Assessment of a Metaviromic Dataset Generated from Nearshore Lake Michigan

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Assessment of a metaviromic dataset generated from nearshore Lake Michigan

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Abstract. Bacteriophages are powerful ecosystem engineers. They drive bacterial mortality rates and genetic diversity, and affect microbiially mediated biogeochemical processes on a global scale. This has been demonstrated in marine environments; however, phage communities have been less studied in freshwaters, despite representing a potentially more diverse environment. Lake Michigan is one of the largest bodies of freshwater on the planet, yet to date the diversity of its phages has yet to be examined. Here, we present a composite survey of viral ecology in the nearshore waters of Lake Michigan. Sequence analysis was performed using a web server previously used to analyse similar data. Our results revealed a diverse community of DNA phages, largely comprising the order Caudovirales. Within the scope of the current study, the Lake Michigan virome demonstrates a distinct community. Although several phages appeared to hold dominance, further examination highlighted the importance of interrogating metagenomic data at the genome level. We present our study as baseline information for further examination of the ecology of the lake. In the current study we discuss our results and highlight issues of data analysis which may be important for freshwater studies particularly, in light of the complexities associated with examining phage ecology generally.

Introduction

The study of viruses in the environment, particularly those infectious to bacteria (bacteriophages) and other prokaryotes, has been driven in recent years by several exciting discoveries (Rice et al. 2004; Zhao et al. 2013). With the advent of improved genomic techniques for molecular-level analyses, microbiologists are slowly uncovering the depth of viral diversity indigenous to areas other than human physiology. Despite the viral renaissance, we still understand very little with regard to how viruses, particularly phages, interact \textit{in situ} with their environment and their hosts.

Phages play a crucial role in the structuring of all microbial communities. They mediate host mortality and drive bacterial genetic diversity and production (Winget et al. 2011) in a constant cycle of antagonistic coevolution (Buckling and Rainey 2002). Arguably, one of the most well characterised bacteriophage habitats is the marine (Suttle 2005). This is due largely to the vast impact that marine cyanobacteria have in the oceans in their role as ecosystem engineers (Berman-Frank et al. 2003), and the subsequent impact of marine cyanophages upon their ecology. The level of diversity in phage populations in freshwater communities is likely to exceed that found in marine environments, if only owing to the known habitat heterogeneity found in these systems (Maranger and Bird 1995; Wang et al. 2012; Berdjeb et al. 2013). Generally, freshwater microbial communities have received less attention from microbiologists in comparison to the marine (Ghai et al. 2014), especially freshwater phage communities. The investigation of environments that are likely to support highly diverse phage populations is an important step towards identifying key bacteria-phage interactions, such as that between \textit{F}SPM-2 and its cyanobacterial host, which allowed Clokie et al. (2006) to identify the virus as being ‘photosynthetic’ in nature. This will, in turn, inform our understanding of how phages drive important environmental processes.

Metagenomic techniques have become the favoured approach for large-scale examination of microbes in the environment. Although they have inherent limitations (as with all experimental techniques), and should be used to draw conclusions with regard to microbial communities within the scope of strong experimental design, they have allowed for a glimpse into phage diversity and relative abundances for a range of niches,
Viral particles were collected by filtering and concentration. The samples were passed through sterile 0.45-μm cellulose acetate membrane filters (Corning Inc., Corning, NY) and 0.22-μm polyethersulfone membrane filter (MO BIO Laboratories, Carlsbad, CA). Viral particles were then concentrated using a 0.10-μm polypropylene filter (EMD Millipore Corp, Billerica, MA) attached to the Labscale tangential flow filtration (TFF) system (EMD Millipore Corp, Billerica, MA). The filter was cleaned extensively between each sample, using a commercially available Tergazyme preparation.

