

Genetic Origins of Mercury Resistance in Great Salt Lake Microorganisms Ashtyn Smith, Austin Wood, Chelsea Lam, Jaimi Butler, Bonnie Baxter

Introduction

Extremophiles are a diverse group of organisms, typically Bacteria and Archaea, that can inhabit extreme environments, such as geysers, deserts, and saline lakes. Their abilities to withstand extremely dry, hot, salinic, acidic, and mercuric conditions have made these microorganisms admirable astrobiological models for life on other planets¹.

Background

Methylmercury (CH_3Hg) is a neurotoxin that accumulates in aquatic environments due to the actions of microorganisms, which can produce this biologically relevant organic form from elemental mercury (Hg) (Figure 1, right).

Many species of microorganisms have shown resistance to Hg and can thrive in polluted waters. Recent studies have shown that Hg resistance in Bacteria and Archaea arises from one of two gene pairs, *mer*AB or *hgc*AB^{3,4}. The *mer*AB system produces gene products that allow the organism to convert CH₃Hg into elemental Hg³. Conversely, the *hgc*AB system coverts Hg into CH₃Hg⁴ (Table 1). Through these mechanisms, microorganisms can play a significant role manipulating the health of aquatic ecosystems.

Mercury Resistant Genotypes					
Gene	Function				
hgcA	Aids in methylation of Hg, gene products currently unknown ⁴				
hgcB	Aids in methylation of Hg, gene products currently unknown ⁴				
merA	Aids in demethylation of CH ₃ Hg through production of mercury reductase ³				
merB	Aids in demethylation of CH ₃ Hg through production of organomercurial lyase ³				

Table 1. Each mercury-resistant genotype consists of two gene pairs. The hgcAB gene pair assists in mercury resistance by methylating Hg⁴. The metabolic pathway in which this occurs is currently being studied and specific gene products have yet to be published. The merAB system, on the other hand, is more defined. The gene products of this system, mercury reductase and organomercurial lyase, work together to demethylate CH₂Hq³.

Due to natural and industrial influences, the Great Salt Lake (GSL) has accumulated Hg within its waters. Although the lake has no fish, mercuric bioaccumulation has extended from the microbial and shrimp populations to terrestrial animals, such as spiders and birds. As previously suggested, the GSL microbes may have a significant influence over the production of CH_3Hg from Hg inputs. Therefore, defining the genotype of mercury-resistant GSL microorganisms is essential to understanding the behavior of CH₃Hg in this ecosystem and may inform future bioremediation attempts on the lake.

Hypothesis

GSL microorganisms that demonstrate a robust resistance to mercuric conditions will express either the hgcAB or merAB genotype.

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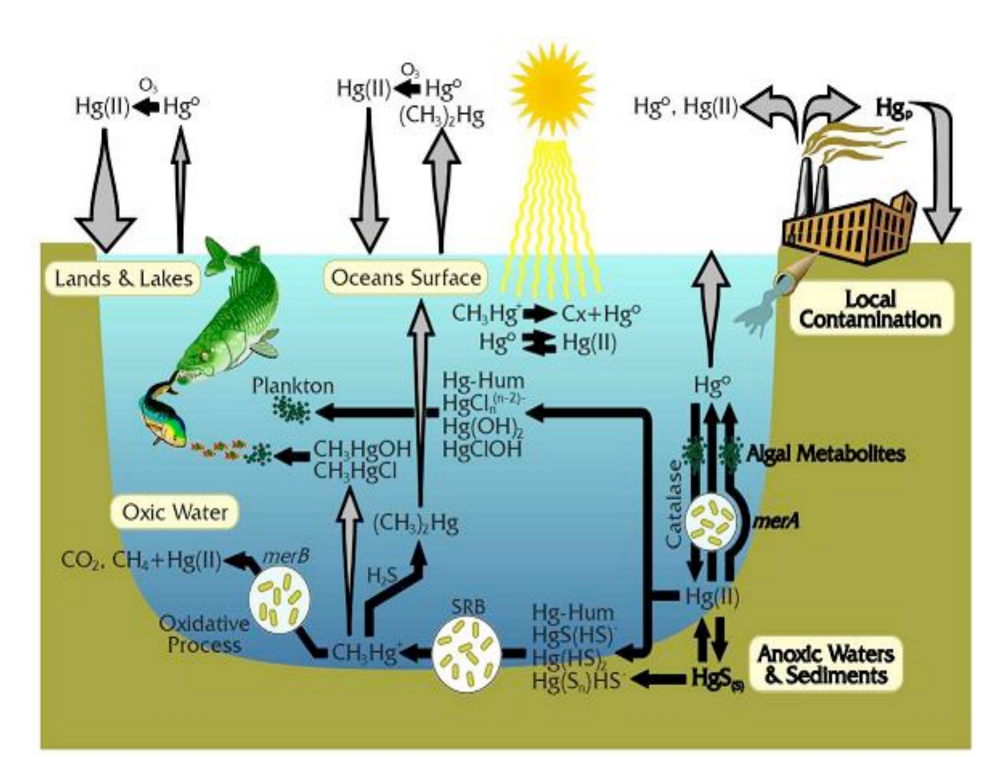


