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## Title

*Dolichospermum* evolution during during cyanobacterial bloom: insights from metatranscriptomics.

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## Abstract

Specific environmental conditions, such as elevated levels of nutrients from human activities, warmer temperatures, still water, and sunlight promote the growth of cyanobacteria, which at high density, form harmfull algae blooms. Each summer, cyanobacteria make an appearance in Lake Champlain (Baie Missisquoi, Qc) and pose health risks to humans and animals due to the production of cyanotoxins. Here, we used a metatranscriptomics approach that targets genes expressed in the water to identify changes in the bacterial community during the summer. At the height of the bloom (mid-September), chlorophyll A levels increased by nearly 100 fold and communities were dominated by the cyanobacteria *Dolichospermum circinale* and to a lesser extent *Microcystis aeruginosa*. We identified 7,000 *Dolichospermum* SNPs (10% of all SNPs) that showed changes in allele frequencies from the begining of the bloom to the end of the season, showing a highly dynamic evolution of the system. Nineteen genes showed the same repeatable patterns of evolution in all three sampling sites (expected: three) and these represent good candidates to predict blooms in the future.

## Introduction

* Specific environmental conditions, such as elevated levels of nutrients from human activities, warmer temperatures, still water, and sunlight promote the growth of cyanobacteria, which at high density, form harmfull algae blooms.
* Each summer, cyanobacteria make an appearance in Lake Champlain and pose health risks to humans and animals due to the production of cyanotoxins.

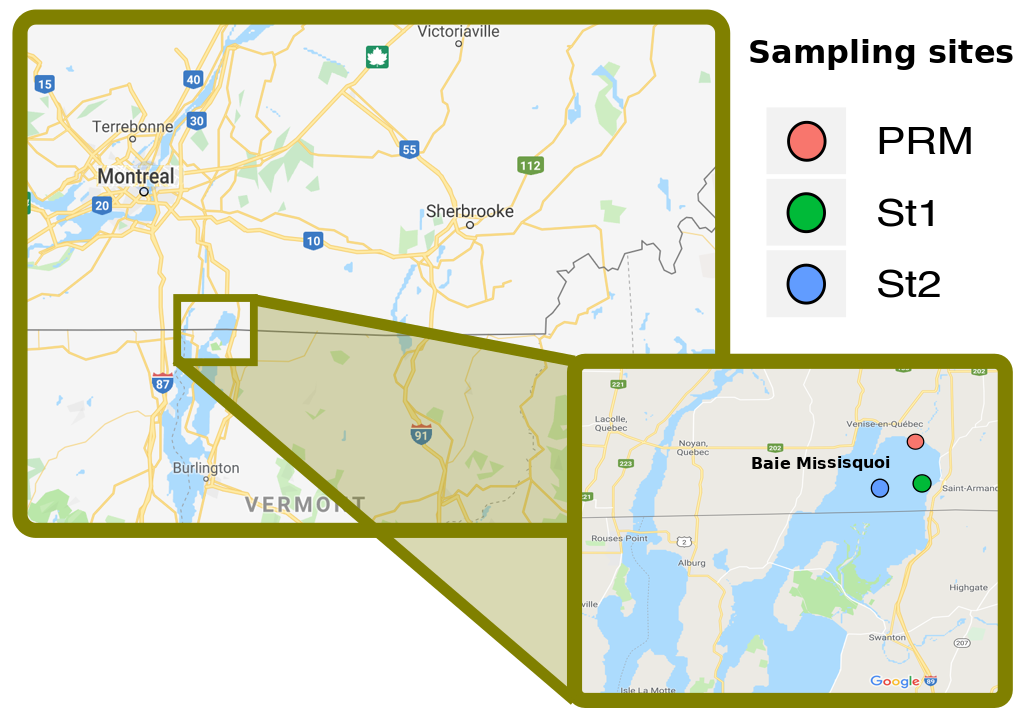
## Objectives

* Use a metatranscriptomics approach that targets genes expressed in the water to identify changes in the bacterial community during the summer.
* Identify *Dolichospermum* (main bloom causing cyanobacteria) genes that change in allele (SNPs) frequency from the beginning to the end of the bloom. These represent candidates genes whose changes in allele frequencies may help to predict blooms in the future.

