-generated H₂ to be a source of energy for subglacial microbes, generation rates would need to be comparable with microbial rates of consumption. Therefore, catchment-scaled rates of H₂ generation by subglacial rock grinding were calculated. These rates were calculated using suspended sediment (SS) export as a proxy for decadal and longer term abrasion rates. Varying rates were obtained depending upon bedrock type and erosion rate. For example, rates of H₂ generation at ML and EG were the same order of magnitude as for LG (Table 5) despite the order of magnitude more H₂ released per gram of the former samples (Figure 1A). This is because LG has a considerably higher sediment export and erosion rate. Similarly, grinding of porphyritic granite from MG produced more H₂ per gram of sample than LG gneiss (Figure 1A), but MG had a lower production rate due to the slower erosion rate at this much smaller glacier (Table 5). Therefore, although the magnitude of H₂ generation per gram is dependent on the bedrock type, the erosion rate is an important influence upon the significance of H₂ generation in a catchment.

To determine if rates of grinding-generated H₂ are sufficient to act as an energy source for subglacial methanogens, we calculated the depth of sediment throughout which measured rates of methanogenesis could be supported, assuming the sediments are anoxic. Rates of methanogenesis were available for two lithologically contrasting catchments: gneissic LG and sedimentary RG. At LG, we calculate that hydrogenotrophic methanogenesis could be supported throughout 17 cm of sediment depth beneath the entire LG catchment. It is important to note that the grinding of basal sediment at LG produces similar quantities of H₂ per gram as grinding bedrock (Figure 1A), indicating that reworking of basal sediment would sustain similar rates of methanogenesis as erosion of bedrock itself. The sediment depth throughout which methanogenesis could be supported by grinding-generated H₂ at RG was much lower (2 cm) due to the significantly lower rate of erosion at RG (~0.1 mm yr⁻¹) than at LG (4.8 mm yr⁻¹; Cowton et al., 2012). The sediment depths calculated here are within estimated sediment depths present at each catchment. At RG estimated depths are on the decimetre or sub-decimetre scale based upon observations that RG sits partly upon till and partly upon bare bedrock (Boyd et al., 2010). At LG, subglacial sediments could be present on the decimetre scale up to the multi-meter scale as indicated by 0 to decimetre thick sediments found at nearby Isunnguata Sermia and spatially-limited seismic evidence of sedimentary basins 10s of meters deep at LG (Dow et al., 2013; Harper et al., 2017; Kulessa et al., 2017).

Telling et al. (2015) calculated that water-Si reactions (Equation 5) generated enough H₂ to support methanogenesis in the top 1 cm of sediment throughout the LG catchment. However, the authors calculated that the rate of CO₂ released from bubbles trapped in the ice (10% of ice volume, CO₂ pre-industrial fraction of 0.0003) by basal melting (assumed rate of 6 mm/yr) was insufficient to act as the sole acceptor for their calculated rates of rock-water H₂ generation. Here, we show that grinding basal sediment at LG generates 8 times the CO₂ required as an electron donor for utilizing grinding-generated-H₂ as an abiotic energy source in hydrogenotrophic methanogenesis (Table 5; section Calculation of supported rates of hydrogenotrophic methanogenesis). At RG, grinding CO₂-rich carbonates and shale generates more than 90 times the CO₂ required for methanogens utilizing grinding-generated-H₂ for the same process. Therefore, hydrogenotrophic methanogenesis could be sustained entirely by the process of glacial grinding throughout LG and RG catchments at sediment depths of 17 cm and 2 cm, respectively. RG is a small glacier with a low erosion rate (~0.1 mm a⁻¹). Glaciers with faster erosion rates and similar bedrock to RG would generate more gases per m⁻² d⁻¹ by grinding and, therefore, could sustain hydrogenotrophic methanogenesis throughout greater depths of sediment, assuming a constant microbial metabolic rate. Further, the orthogneiss sample from LG is broadly representative of much of the basement rock that underlies the Greenland Ice Sheet. We therefore suggest that mechanical glacial erosion could provide an abiotic source of energy, in the form of H₂ and CO₂, for subglacial microbial metabolism beneath the Greenland Ice Sheet and especially in marginal areas with higher ice flow and bedrock erosion rates.

Like H₂ and CO₂, CO was another gas emitted in our grinding experiments that may have similar potential to provide carbon and energy to subglacial microbes where other sources are limited. Several groups of bacteria and archaea can utilize CO as a source of carbon and/or energy. However, CO-utilizing microbes have been little explored in the subglacial or sub-ice-sheet literature to date. Groups of microbes that can utilize CO include anaerobic sulfate reducers, hydrogenogens, methanogens and acetogens (Oelgeschläger and Rother, 2008). Given the wide range of anaerobic metabolisms that have been reported in subglacial systems (e.g. see Skidmore, 2011 for review) it is plausible that CO-metabolizing species are present in subglacial environments. CO was measured after grinding rocks from two glaciated catchments with contrasting geologies: sedimentary RG, and igneous MG. The estimated long-term catchment-scaled rates of CO production at RG and MG were similar at 6 and 8 nmol m⁻² d⁻¹, respectively. At both catchments, CO generation rates are an order of magnitude lower than H₂ generation rates (Table 5). However, it is still plausible that grinding-generated CO could be metabolized beneath glaciers. Further work would be needed to establish the production of CO by subglacial erosion at additional catchments and to establish the existence and utilization of CO by subglacial microbes to determine if this finding is significant.