Figure 1. Through the actions of microorganisms and local industry, Hg and CH₃Hg concentrations are manipulated and bioaccumulated in aquatic ecosystems².

Objectives

- 1. Identify mercury-resistant microorganisms in the GSL.
- 2. Establish the range of mercury resistance.
- 3. Determine the mercury-resistant genotype.

Materials & Methods

Overview: Microorganisms will be harvested from the GSL and cultivated on increasing concentrations of mercury. The most resistant microbes will undergo genetic analysis to identify resistant genotypes in the population.

Harvest, Cultivation, & Isolation

Anaerobic halophiles, "salt-loving" microorganisms, were obtained from the deep brine layer of the GSL. This method was repeated in 8 different locations of the lake in order to observe a broader sample of the GSL microbial community. Locations are listed in Tables 2, 3 & 4.

Samples were cultivated in broth culture before being transferred to petri dishes containing modified growth medium (MGM)⁵ infused with 5 ppm mercury chloride (Hg(II)Cl₂), a form of Hg represented in the GSL. The plates were representative of a range of salinities, 12%, 18%, 23%, and 25%, as well. Cultures were incubated at 37° C in anaerobic chambers.

Determining Mercury Resistance

Individual colonies were selected from the 5 ppm Hg(II)Cl₂ plates and transferred to 10 ppm Hg(II)Cl₂ plates with corresponding salinities. After sufficient colony growth was observed, the specimens were transferred to 20 ppm Hg(II)Cl₂. In this way, mercury concentrations were slowly increased in order isolate the most resistant microorganisms. All plates were incubated at 37° C in anaerobic chambers.

The microorganisms that demonstrated the greatest resistance to Hg(II)Cl₂ are being analyzed for *hgc*AB and *mer*AB. Additionally, the 16S rRNA gene, which will determine the identity of these microbes, will be assessed, as well. Analysis of DNA extracts will be carried out with PCR amplification, gel electrophoresis, and genetic sequencing.

Mercury-resistant halophiles have successfully been cultivated on 5, 10, and 20 ppm Hg(II)Cl₂ plates across various salinities, distributions represented Tables 2, 3, & 4. Of the eight sampling sites, 46, 26, and 25 distinct colonies were exhibited on 5, 10 and 20 ppm Hg(II)Cl₂ plates, respectively.

Genetic Analysis

Results

Isolates on 5 ppm Hg(II)Cl ₂ MGM								
Site	12%	18%	23%	25%				
Bear River Bay	1	1	2	1				
Bear River Pond 3E	3	1	1	1				
Bear River Pond 5C	1	1	0	0				
South Shore	1	1	1	1				
North Basin	3	3	2	1				
armington Bay	2	1	1	1				
North Arm/Gunnison sland	2	2	2	2				
Ogden Bay	5	1	0	1				

Table 2. From the 8 sampling locations, 46 distinct colonies were presented on plates containing 5 ppm Hg(II)Cl₂.

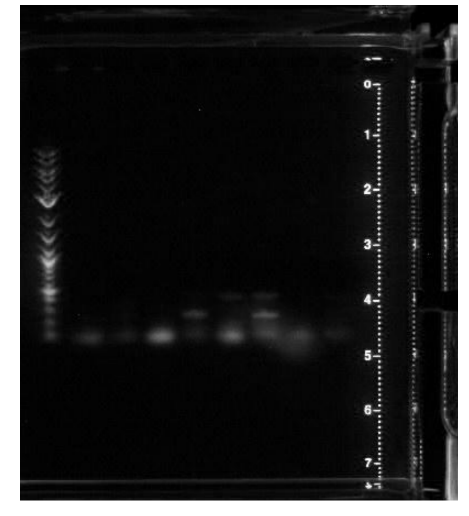
Isolates on 10 ppm Hg(II)Cl ₂ MGM								
Site	12%	18%	23%	25%				
Bear River Bay	0	1	1	1				
Bear River Pond 3E	2	1	0	1				
Bear River Pond 5C	0	1	0	0				
South Shore	1	0	0	1				
North Basin	3	0	0	1				
Farmington Bay	1	1	0	1				
North Arm/Gunnison sland	2	1	1	2				
Ogden Bay	2	0	0	1				

Table 3. Of the 46 colonies, only 26 colonies continued to grow on 10 ppm Hg(II)Cl₂ plates.