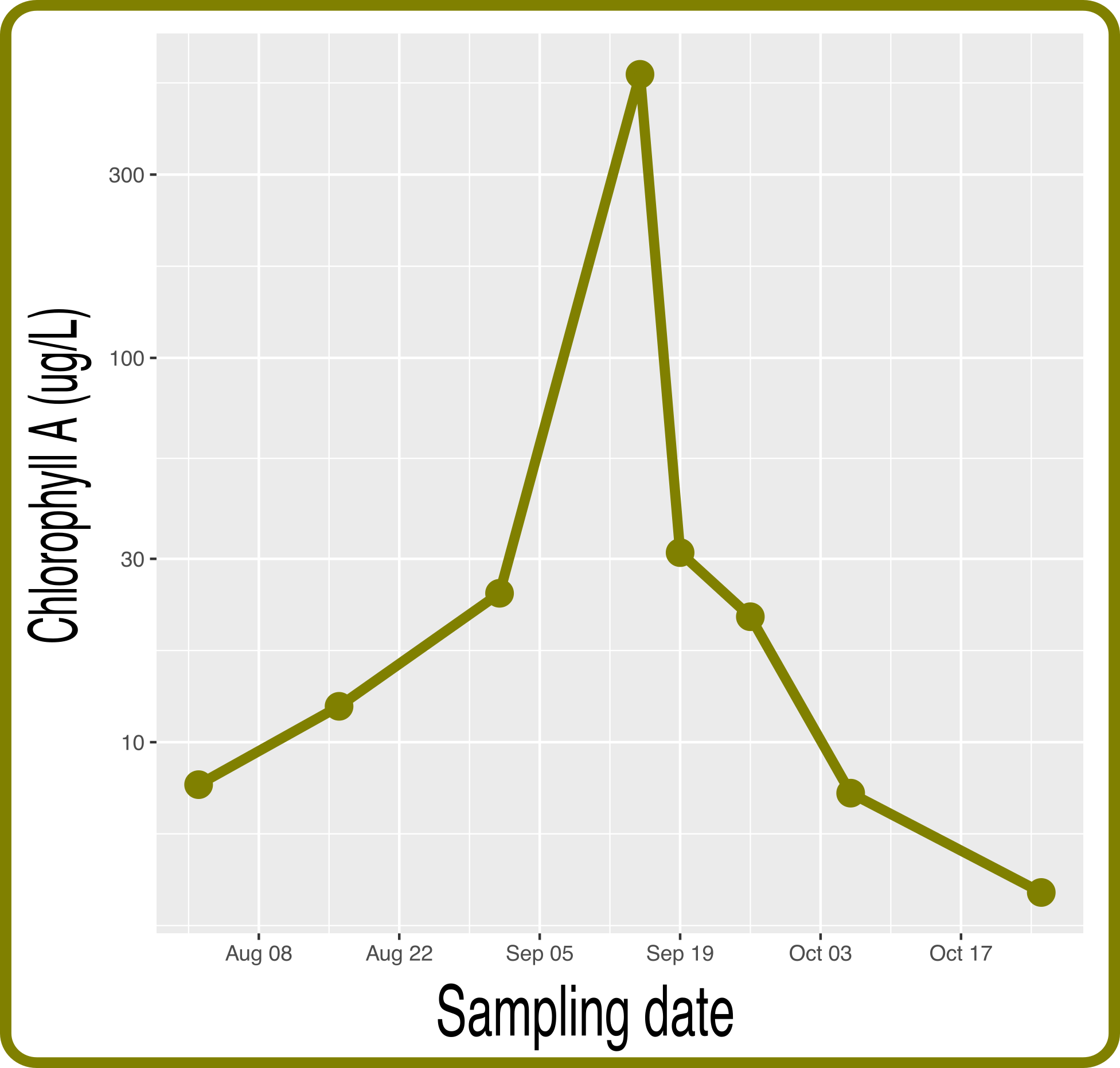
## Method

1. Water sampled at three location in Lake Champlain (Baie Missisquoi: St1, St2, PRM, **see Figure 1**) during the cyanobacterial bloom in the summer of 2016 (June 1st - October 10th).
2. RNA extracted, followed by High Throughput sequencing (100bp paired end Illumina HiSeq).
3. Metatranscriptomes processed (trimming, merging, removal of rRNA, annotation) using SAMSA2 pipeline1.
4. Sequences specifically annotated to *Dolichospermum* (the main cyanobacteria responsible for the bloom) used to produce a *de novo* reference transcriptome assembly using Trinity2
5. *Dolichospermum* sequences aligned to transcriptome using bowtie23. SNP called using SAMtools4.
6. Changes in allele frequency quantified using regressions (logistic for St1 & Ste, linear for PRM) in R.
7. Candidate genes showing significant changes (FDR corrected *p*-value < 0.05) in allele frequency matched to reference database and compared against reference transcriptome.

