



Conference Abstract

The Undiscovered Biosynthetic Potential of the Greenland Ice Sheet Microbiome

Ate H. Jaarsma[‡], Katie Sipes[‡], Athanasios Zervas[‡], Francisco Campuzano Jiménez[‡], Andrea Claire Smith[‡], Liessel Victoria Svendsen[‡], Lea Ellegaard-Jensen[‡], Mariane S. Thøgersen[‡], Peter Stougaard[‡], Liane G. Benning[§], Martyn Tranter[‡], Alexandre M. Anesio[‡]

‡ Aarhus University, Roskilde, Denmark

§ German Research Centre for Geosciences, Helmholtz Centre, Potsdam, Germany

Corresponding author: Ate H. Jaarsma (ahj@envs.au.dk)

Received: 14 Jun 2023 | Published: 13 Oct 2023

Citation: Jaarsma AH, Sipes K, Zervas A, Jiménez FC, Smith AC, Svendsen LV, Ellegaard-Jensen L, Thøgersen MS, Stougaard P, Benning LG, Tranter M, Anesio AM (2023) The Undiscovered Biosynthetic Potential of the Greenland Ice Sheet Microbiome. ARPHA Conference Abstracts 6: e108004. https://doi.org/10.3897/aca.6.e108004

Abstract

The Greenland Ice Sheet is a biome primarily driven by microbial activity. Despite the harsh conditions, such as cold temperatures, low nutrient levels, high UV radiation in summer, and long dark winters, various niches can be found on the ice sheet that can support organisms capable of withstanding these challenges. During the summer, eukaryotic glacier ice algae grow in large quantities on the ice surface, accompanied by a community of bacteria, fungi, and viruses. Additionally, cryoconite holes and snow serve as habitats with their own distinct microbial communities. Nevertheless, the microbiome of supraglacial habitats remains poorly studied, leading to a lack of representative genomes from these environments. In this study, we conducted a comprehensive investigation of the supraglacial microbiome using both culturing-dependent and -independent methods. We compared genomes obtained through metagenomic sequencing (133 high-quality metagenome-assembled genomes or MAGs) and whole genome sequencing (73 bacterial isolates) to the metagenome assemblies to determine their abundance within the total environmental DNA. Interestingly, the isolates obtained in this study were not dominant taxa in their respective habitats, unlike the MAGs.

[©] Jaarsma A et al. This is an open access article distributed under the terms of the Creative Commons Attribution License (CC BY 4.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Under-investigated extremophiles, such as those inhabiting the Greenland Ice Sheet, may offer an untapped reservoir of undiscovered chemical diversity. We cataloged the biosynthetic potential of these organisms by examining the presence of biosynthetic gene clusters (BGCs) in the obtained genomes. To accomplish this, we utilized tools like the Antibiotics and Secondary Metabolites Analysis Shell (AntiSMASH) and the Biosynthetic Gene Similarity Clustering and Prospecting Engine (BiG-SCAPE) to mine these genomes and subsequently analyze the resulting predicted BGCs. We identified a total of 849 BGCs, which were organized into 411 gene cluster families (GCFs). Notably, the MAGs and isolate genomes exhibited distinct pools of biosynthetic diversity, with only 5 GCFs shared between the two groups. The cryoconite genomes yielded the most unique GCFs. Furthermore, we found evidence for the capacity of these microbes to produce antimicrobials, carotenoids, and osmoprotectants. However, many of the obtained BGCs could not be matched to similar, previously described BGCs, highlighting the vastness of the undescribed biosynthetic potential present in microbes from the Greenland Ice Sheet.

Keywords

supraglacial habitats, extremophiles, metagenomic sequencing, biosynthetic gene clusters

Presenting author

Ate H. Jaarsma

Presented at

Preferably oral, otherwise poster presentation. Theme: part 1. Natural settings (Glaciers, ice sheets, and permafrost)

Funding program

The <u>Deep Purple project</u> received funding from the European Research Council (ERC) under the European Union's Horizon 2020 research and innovation programme under grant agreement No 856416.

Conflicts of interest

The authors have declared that no competing interests exist.