Novel Biogenic Aggregation of Moss Gemmae on a Disappearing African Glacier

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Abstract

Tropical regions are not well represented in glacier biology, yet many tropical glaciers are under threat of disappearance due to climate change. Here we report a novel biogenic aggregation at the terminus of a glacier in the Rwenzori Mountains, Uganda. The material was formed by uniseriate protonemal moss gemmae and protonema. Molecular analysis of five genetic markers determined the taxon as Ceratodon purpureus, a cosmopolitan species that is widespread in tropical and polar regions. Given optimal growing temperatures of isolate is 20–30°C, the cold glacier surface might seem unsuitable for this species. However, the cluster of protonema growth reached approximately 10°C in daytime, suggesting that diurnal increase in temperature may contribute to the moss’s ability to inhabit the glacier surface. The aggregation is also a habitat for microorganisms, and the disappearance of this glacier will lead to the loss of this unique ecosystem.

Introduction

Many psychrophilic and psychrotolerant microorganisms inhabit supraglacial environments, which have recently been recognized as an important biome [1]. Cryonite granules, dark spherical aggregates typically 1 mm in diameter have been frequently observed on ablation zones of glaciers in many parts of the world [2]. These cryonites consist of mineral particles, organic matter, and microorganisms, which are mainly formed by the aggregation of filamentous cyanobacteria [3]. These cryonite harbor a diverse range of microorganisms, and studies of their molecular diversity have revealed microbial communities of bacteria [4] and archaea [5,6], as well as algae, fungi, amoebas, and invertebrates such as tardigrades [5]. These microbial communities play an important role in the carbon and nitrogen cycles on the glacier [1,7].

Other types of biological aggregations have been reported from supraglacial ecosystems in Iceland and Alaska, namely, globular moss aggregations known as ‘glacier mice’ [8,9] or ‘moss polster’ [10]. These are lenticular moss cushions (0.02 to 0.1 m in diameter) and are composed of a moss envelope covering an internal mass formed from glacial sediment and airborne particles [11]. These moss cushions are expected to impact the ecology and nutrient cycle of the supraglacial ecosystem [11], and also provide a favorable habitat for a variety of invertebrates, including Collembola, Tardigrades, and Nematoda [12].

Previous biological studies have frequently examined mid-latitude and polar glaciers, however, the tropical glaciers are have been studied rarely, except for New Guinea [13]. In equatorial Africa, glaciers persist in three major mountain regions (Mt. Kilimanjaro in Tanzania, Mt. Kenya in Kenya, and the Rwenzori Mountains in Uganda), which have not been previously been targeted in surveys of glacier biology. The Rwenzori glaciers are shrinking rapidly and are expected to disappear by 2020 due to climatic warming [14] and/or lowered humidity and lowered cloudiness [15], as measured by aerial photography and satellite imagery [14].

During a biological field survey on a glacier near the summit of Mt. Stanley, the highest peak in Uganda and in the Rwenzoris, we found a large, black bioaggregation (average long and short axes: 10.1 mm and 12.7 mm) in the supraglacial environment, greater than cryonites. Examination revealed that these granular structures were formed by filamentous moss gemmae and protonema, not cyanobacteria. This is the first report to describe this habitat for such a structure, which we classified as a “glacial moss gemmae aggregation” (GMGA). In order to identify the material we measured the structure (size and mass) and isolated the dominant moss species using both culture and molecular techniques. Furthermore, we examined the photosynthetic activity of isolates under various temperature and radiation conditions.

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Elena glacier was estimated to cover 6.5 km² [16] and by 2003 it had decreased to approximately 1 km² [14]. During this period, glaciers on Mts. Emin, Gessi, and Luigi disappeared completely, leaving only glaciers remaining on three major peaks: Ms. Speke, Baker, and Stanley. In this study, we surveyed Stanley Plateau, the largest glacier on Mt. Rwenzori. Stanley Plateau is a flat sloped glacier that flows from Mt. Stanley’s Alexandra Peak, and is around 1 km long and 0.1–0.3 km wide (Fig. 2a).

