

Biodegradation of microcystins in Lake Erie source waters and sand filters from drinking-water plants

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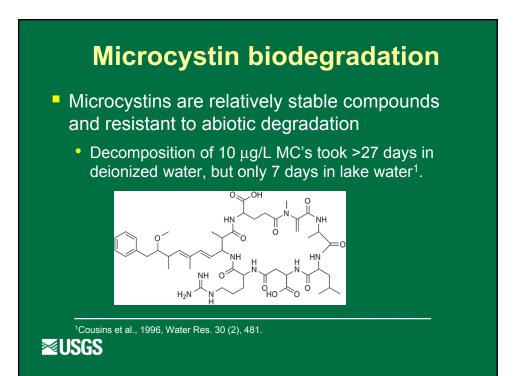
U.S. Department of the Interior U.S. Geological Survey

Addressing cyanoHABs requires multiple strategies

- 1. Reduce sources and causes
- Monitor for and predict cyanoHAB occurrence and toxin concentrations
- Water treatment and bloom control strategies to reduce or eliminate toxins

Microcystin removal in water

- Chemical and physical methods for removal of MCs have efficiency and cost limitations
 - Cell bound—coagulation, sedimentation, filtration
 - Extracellular—activated carbon, chlorination, ozonation, permanganate, UV, membranes
- Biodegradation is environmentally friendly and could augment existing treatments



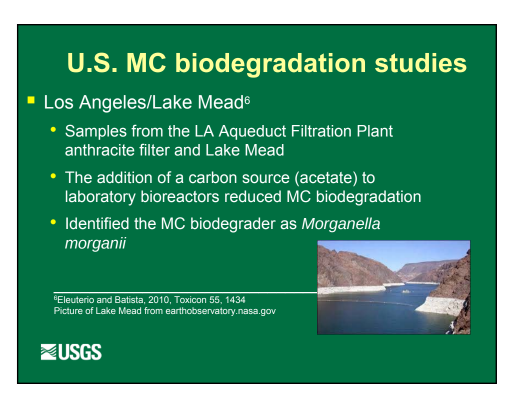
Microcystin biodegradation studies

Indigenous MC-degrading bacteria identified in lakes, rivers, and water-treatment plant biofilms

- Studies in the 1990's in Australia and Japan
- Identified as Pseudomonas sp³ and later reclassified as Sphingomonas sp⁴
- Other genera identified in Europe, Australia², and Asia
- Bacteria may contain the *mlrA* gene⁵

²Ho, L. et al., 2006, Water Research 40, 768
 ³Jones et al., 1994, Nat. Toxins 2 (4), 228
 ⁴Bourne et al., 1996, Appl. Environ. Microbiol. 62 (11), 4086
 ⁵Saito et al., 2003, FEMS Microbiol. Lett. 229 (2), 271.
 Sphingomonas from Wikepedia.

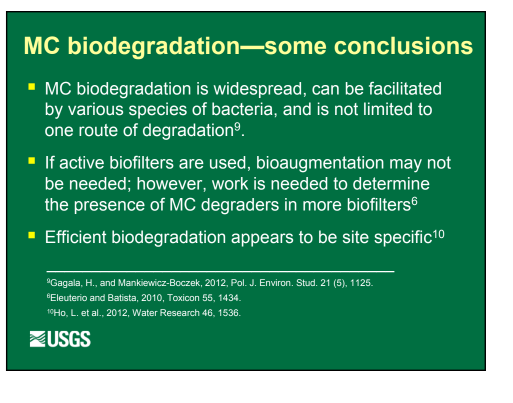




U.S. MC biodegradation studies

- Lake Okeechobee⁷
 - Phosphorus levels did not affect biodegradation.
 - Identified *Micobacterium* and *Rhizobium* gallicum as MC biodegraders
- Lake Erie⁸
 - Offshore samples
 - Used metagenomics to identify a diverse array of bacterial phyla as potential MC biodegraders

⁷Ramani et al., 2011, Biodegradation 23 (1), 35.
⁸Mou et al., 2013, Appl. Environ. Microbiol. 75(21), 6924.