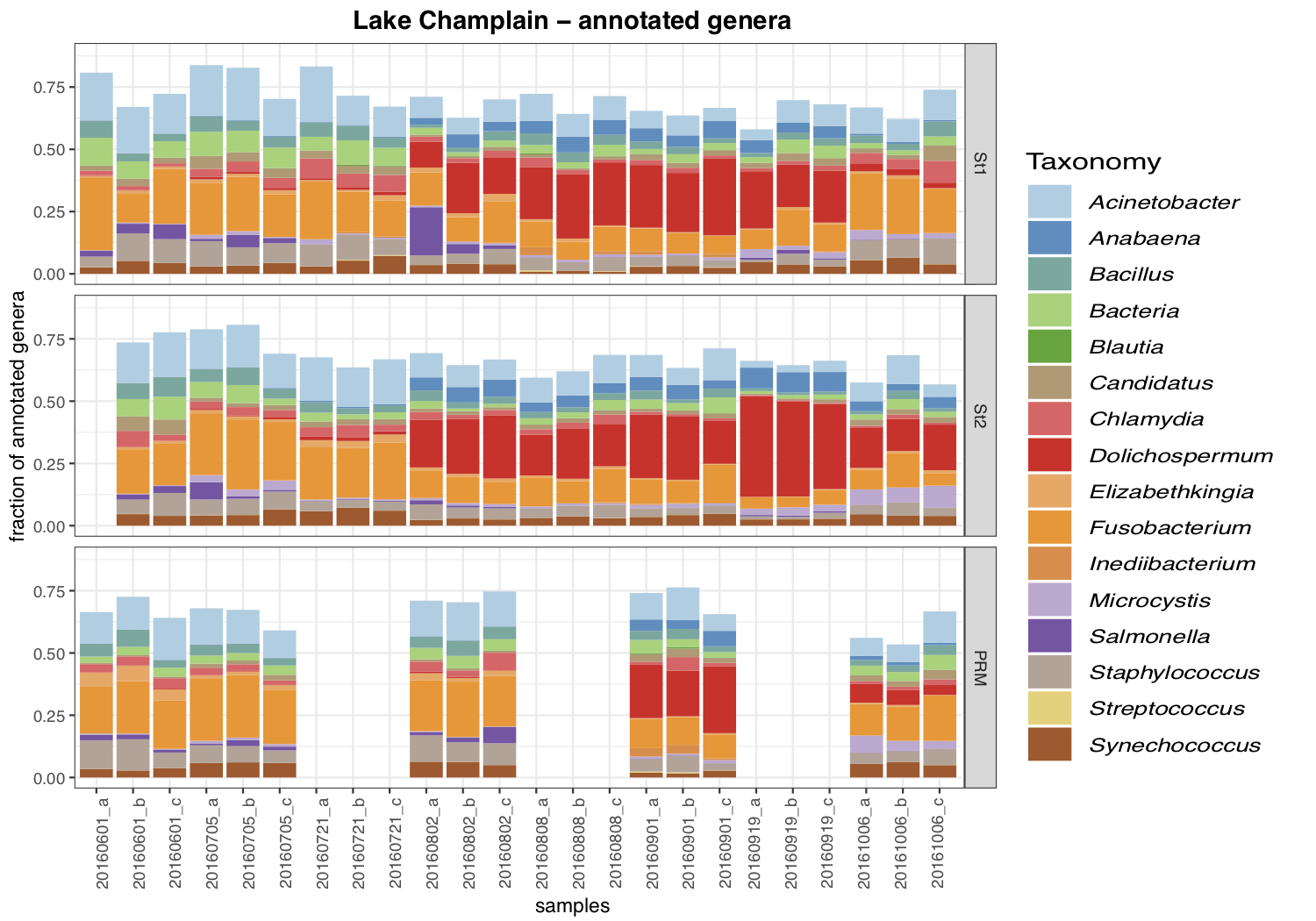
## Results



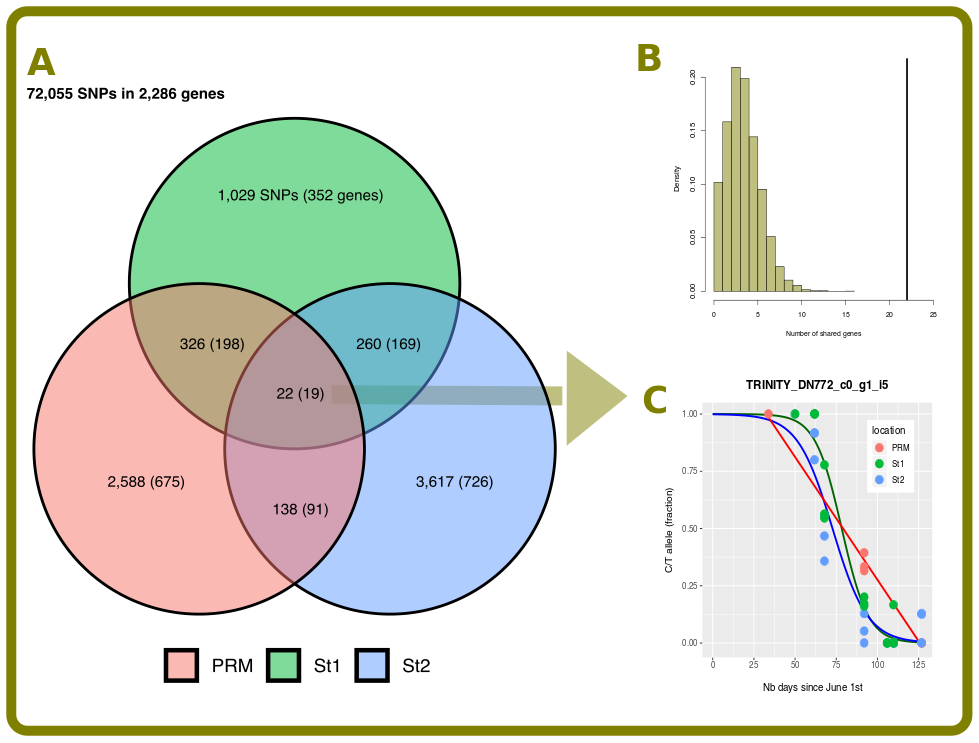
**Figure 1: A map of sampling sites**



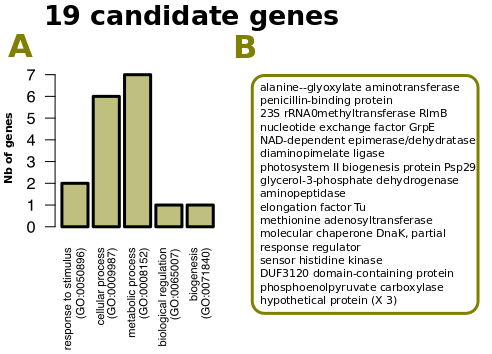
**Figure 2: Chlorophyll concentration peaked in mid-September (August 1st: 7.8 ug/L, Sept 15th: 547 ug/L) resulting in a cyanobacterial bloom**



**Figure 3: Communities dominated by Gram negative *Fusobacterium* & *Acinetobacter* prior to bloom. During Bloom, *Dolichospermum circinale* rose in frequency, followed to a lesser extent by *Microcystis aeruginosa* in September.**



**Figure 4: 72,000 SNPs identified in 2,286 *Dolichospermum* genes. A.Venn diagram: Number of SNPs (genes) showing significant changes in allele frequency from beginning (June 1st) to end of bloom (October 6th). C. Null distributions of the number of SNPs expected to show significant allele frequency change in all three sampling sites compared to observed (black vertical line, 22 SNPs observed, *p*-value < 0.0001). B. Example of a significant SNP in both St1, St2 (logistic regressions) and PRM (linear regression) sites.**



**Figure 5: A. Gene ontology groups for candidate genes (no significant over-represented groups compared to reference transcriptome). B. List of candidate genes.**

## Conclusions

* Metatranscriptomics can identify cyanobacteria known to cause blooms (*Dolichospermum circinale*, *Microcystis aeruginosa*).
* ~7,000 SNPs (10% of all SNPs) show changes in allele frequencies during the bloom in at least one of the sampling site, showing a highly dynamic evolution of the system. Nineteen genes show repeatable changes in allele frequencies in all three sampling sites. These represent good candidates to predict bloom.
* Further sampling and sequencing in progress in other lakes in order to confirm patterns observed here.

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## References

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4. Li, H. *et al.* The sequence alignment/map format and samtools. *Bioinformatics* **25**, 2078–2079 (2009).