In February 2012 and 2013, we collected surface ice samples, including biological debris, at three sites: ST1 (N00°22′31.3″, E29°52′40.2″), ST2 (N00°22′34.7″, E29°52′37.2″) and ST3 (N00°22′52.3″, E29°52′24.6″). At each site, 5 samples were collected from different areas (samples size not measured) and placed in 8 ml plastic bottles for DNA analysis. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for cell counts, and 5 samples from different areas until transport to Kasese, Uganda, the closest city to the Rwenzori Mountains National Park. There, samples for molecular analysis were kept frozen around −20 °C and samples for isolation were kept cold around 0 °C and samples for molecular isolation. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for DNA analysis. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for DNA analysis. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for DNA analysis. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for DNA analysis. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for DNA analysis. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for DNA analysis. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for DNA analysis. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for DNA analysis. In the glacier foreland, located about 10 m from ST1, we collected shoots of bryophytes on dried GMGAs and placed them in 8 ml plastic bottles for DNA analysis.

Samples for cell counts were fixed with 3% formaldehyde and stored at room temperature. All other samples were kept cold around 0 °C in large stainless steel vacuum flasks with glacial ice samples until transport to Kasese, Uganda, the closest city to the Rwenzori Mountains National Park. There, samples for molecular analysis were kept frozen around −20 °C and samples for isolation were kept cold around 0 °C, until they could be transported to the lab for analysis at the National Institute of Polar Research (Tokyo, Japan).

In the field, the internal temperature of the GMGAs at ST1 was measured using a waterproof temperature logger (R-52i; T&D, Matsumoto, Japan). Probes were inserted into the center of two GMGAs and monitored by camera (Optio WG-2; Ricoh, Tokyo, Japan) at intervals to ensure that the measuring apparatus was not disturbed.

Microscopic observation of biological materials on ice surface

The samples were cold-preserved prior to isolation and identification of species. After formaldehyde fixation, 0.1–0.4 ml of 12–60 fold diluted samples were filtered through a hydrophilic polytetrafluoroethylene membrane (Omnipore JGWP1300; Merck Millipore, Billerica, USA) with diameter 13 mm and pore size 0.2 μm. We observed and counted cell concentrations from one-quarter of the membrane, using a fluorescent microscope (IX71 and 81; Olympus, Tokyo, Japan).

18S rRNA gene molecular cloning

DNA of approximately 0.5 g was extracted from samples ST1, ST2, and ST3 using the Fast DNA SPIN Kit for soil (MP Biomedicals, Santa Ana, USA) according to the manufacturer’s instructions. Extracted DNA was diluted to 1.62 ng/μl with water (Ambion Nuclease-Free Water; Life Technologies, Carlsbad, USA). Five aliquots from each site were combined for DNA amplification. Thermal cycling was performed with an initial denaturation step at 98 °C for 3 min, followed by 25 cycles of denaturation at 98 °C for 10 s, annealing at 55 °C for 30 s, and elongation at 72 °C for 1.5 min, using Ex Taq HS DNA polymerase (Takara, Shiga, Japan) and the primer pair of Euk A (5′-ACCTGTGTTGACCTCTGCAGAT-3′) and Euk B (5′-GATCCTCTGAGGTTACCTACCTAGC-3′). Cycling was completed by a final elongation step at 72 °C for 3 min. The PCR-amplified DNA fragments were cloned into the pCR4 vector of the TOPO TA cloning kit (Invitrogen, Carlsbad, USA). Clones obtained from the libraries were sequenced using the 3130xl Genetic Analyzer (Life technologies, Carlsbad, USA) at the National Institute of Polar Research. All sequences were assembled using CodonCode Aligner (CodonCode Corporation, Centerville, USA) and assembled full-length sequences of 18S rRNA were aligned with the eukaryotic Silva database [17] using mothur ver. 1.27.0 [18]. Tentative chimeric sequences were removed using both the reference and de novo modes of Uchime [19] implemented in mothur software package. All good-quality sequences with more than 97% similarity were clustered into operational taxonomic units (OTU).