Isolates on 20 ppm Hg(II)Cl ₂ MGM								
Site	12%	18%	23%	25%				
Bear River Bay	0	1	1	1				
Bear River Pond 3E	2	1	0	1				
Bear River Pond 5C	0	1	0	0				
South Shore	1	0	0	1				
North Basin	3	0	0	1				
Farmington Bay	1	1	0	1				
North Arm/Gunnison sland	2	1	1	1				
Ogden Bay	2	0	0	1				

Table 4. Of the 26 colonies grown in 10 ppm Hg(II)Cl₂ all but one colony thrived on the 20 ppm $Hg(II)CI_2$ plates.

Four 20 ppm Hg(II)Cl₂ isolates were analyzed for hgcA, merAB, and 16S rRNA. Gel electrophoresis and sequencing of the resulting PCR products did not yield conclusive results. According to the data shown in Figures 2 and 3, two strains demonstrated both genotypes. However, gene sequencing was not successful and could not confirm these findings. More tests will be needed before drawing any conclusions.



respectively

GSL microorganisms demonstrate a robust resistance to mercury chloride, as was shown in Tables 2, 3, and 4. The mercury-resistant genotype of these halophiles, however, were not identified. Initial analysis of four 20 ppm Hg(II)Cl₂ isolates suggests that two microbes express the *hgc*AB genotype while two others seem to demonstrate both hgc and mer genotypes (Figures 2 & 3). Despite these results, genetic sequencing was not conclusive. More experiments will be needed before extending any conclusions.

project:



Results Continued

Figure 2. From left to right, hgcA and B PCR product alternates after the DNA ladder shown in lane 1. Lanes 2 & 3 and 4 & 5 correspond with the organisms represented in lanes 2 & 3 and 4 & 5 of Figure 3,

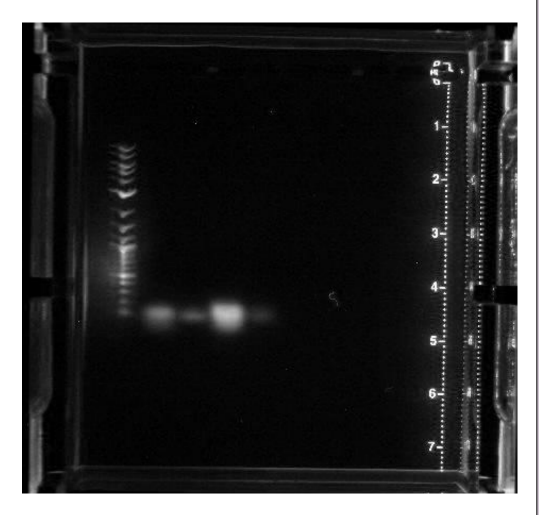


Figure 3. From left to right, *merA* and B PCR product alternates after the DNA ladder shown in lane 1. Lanes 2 & 3 and 4 & 5 correspond with the organisms represent in lanes 2 & 3 and 4 & 5 of Figure 2, respectively.

Conclusions

Acknowledgments

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References

[1] Baxter BK, Eddington B, Riddle MR, Webster TN, Avery BJ. 2007. Great Salt Lake halophilic microorganisms as models for astrobiology: Evidence for desiccation tolerance and ultraviolet irradiation resistance. SPIE Proceedings, 6694. doi: 10.1117/12.732621

[2] Barkay T. n.d. Barkay Research. Retrieved from http://aesop.rutgers.edu/~barkay/TBRESP.htm [3] Narita M, Huang CC, Koizumi T, Yamagata T, Endo G. 2000. Identification and characterization of anaerobic mercury-resistant bacteria from mercury-polluted sediment. Water Science and Technology: A journal of the International Association on Water Pollution Research. 42(3/4):109-114. [4] Parks JM, Johs A, Podar M, Bridou R, Hurt Jr. RA, Smith SD, Tomanicek SJ, Qian Y, Brown SD,

Brandt CC, Palumbo AV, Smith JC, Wall JD, Elias DA, Liang L. 2013. The Genetic Basis for Bacterial Mercury Methylation. Science. 339:1332-1335.

[5] Dyall-Smith M. 1998-2009. The Halohandbook. n.p.