Glacial Erosion Liberates a Source of Acidity for Chemical Weathering

The generation of CO₂ by subglacial erosion has the potential to influence glacial chemical weathering rates because CO₂ dissolves in water to form carbonic acid which dissociates to generate
protons. This drives the chemical dissolution of carbonate (9) and silicate (10) minerals.

\[
\begin{align*}
\text{CO}_2\text{(aq)} + \text{H}_2\text{O}\text{(l)} + \text{CaCO}_3\text{(s)} & \rightarrow \text{Ca}^{2+}\text{(aq)} + 2\text{HCO}_3^-\text{(aq)} \\
2\text{CO}_2\text{(aq)} + 3\text{H}_2\text{O}\text{(l)} + \text{CaAl}_2\text{Si}_2\text{O}_8\text{(s)} & \rightarrow \text{Al}_2\text{Si}_2\text{O}_5\text{(OH)}_4\text{(s)} + \text{Ca}^{2+}\text{(aq)} + 2\text{HCO}_3^-\text{(aq)} \\
\end{align*}
\]

Proton sources in the subglacial environment can become depleted due to isolation during glaciation, and the supply of protons can limit chemical weathering rates (Tranter et al., 1996). CO$_2$ is supplied to the subglacial environment by: periodic inputs of supraglacial meltwaters carrying atmospheric gases (isolated to small areas of the ice base where surface waters reach these depths); the release of palaeo-CO$_2$ from gas bubbles in the ice during basal melting (which occurs at a rate determined by basal melt rates which varies across the bed); and from microbial respiration (Sharp et al., 1999; Montross et al., 2013). The production of CO$_2$ from glacial crushing is therefore an additional source that can provide acidity to drive chemical weathering.

At ML, we calculated gas generation rates using suspended sediment fluxes as an estimate for the long term abrasion rate. Assuming that the present day rates match the longer term abrasion rate, we calculate that 18–21% of CO$_2$ required to sustain chemical weathering rates measured by Hodson et al. (2000) could be generated by grinding of the bedrock and sediment. Although this may be an overestimate if there are significant rates of sediment redistribution relative to abrasion beneath the glacier, the calculation shows the clear potential for CO$_2$ released by grinding to have a significant impact upon CO$_2$ weathering budgets beneath glaciers. The data was not available to calculate weathering rates at RG but given the similarly high magnitudes of CO$_2$ production rates to ML, it is likely that grinding-generated CO$_2$ would be a potential influence on weathering rates at that catchment as well. The trace CO$_2$ generated by grinding igneous and some metamorphic rocks at LG, MG and EG is unlikely to significantly influence dissolution rates in comparison to other sources. However, for glaciers located on sedimentary and some metamorphic bedrock types, which can be found in the Rockies, the Alps, the Himalayas and Antarctica, grinding-generated CO$_2$ has the potential to decrease the amount of CO$_2$ that has previously been interpreted to have been drawn down from the atmosphere by glacial weathering (Hodson et al., 2000).

Glacial Erosion as an Abiogenic Source of Methane

It has been theorized that microbial breakdown of organic matter in overridden sediments has the potential to generate CH$_4$ clathrate beneath ice sheets (Wadhams et al., 2008, 2012). Evidence of subglacial CH$_4$ oxidation has also been demonstrated indicating that methanotrophy can potentially remove large quantities of CH$_4$ in areas of the subglacial environment that areoxic, such as subglacial lakes and upper sediment layers (Dierer et al., 2014; Michaud et al., 2017). CH$_4$ has recently been reported in waters emerging from the Greenland Ice Sheet with some indication that net production is occurring in large areas beneath ice sheets (Dierer et al., 2014; Lamarche-Gagnon et al., accepted). These studies have inferred the source of this CH$_4$ to be microbial. CH$_4$ released by the mechanical subglacial erosion of bedrock has not previously been considered as a potential source. This abiogenic source could be significant in some catchments depending upon the catchment’s bedrock type and erosion rate.