Isolation of moss and molecular identification

Fragments of cold-preserved GMGA samples were inoculated in liquid Bold’s basal medium (BBM) [20] in a laminar flow bench and incubated at 4 °C for 1 month. Protonemata that grew directly from observed gemmae (Fig. 2 c,d) were transplanted to fresh BBM liquid medium and a 1-month-incubation was repeated. Isolated protonemata were kept in BBM and 1.5% agar medium before extraction and analysis. DNA of a single cluster of protonema in liquid medium was extracted with the Fast DNA SPIN Kit for soil, and 4 different regions (18S rRNA; chloroplast genes, trnL, rps4 and atpB-rbcL intergenic spacer; and mitochondria gene, nad5) were amplified by using Ex Taq HS DNA polymerase (Takara, Shiga, Japan). Thermal cycling for 18S rRNA was carried out following Remias et al. [21], with 35 cycles of denaturation at 98 °C for 10 s, annealing at 54 °C for 30 s, and elongation at 72 °C for 1 min 45 s using the primer pair NS1 (5′-GTAGCTATATGCTTGCTCTC-3′) and 18LS5′-CACCCTACGGAACCTTGTACGCAT-3′). For chloroplast trnL, thermal cycling was performed with 35 cycles of denaturation at 98 °C for 10 s, annealing at 60 °C for 30 s, and elongation at 72 °C for 1 min 45 s and primer pair trnC (5′-CGAAATCGGTAAGCCGTTACG-3′) and trnF (5′-ATTTGAACGTTGACCGGAC-3′), following Taberlet et al. [22]. For chloroplast rps4, thermal cycling was performed according to Nadot et al. [23] and Souza-Chies et al. [24], with 35 cycles of denaturation at 98 °C for 10 s, annealing at 55 °C for 30 s, and elongation at 72 °C for 1 min 43 s using primer pair rps5 (5′-ATGCTCCGGATTACGAGGACGT-3′) and rtm (5′-TACCGAGGTTTTCGAATGC-3′). For chloroplast atpB-rbcL intergenic spacer, thermal cycling was performed according to Chiang et al. [25], with 35 cycles of denaturation at 98 °C for 10 s, annealing at 49 °C for 30 s, and elongation at 72 °C for 1 min 30 s using primer pair atpB-1 (5′-ACATCKKAT-3′) and rbcL-1 (5′-ACACCGAGTTCCTAATTCC-3′), following Lia et al. [26]. Lastly, for mitochondria nad5, thermal cycling was performed according to Shaw et al. [27], with 35 cycles of denaturation at 98 °C for 10 s, annealing at 52 °C for 30 s, and elongation at 72 °C for 1 min 45 s using primer set nad5F4 (5′-GAAGGAGTAGGTCGTGCTCT-3′) and nad5R3 (5′-AAACCGCTGGTGTTACCAT-3′), following Lia et al. [26]. Some of shoots of Bryophyta on dried GMGAs were picked up by tweezers and DNA was analyzed by same method as isolated protonema.
Photosynthetic rate of GMGA and isolate

Photosynthetic rate at 7 different incubation temperatures (5, 10, 15, 20, 25, 30, and 40°C) was measured using a pulse amplitude modulation (PAM) fluorometer (Water-PAM, Waltz, Effeltrich, Germany) with Win-control software for control and analysis following Tanabe et al. [27]. PAM fluorometer is useful to measure the electron transport rate (ETR) of isolate under different incubation factors [27]. For incubation temperatures of 5, 10, and 15°C, photosynthetically active radiation (PAR) intensities were 3, 64, 94, 144, 215, 305, 422, 687, and 1000 mmol photons/m²/s. After these measurements, we had to change to another PAM device, because this device was obviously unstable only under 40°C due to mechanical trouble. Then, we used another device and measured again from 20°C. Results of 20, 25, 30°C are almost the same as previous analysis, and measurement was stable at 40°C in next time. For incubation temperatures of 20, 25, 30, and 40°C, PAR intensities were 8, 62, 92, 140, 209, 297, 412, 674, and 986 mmol photons/m²/s. After a 30 s exposure, a saturating pulse of > 2000 mmol photons/m²/s was applied for 0.4 s at 5°C in a temperature-controlled incubator. The gain value of the photomultiplier (PM-Gain) was set to 3 for all measurements. After incubation of each sample at 5°C for 60 min in dark conditions, a tissue sample of GMGA and isolated protonemata were transferred to the quartz cuvette of the fluorometer. After measurement, incubation temperature was raised by 5°C to 10°C and incubated for 1 h, after which the temperature was raised by 5°C again and incubated for 1 h repeatedly until incubation temperature was 40°C. Light curves were obtained by running a rapid light curve protocol in Win-control software. The photosynthetic rate expressed as relative electron transport rate $rETR$ [28] was as follows:
Here, F and Fm’ are the transient and maximum fluorescence levels at certain actinic light intensities at a given time and (Fm’–F)/Fm’ indicates Photosystem II (PSII) yield. Non-photochemical quenching (NPQ) was calculated by the following equation:

\[ NPQ = \frac{(Fm - Fm')}{Fm'} \times PAR, \]

where Fm is the maximum fluorescence level of non-illuminated samples.

Ethics Statement

Uganda Wildlife Authority and Uganda National Council for Science and Technology authorized all field researchs in Rwenzori Mountains National Park.

Results

Morphological features of GMGA

We found ellipsoidal blackish bioaggregations covering the glacier surface at ST1 (Fig. 2b). We collected and sampled 96 of these bioaggregations, as well as measured their long and short axes, thickness, and mass (Fig. 2c,d). The average long axis, short axis, thickness, and mass were 18.7 mm, 12.7 mm, 8.3 mm, and 1.6 g, respectively (Fig. 3), and these aggregations were clearly larger than cryoconites (average diameter: 1.1 mm; 3). Short axis length was well correlated with long axis length (\( R^2 = 0.705 \)), but not as well correlated with thickness and mass (\( R^2 = 0.543 \) and 0.616, respectively). This means that the structure of this bioaggregation is not spherical but is instead flattened. The bioaggregation was composed of many gemmae and protonema. The gemmae were germinating and developing filamentous protonema, and gemmae were formed repeatedly on protonema. Many moss gemmae were observed especially on the surface of the bioaggregation (Fig. 4a,b), from the top 1–2 mm of the cross section (Fig. 2d). The main framework of these structures was formed by moss gemmae, so we named this structure as “glacial moss gemmae aggregation (GMGA)”.

Figure 2. Research site and glacier moss gemmae aggregation (GMGA). a) Stanley Plateau glacier and Margarita Peak from Mt. Baker, b) Glacier ice surface covered by GMGAs, c) GMGAs (grid cells beneath GMGA are 1×1 mm), d) Cross section of GMGA (scale bar: 2 mm).

![Figure 2](https://example.com/figure2)

Figure 3. Size (long and short axes, thickness) and mass distribution of GMGAs.

![Figure 3](https://example.com/figure3)
The gemmae are filamentous, composed of 1–2 rows of 2–20 cells with slightly thickened brownish cell walls, 100–200 μm long in maximum. These morphological characteristics were well agreed with rhizoidal gemmae of cosmopolitan moss, *Ceratodon purpureus* (Hedw.) Brid. described by Imura and Kanda (1986) based on Antarctic specimens [29].

**Molecular identification of moss species**

We obtained a total of 81 clones from the GMGAs by 18S rRNA gene PCR-cloning. Sixty-three clones (77.8%) were clustered into the same OTU (AB858433: Table 1). The remaining 18 clones were of cercozoa, green algae, and fungi. The 18S rRNA, *rps4*, *trnL*, *atpB-rbcL* intergenic spacer and *nad5* gene sequence of the isolated protonemata (Fig. 4c,d) that grew from the observed gemmae and shoots on dried GMGA were summarized in Table 1. These high-percentage matches (more than 99.9% similarity) from five different regions of the protonemata show that the isolated moss is indeed *C. purpureus*. Also results from four different regions of shoots on dried GMGA show that this specie belonging to genus: *Bryum*, however we could not identify species level from these regions.

**Optimum temperature and PAR of GMGA and isolated protonemata of C. purpureus**

Internal (center) temperature changes was measured during the 2013 field season of two *in situ* GMGAs in ST1 and one dried GMGA found on a rock in glacier foreland (Fig. 5). Temperature change data show clear diurnal cycles with daily exposure to below-freezing temperatures daily. Maximum temperatures reached 8–10°C for *in situ* GMGAs and above 20°C for dried GMGA. The daytime increase in temperature was due to absorption of thermal radiation, but was variable due to decreases in radiation from frequent cloud cover and cooling by glacier ice.