USGS Western Lake Erie study Objectives, 2015–17

- Identify naturally occurring microcystin-degrading bacteria from source waters and sand filters in drinking-water plants in the Western Basin of Lake Erie and watershed.
- Identify whether a molecular target, such as the *mlrA* gene, can be used to quantify biodegradation of microcystins.
- Compare rates to those of a known microcystin biodegrader.

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Timeline

- Aug–Dec, 2015 and July–Aug, 2016—Samples from source waters and sand filters
- Aug 2015–Sept 2016—Laboratory work
 - Microcosms to determine biodegradation potential
 - Isolate and identify biodegraders
 - Compare the rate of biodegradation
- 2017—Data analysis and report

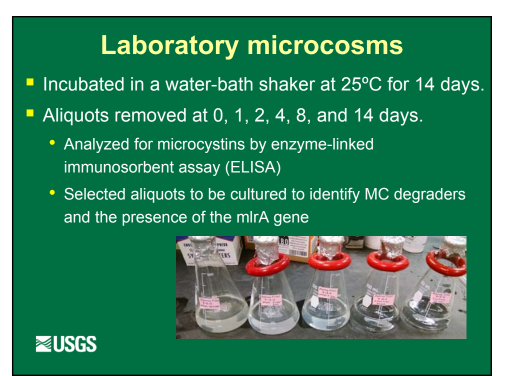


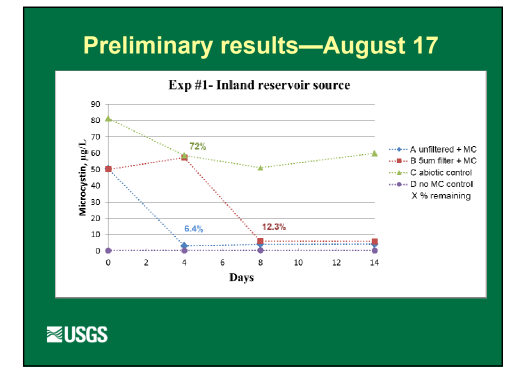
Laboratory microcosms

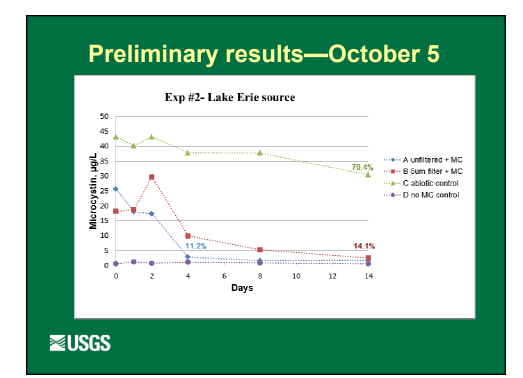
- A. MC + unfiltered source water
- **B**. MC + 5- μ m filtered source water
- C. MC + 0.2-μm filtered source water (abiotic control)
- D. Unfiltered source water (background control)

For sand filter experiments, used buffered water with 20 g sand









Preliminary results – Culture Work

- Samples cultured on R2A media at time 0 and at the time when degradation was seen (ie: time 4 and time 8 from flasks A and B from Exp #1)
- Colony morphologies that appeared to be enriched at the later time points were selected for further analysis
- Three-phase streaks done on LB media to generate pure cultures
- Glycerol stocks of each isolate are made for further testing and analysis.

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Preliminary results – Culture Work

Isolate from sand filter at time 0 viewed under UV light (right) and a non-fluorescent isolate (left)





Clyde BT, 1.10

1:1004

Spread plates prepared from glycerol stocks collected during microcosm incubation. Isolated colonies with varying morphologies collected from T8 for pure cultures.

Three-phase streak of an isolate

Preliminary results – Culture Work



