A COMPARATIVE STUDY OF ENDOLITHIC MICROBIAL COMMUNITES BETWEEN DIFFERENT TYPES OF ROCKS IN THE NORWEGIAN HIGH ARCTIC

Korea Polar Research Institute

Y. H. Choe, Y. K. Lee*, M. Kim, J. S. Woo, M. J. Lee, **Korea Polar Research Institute**

yhchoe@kopri.re.kr, yklee@kopri.re.kr*

Introduction

In extreme environments, such as the Arctic, endolithic communities are most of the extant life. The endolithic environment is thought to buffer microbial communities from intense solar radiation, temperature fluctuations, wind, and desiccation in environments where such environmental factors inhibit epilithic growth. Therefore, the endolithic environment is a critical habitat to explore in exobiological research. The abundance of endolithic life in the high Arctic, combined with the potential for biosignature preservation, suggests that rocks associated with endolithic ecosystems may be the best hope for finding fossil evidence of past life on the Martian surface. The goal of this study is to assess the activity and community composition of the endolithic community inhabiting rocks in the Norwegian high Arctic (Fig. 1). To achieve this goal, we combined techniques of DNA sequencing, microscopy and spectroscopy. This work provides an overview of the polar microbial community and a new recognition of how environmental stresses such as prevailing polar desert conditions may affect the biogeochemical dynamics of high Arctic endolithic microorganisms.

Results

SEM images allow one to describe morphotypes and to estimate roughly the dimensions of the structures, but it is not possible to obtain information on the spatial organization of the organisms within the endolithic band. The images obtained illustrated bacterial colonization in the rocks. They showed a wide distribution of single coccoid cells of about 5 µm in diameter attached to surface, small globules forming aggregates covered grains of rock, and fine filamentous structures of 0.3 to 0.5 mm in width, lead to a network in the crevices of the rock (**Fig. 3**).





Figure 3. Colony of bacterial cells visualized by scanning electron microscopy growing along the rock surface

At the phylum level, the rock bacterial community was dominated by five bacterial phyla, Actinobacteria, Proteobacteria, Bacteroidetes, Acidobacteria, and Chloroflexi (Fig. 4). Other phyla such as Deinococcus-Thermus, Verrucomicrobia, Gemmatimonadetes, and Planctomycetes were also present consistently across all samples, but in lower abundance. The relative abundance of major bacterial phyla varied considerably among samples. Specifically, the largest variation was found in Proteobacteria and Deinococcus-Thermus, and to a lesser extent Acidobacteria and Bacteroidetes.

(A)

(B)







Figure 1. Study sites on Svalbard (A) Troll spring, (B) Halvdanpiggen woodfjord, (C) Dasan station-southeast, (D) Scheteligfjellet

Materials and Methods

Endolithic samples were collected in August 2014 at various sites in Spitsbergen on Svalbard, Norway (Fig. 2). Samples were collected with hardened steel chisels that were flame sterilized in the field with 100% ethanol and a butane torch. Duplicate rock samples at each site were collected into sterile, plastic tubes and bags, and transported to the laboratory in an icebox keeping under ~4°C. For scanning electron microscopy (SEM), rock pieces were sputtercoated with gold and examined with scanning electron microscope at 20 kV. Sequences generated from pyrosequencing of bacterial 16S rRNA gene amplicons were processed using the mothur pipeline. Statistical analyses were performed using the vegan R package.



Figure 4. Bacterial Phyla composition in various rocks (A) Carbonate, (B) Basalt, (C) Sandstone, (D) Chert

Discussion and Implications

Figure 2. Various rock samples (A) Troll spring – Carbonate, (B) Halvdanpiggen woodfjord - Basalt, (C) Dasan station-southeast – Sandstone, (D) Scheteligfjellet - Chert

Phylogenetic and statistical comparison of endolithic communities from this study with those from previous studies in similar environments support the hypothesis that patterns of microbial diversity are governed by similar principles observed in macro-ecological systems. The majority of the endolithic bacterial sequences were most closely matched to those of isolates or clones derived from cold/dry deserts (i.e., Antarctica, Atacama, and cold alpine environments). This indicates that endolithic bacteria in this region have little site variations and are under similar degrees of environmental constraints of the cold and dry Arctic. Results also provide insight into geobiological processes that shape the biosphere and help us understand the cold and dry envionments possibly elsewhere in the Solar System.

References

[1] Starke V. et al. (2013) Environmental Microbiology Reports, 5, 648-659. [2] Smith H. D. et al. (2014) International Journal of Astrobiology, 13 (3), 271–277. [3] Horath T. et al. (2005) Microb Ecol. 51(3), 353-64. [4] Cary S. et al. (2010) Nature Reviews. Volume 8. [5] Ziolkowski L. A. et al. (2013) Biogeosciences, 10, 7661–7675

Acknowledgements

This study was supported by the "Environmental Change Studies based on the Arctic Dasan Station: in terms of Geology, Atmospheric Science, and Ecology (PE15030)" funded by the Korea Polar Research Institute.