A photosynthetic light curve was measured using a PAM fluorometer under different temperatures of GMGA and the two isolates (Fig. 6). The ETR of GMGA and isolates is high between 20–30°C, and highest at 25°C for GMGA. Electron transport was detected in all three samples even at temperatures as low as 5°C, but ETR was zero or extremely low at 40°C. These results indicate that the optimum temperatures of GMGA and isolated *C. purpureus* are around 25°C. The ETR of GMGA and the isolates was high at low PAR levels (305 and 422 μmol/m²/s) at 5–15°C; however, ETR was high at a medium PAR level (687 μmol/m²/s) at 20–30°C (Fig. 5). Therefore, the optimal PAR value for *in situ* temperatures (0–10°C) is likely between 305–422 μmol/m²/s. The highest ETR value in GMGAs was 674 μmol/m²/s, which was approximately twice that of the other two isolates (Fig. 6).

**Distribution of GMGAs on the glacier**

GMGAs of *Ceratodon purpureus* were observed only at site ST1, the glacier terminus (Fig. 7). The organic carbon mass (62.72±19.39 g/m²) was highest at the terminus. This record is higher than current highest glacial organic carbon mass (38.5±12.4 g/m²) from Qiyi Glacier, China [30]. Moreover, organic carbon mass at our sites without observed GMGAs (ST2:20.73±8.91 g/m², ST3:23.55±6.89 g/m²) were roughly equal to the average high organic carbon mass at Qiyi Glacier (mean: 23.4±16.5 g/m²) [26].

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**Figure 4. Cells of moss gemmae and protonemata.** a,b) Moss protonemal cells formed the main frame of the GMGA (scale bar: 100 μm), c) Moss protonema grew from gemmae below 4°C (scale bar: 100 μm), d) protonemal cells for molecular identification after incubation below 4°C for 1 month in liquid Bold’s basal medium.
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| Gene type          | Genetic region | Sample type       | Accession number | Length (bp) | Closest relative specie | Accession number | Identity (%) | Sequence match (bp) |
|-------------------|----------------|-------------------|------------------|-------------|------------------------|------------------|--------------|---------------------|
| Ribosomal RNA gene | 18S rRNA       | 1: GMGA_cloning   | AB858433         | 1819        | Ceratodon sp.           | KC291530         | 99.8         | 1721/1724           |
|                   |                |                   |                  |             | Ceratodon purpureus     | Y08989           | 99.7         | 1751/1757           |
|                   |                |                   |                  |             | Ceratodon purpureus     | KC291530         | 99.9         | 1677/1678           |
|                   |                | 2: isolate protonema |               |             | Ceratodon purpureus     | Y08989           | 99.7         | 1673/1679           |
|                   |                |                   | A8872997         | 1697        | GMGA_cloning (this study) | A8858433 | 100          | 1679/1679           |
|                   |                | 3: dried GMGA_cloning |            |             | Bryum caespiticium      | AFO23703         | 100          | 1697/1697           |
| Chloroplast gene  | rps4           | 2: isolate protonema | A8848717        | 674         | Ceratodon purpureus     | FJS72605         | 100          | 623/623             |
|                   |                |                   |                  |             | Ceratodon purpureus     | FJ572589         | 100          | 625/625             |
|                   |                |                   |                  |             | Ceratodon purpureus     | AF435271         | 100          | 561/561             |
|                   |                |                   |                  |             | Ceratodon purpureus     | AY908122         | 100          | 652/652             |
|                   |                |                   |                  |             | Trichodon cylindricus*  | AY908125         | 94.1         | 622/661             |
|                   |                | 3: dried GMGA_cloning | A8872999        | 684         | Bryum caespiticium      | AFS21683         | 99.9         | 667/668             |
|                   | trnL           | 2: isolate protonema | A8848718        | 482         | Ceratodon purpureus     | FJS72485         | 100          | 482/482             |
|                   |                |                   |                  |             | Ceratodon purpureus     | AF435310         | 100          | 482/482             |
|                   |                |                   |                  |             | Ceratodon purpureus     | AF435271         | 100          | 482/482             |
|                   |                |                   |                  |             | Glyphomitrium humilellum* | EU246911 | 94.2         | 438/465             |
|                   |                | 3: dried GMGA_cloning | A8873000        | 520         | Bryum caespiticium      | AY150351         | 100          | 492/492             |
|                   | apb-rbcL intergenic spacer | 2: isolate protonema | A980065        | 637         | Ceratodon purpureus     | AY881031         | 100          | 621/621             |
|                   |                |                   |                  |             | Ceratodon purpureus     | AY881034         | 100          | 621/621             |
|                   |                |                   |                  |             | Ceratodon purpureus     | AY881052         | 100          | 598/598             |
|                   |                |                   |                  |             | Cheilothela chloropus*  | AY881063         | 89.9         | 571/635             |
| Mitochondrial gene | nad5           | 2: isolate protonema | A8848719        | 1112        | Ceratodon purpureus     | AY908859         | 99.9         | 1093/1094           |
|                   |                |                   |                  |             | Ceratodon purpureus     | AY908862         | 99.9         | 1090/1091           |
|                   |                |                   |                  |             | Trichodon cylindricus*  | AY908863         | 99.5         | 1089/1095           |
|                   |                | 3: dried GMGA_cloning | A8872998        | 1107        | Bryum argenteum         | AY908945         | 100          | 1082/1082           |