To investigate the significance of grinding as a source of CH$_4$, we compared estimated generation rates at two geologically contrasting catchments; sedimentary RG and gneissic LG. Grinding at RG generated an order of magnitude more CH$_4$ than at LG despite the considerably lower erosion rate (Table 5), likely due to the order of magnitude higher CH$_4$ production per gram of sample (Figure 1D) from RG’s comparatively carbon-rich bedrock. At LG, subglacial CH$_4$ production has recently been estimated to be at least 6 tons per year, for the 600 km$^2$ catchment (Lamarche-Gagnon et al., accepted). This flux was inferred to be from net microbial production of CH$_4$ based on stable isotope analyses of the CH$_4$ exported in subglacial meltwaters. We used catchment rates of CH$_4$ generation by grinding to estimate abiogenic CH$_4$ exported from the LG catchment. Our estimated theoretical CH$_4$ flux from subglacial grinding amounted to 0.14 t y$^{-1}$, an order of magnitude lower than the measured flux of 6 t y$^{-1}$ at LG (Lamarche-Gagnon et al., accepted). Thus, rock grinding is less significant than microbial generation as a source of CH$_4$ in catchments with carbon-poor bedrock.

There are no direct measurements of CH$_4$ in subglacial meltwaters at RG. The estimated grinding-generated abiogenic CH$_4$ export at RG was calculated to be 0.005 t y$^{-1}$, an order of magnitude lower than at LG. However, if the size of the glaciers is taken into account, the export of grinding-generated CH$_4$ per km$^2$ per year is an order of magnitude higher at RG (0.004 t km$^{-2}$ y$^{-1}$) than at LG (0.0002 t km$^{-2}$ y$^{-1}$). Methanogenesis rates measured in incubated subglacial sediment from both catchments are similar at 0.0002–0.0012 nmol CH$_4$ g$^{-1}$ d$^{-1}$ at RG (Boyd et al., 2010) and 0.00018 nmol CH$_4$ g$^{-1}$ d$^{-1}$ at LG (Stibal et al., 2012). The similar methanogenesis rates combined with the order of magnitude more CH$_4$ generated by grinding at RG than LG, both per gram of rock and per km$^2$, could indicate that grinding would release more CH$_4$ than methanogenesis at RG. Therefore, abiogenic grinding-generated CH$_4$ release could be a more significant source of CH$_4$ than methanogenesis in catchments with carbon-rich bedrock.

Our results show that abiogenic CH$_4$ formation from the subglacial grinding of bedrock likely represents a previously unaccounted CH$_4$ source in subglacial catchments. Grinding of bedrock could even constitute the bulk of CH$_4$ released from subglacial systems in regions with carbon-rich bedrock. As such, grinding-generated CH$_4$ should be considered when analyzing CH$_4$ export from regions experiencing high erosion rates or with bedrock with a high carbon content.
CONCLUSION

We demonstrate that mechanical erosion of rocks and minerals beneath glaciers generates a previously unappreciated source of gases relevant to subglacial microbial processes and chemical weathering. We present the first experimental evidence that the grinding of rock and sediment samples from glaciated catchments with widely varying geologies have the potential to generate significant quantities of CO\(_2\), CO, CH\(_4\), other short chain hydrocarbons and, consistent with previous work, H\(_2\) (Telling et al., 2015). The significance of grinding-generated gases for microbial and chemical dissolution processes depends on three primary factors. First, the amount of gas emitted during the mechanical breakdown of a rock will depend upon its mineral composition, the presence of pore spaces in the rock and the presence of fluid inclusions and their composition. Second, the erosion rate in the glaciated catchment will determine how rapidly rocks, sediment and minerals are mechanically broken down and thus, the rate at which gas is emitted. Third, the significance of gas generation via erosion will be determined in part by the comparative rates of supply from other sources in the catchment. Our experiments have shown that glacial erosion has the potential to release CO\(_2\) in sedimentary and some metamorphic catchments at rates that can influence the magnitude of the atmospheric CO\(_2\) budget required to drive weathering. Beneath glaciers of all bedrock types, we would expect to see generation of H\(_2\) and CH\(_4\) by glacial erosion. For catchments where CO\(_2\) is generated in addition to H\(_2\) such as catchments with sedimentary bedrock or basement bedrock as found in Greenland, some portion of hydrogenotrophic methanogenesis could be supported by grinding-generated gases beneath the ice. CH\(_4\) generation from glacial erosion was common to all bedrock types and was shown to be particularly important in sedimentary catchments where rates of grinding-generation could be greater than in situ microbial production. Future studies should determine whether the gases produced in this study are directly utilized by microbial processes or weathering reactions through laboratory incubations with grinding-generated gases. Our findings show that mechanically-generated H\(_2\), CO\(_2\), CO, and CH\(_4\) should be considered in future studies which investigate microbial or geochemical subglacial processes, particularly energy sources for microbes, CH\(_4\) cycling beneath ice, and chemical weathering processes.

DATA AVAILABILITY

Raw data supporting the conclusions of this manuscript are available at doi: 10.6084/m9.figshare.7350041.

AUTHOR CONTRIBUTIONS

JT conceived the study; MM, JT, and MS collected samples and MS supplied data; MM conducted the experiments and analyzed the data. JT and JW assisted with data analysis. MM wrote the first draft of the manuscript with contributions from JW. All authors contributed to the final manuscript.

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