*show second highly related species.
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Discussion

Glacial Moss Gemmee Aggregation (GMGA) is a novel moss aggregation

Mosses, in the form of “glacier mice”, have been previously recorded from supraglacial habitats [11,12]; however, the structure of GMGA is completely different from that of “glacier mice”. Whereas “glacier mice” are formed by the moss shoots, that level of cellular differentiation was not detected in GMGAs. These findings report first description of developing moss gemmee and protonema in the supraglacial environments. *Ceratodon purpureus*, which formed the GMGA observed in this study, is a cosmopolitan moss species widely distributed throughout entire continents [31] and is known to grow in extreme environments (i.e. polluted sites including highway shoulders and on coal and heavy metal mine tailings) [32]. *Ceratodon purpureus* also occurs in the cryosphere in high alpine areas, Antarctica [33–35]. In the Rwenzori Mountains, unfortunately inhabitation of *C. purpureus* around glacier had not directly observed by authors, however, *C. purpureus* has been detected at elevations from 2900 m to 3700 m [36] and *C. purpureus* specimen (PC0106302) taken at just below the Speke Glacier (4480 m a.s.l.) are stored in Muséum National d’Histoire Naturelle, Paris, France. These evidences would show that possibility of dispersal of spore or gemma from near glacier and depositions on the glacier surface by local wind circulation.

Adaptation of GMGA isolate to warmer temperature

The optimum temperature of polar mosses are widely distributed from 2°C to 35°C according to species [37]. The optimum temperature for the *C. purpureus* isolates (25°C; Fig. 6) is normal value even in polar region, but this was higher than that for *C. purpureus* in Antarctica as previously reported, which was 15°C in the liquid and agar cultures [33]. Moreover, another study showed that the optimum temperature for photosynthesis in *C. purpureus* is around 15°C, but significant carbon fixation occurs at 5°C [35]. Although the measurement of optimum temperature by measuring fluorescence of chlorophyll used in the present study is an indirect measurement of growth (e.g., [39]), these values reflect photosynthetic ability at each temperature (e.g., [35]). Therefore, the populations of *C. purpureus* from this Ugandan glacier have likely adapted to a higher optimum temperature than Antarctic populations.
Cold and light stress in isolated *C. purpureus*

Polar mosses tend to adapt broad range of favorable temperature. For example, relationship between net assimilation rate (NAR) and temperature of *Drepanocladus uncinatus* in Signy Island show that optimum temperature is 20°C and NAR are more than 40% from 0°C to 30°C (5°C interval) by using cold incubation sample (5°C in light/−5°C in dark) [37]. Relatively higher temperature optimum and broad range of favorable temperature is similar to our result of GMGA and *C. purpureus* isolates. Optimum temperature shift with environmental temperature change were reported from some of experiments [38,39], but *Drepanocladus uncinatus* in Signy Island [37] did not correspond to these studies. Also in our study, optimum temperature did not shift to lower temperature inspite of preincubation temperature at 4°C. The internal temperature of the GMGAs (Fig. 5) was below the optimum (Fig. 6), therfore, the daytime internal temperature of GMGA (10°C) is not optimum but favorable for growth of Ugandan glacial *C. purpureus*. In this temperature range of internal GMGAs (−5°C to 10°C), the optimum PAR is below the warmer temperature (Fig. 6). This phenomenon may be able to explain photoinhibition from cold stress. In low-temperature conditions, PSII is inhibited due to decreased rate of repair of damaged D1 protein and increased excitation pressure [40].

Polar mosses are able to survive short-term freezing and thawing cycles in summer and prolonged freezing in winter. *Bryum argenteum* taken from tropical to polar origin showed no damage after 10 days with a temperature regime of 5°C in light/−5°C in dark, and grew slowly under these conditions [37]. In Rwenzori, *C. purpureus* live under similar diurnal cycle (from −5°C to 10°C) through year and GMGAs structure would be formed slowly. After being deposited and frozen at below −20°C for 2.5 years, regeneration of the *C. purpureus* from east Antarctica is very active [33]. *Ceratodon purpureus* protonemata from GMGA can be isolated and grown at 4°C after half a year of cryopreservation at −80°C. Therefore, isolate of *C. purpureus* in this study have potential to survive both short-term and prolonged freezing stress.

PAR in this natural environment is higher than optimum PAR at 5–15°C. Although we did not measure PAR directly, an automatic weather station with a radiation meter was installed beside the Stanley Plateau (N0°22′34.55″, E 29°52′43.24″, 4750 m above sea level) by the Stations at High Altitude for Research on the Environment project [41]. According to their data, and assuming no significant divergence with today’s conditions, the maximum diurnal seasonal shortwave radiation (c.a. 400 W/m²) occurs around 2:00 PM. PAR (400–700 nm) generally comprises 50% of total solar radiation reaching the Earth’s surface. Assuming a conversion rate of 1 W/m² to 4.57 μmol/m²/s, our estimated maximum PAR is 914 μmol/m²/s when maximum shortwave radiation is 400 W/m². This value is higher than the optimum PAR in any temperature and this may cause photoinhibition in low temperature.

These results indicate that both low temperature and high radiation on the glacier are stress factors for *C. purpureus*. In acidic rivers in Japan, *Dicranaella heteromalla* (Hedw.) Schimp. remains in a prolonged protonema stage for several growing seasons without producing shoots or sporophytes [42], which researchers concluded was due to the water’s extraordinarily low pH (1.9–2.1). A similar prolonged protonema phase of the moss *Scopelophila catacratace* (Mitt.) Broth was reported in copper-rich sites as well in Japan [43]. Therefore, the low temperature and high radiation stress on the glacier, may keep *C. purpureus* in the gemmace and protonemal stage instead of developing into shoots.

Higher photosynthesis activity of GMGA than of isolates

The value of ETR from GMGAs was twofold that of isolates (Fig. 6), possibly due to differences in nutrient condition and the effects of other photosynthetic microorganisms. Growth conditions of moss may be more suitable in GMGA than in the artificial medium (liquid BBM) used in this study, because GMGA contains sufficient nutrients for effective growth. Yet, GMGA is not a simple aggregation of only moss, but also contains many other microorganisms. For example, we observed *Cylindrocystis brebissonii* cells and red snow algae [44], which is related to the green algae commonly found in supraglacial environments, in the GMGAs. Both moss and green algae affect the total photosynthetic activity of GMGAs.

Possible process of GMGA formation

If biological material in sites without GMGAs form a thick deposition layer (more than a few millimeters), the temperature below the surface would increase to above 0°C, the same as in GMGAs. If this is so, then the invasion of gemmace of *C. purpureus* adapted to warmer temperatures on the cold glacier surface can be attributed to this increased subsurface temperature. Similar temperature increases occur in other glaciers (e.g., Qiyi Glacier); however, GMGA-like structures and growth of moss gemmace have not been found on any other glacier, despite studies of glacier biology being conducted around the world [45]. This may relate to possible inhabitation of *C. purpureus* near the glacier and unique features of the Rwenzori; namely, the lack of a clear seasonal temperature cycle. On Stanley Plateau, diurnal temperature change in all seasons is in a range of approximately 0°C to 5°C [41]. Consequently, long periods of freezing do not exist, permitting microorganisms to grow throughout the year. Therefore, we suppose that at least these two factors (the internal temperature rise and the long growth season) may contribute to the formation of GMGAs.

However, GMGAs are disproportionally dominant near the glacier terminus. The high number of GMGAs observed at ST1 must be supported by factors specific to that site. Although our data does not let us reach firm conclusions about the distribution of GMGAs, downward transporation by surface melting water and availability of sunlight is a likely candidate factor.

Glacier surface melt of Stanley Plateau homogenously spread over the ice area and remarkable water channels on surface are few. Takeuchi [46] speculated cryoconite granules (around 1 mm diameter) are more stable from meltwater than unicellular microorganisms due to larger size character. GMGAs are much larger diameter than typical cryoconite (Fig. 3) seems to be more stable on the ice. Also GMGAs penetrate few mm into ice due to radiation warming (Fig. 2b). These also prevent to wash this material to downward. Therefore, downward transportation of well-developed GMGAs seems unlikely happened.

During the biological growth season, supraglacial light conditions generally change based on depth of snow cover. In the early melt season snowfree season is removed by melting to expose the glacier ice surface to sunlight at lower elevations only. The snow line then retreats until reaching equilibrium line altitudes at the end of melt season. This leaves the entire surface of the ablation zone snow-free, making it an available habitat for photosynthetic microorganisms. As a result, the biodiversity of glacier microbial communities changes with elevation [44,46–48].

In early February 2013, the entire glacier surface was covered by snow except for a steep slope at the glacier terminus, where ST1 is located. The snow cover near ST2 was 0.85 in deep, but near ST1, zero or a few centimeters of snow cover were observed. Because snow blocks radiation, these differences in snow depth
cause variable light conditions and GMGA internal temperatures. Although the precise factors causing this difference in snow depth are unknown, slope angle and wind erosion likely to cause gradients in snow cover. We observed similar types of steep slopes on all edges of this glacier and other glaciers located beyond the ridge (Margarita Glacier: Fig. 2a); however, further observations and measurements must be necessary.

Direct ecological linkage between glacier and glacier foreland

In the glacier foreland immediately adjacent to the glacier terminus, we found an abundance of dried GMGAs on rock surfaces, which were likely left on the freshly barel subglacial rocks after glacier retreat. The temperature of these dried GMGAs on the rocks reaches approximately 20°C and conditions appear much drier than on the glacier, where water is supplied by melting (Fig. 5). Dried GMGAs create a soil-like structure on the abiotic rock surface with gametophyte of different dominant Bryophyta (Bryum sp.). The successional of dominant bryophyta species from C. purpureus gemmae to Bryum sp. shows that GMGAs had changed after leaving the glacier. Previous studies conceptually proposed the linkage between glacier and glacier foreland by nutrient connection [49] and outwash of cryoconite granules [2]. Otherwise, our findings indicate that GMGAs accumulate as a soil-like structure on the abiotic rock surface, directly linking the glacier and glacier foreland ecosystems.

Furthermore, another linkage between glacier and glacier foreland was found from subglacial environment [50,51]. In Canadian Arctic, varieties of mosses regenerate from old glacier and glacier foreland ecosystems. Soils-like structure on the abiotic rock surface, directly linking the nutrient connection [49] and outwash of cryoconite granules [2]. Otherwise, our findings indicate that GMGAs accumulate as a soil-like structure on the abiotic rock surface, directly linking the glacier and glacier foreland ecosystems.

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Author Contributions

Conceived and designed the experiments: JU YT. Performed the experiments: JU ST KY YT DS. Analyzed the data: JU YT. Contributed reagents/materials/analysis tools: JU YT HM SI SK. Wrote the paper: JU YT DS SK.

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