**Viral DNA extraction**

Extraction of viral DNA was attempted from the 16 samples using the MO BIO Laboratories PowerWater DNA Isolation Kit (Carlsbad, CA), according to the manufacturer’s instructions with one alteration: after optimisation and validation experiments, an additional heat treatment at 70°C for 10 min before initial vortexing was added. It was not possible to extract DNA from all the viral fractions (a common problem encountered during the examination of environmental viruses (Hurwitz and Sullivan 2013)), and nine viral fractions yielded high concentration, high quality DNA. DNA was prepared without any additional amplification steps, therefore favouring the capture of dsDNA viruses (Kim and Bae 2011). To test for bacterial contamination, each extraction underwent PCR testing using primers for the bacterial 16S rRNA gene (Marchesi et al. 2001), alongside an *Escherichia coli* positive-control. No amplicons were produced by the viral DNA template.

**Sequencing and assembly**

Libraries were prepared from the nine samples using the Nextera XT kit for Illumina MiSeq library preparation. The nine samples were multiplexed for a single run of paired-end reads and sequenced by Genewiz (South Plainfield, NJ). Over 15 000 000 raw reads were generated: the raw FASTQ files can be found using NCBI’s BioProject database (Accession: PRJNA248239).

Paired-end assembly and contig assembly were performed for each sample using MetaVelvet (Namiki et al. 2012) with a hash length of k = 29. The numbers of contigs and paired-end reads generated for each sample are detailed in Table 1.

**Taxonomic classification and BLAST-based comparative viromics**

Taxonomic classification was conducted on the nine samples using the web server Metavir (Roux et al. 2011). Owing to input restrictions of Metavir, the longest 300 000 paired reads (Table 1) were selected for analysis for each of the nine. Metavir assesses virome composition using the GAAStool (Angly et al. 2009). BLAST analyses by Metavir were performed using the RefSeq complete viral genomes protein sequences database (ftp://ftp.ncbi.nlm.nih.gov/refseq/release/viral/, accessed 27 August 2015); the release of the database used for comparison was that of 18 January 2014 and contained 173 122 protein sequences. The e-value cut off was 10e-5. 65% of the data generated by our study and processed through Metavir was determined to have originated from phage genomes. In addition to taxonomic classification, Metavir was used to compute rarefaction curves (Fig. S1 of the Supplementary material).
Comparative analysis of viromes based on contig assembly

Comparisons to publicly available freshwater viromes were performed in Metavir by performing tBLASTx comparisons using a subsample of the sequences taken from a virome, after data normalisation. Each of the nine viromes were separately assembled in a pair-wise fashion: reads generated from one sample were assembled to the reads generated from another sample, using MetaVelvet. This cross-sample assembly strategy can identify any similarities in the same taxa present within two samples.

ORF generation

The 20 longest contigs assembled by MetaVelvet for each of the nine samples were examined further. Open reading frames (ORFs) within each contig were predicted using GeneMarkS (Besemer et al. 2001). Each predicted ORF was then BLASTed against all records in the non-redundant (nr) protein sequences database, using the blastx algorithm from the NCBI web interface (results not shown). BLAST homologies having an e-value less than 10e-3 were considered putative hits.

Results

Overview of Lake Michigan viromes

Nine viromes were generated from samples collected from the nearshore waters of two Chicago area beaches, Montrose Beach and 57th Street Beach, and analysed to assess viral diversity. Samples were evaluated without reference to spatial or temporal difference, and were ultimately pooled in order to assess the range of diversity observed (i.e. as a composite group). Isolated DNA produced >15 000 000 raw reads (see Methods).

Rarefaction curves (Fig. S1) generated from the raw data suggested that total viral diversity was captured in only one of the nine viromes. As with most studies examining viral metagenomics, most of raw reads from Lake Michigan did not show any significant sequence similarity to current viral data: only 6.9% were found to produce a BLAST score of ≥50 against entries in the NCBI RefSeq viral protein database.

Comparison of Lake Michigan viromes

The Lake Michigan viromes were compared directly to 20 viromes generated from freshwater environments (publicly available) using the Metavir virome-comparison function which is based on the BLAST results for each sample (data normalisation is performed, by genome length, as part of the Metavir package (Roux et al. 2011)). The Lake Michigan samples clustered together, largely distinct from the other Metavir viromes (Fig. 1) but demonstrated very slight overlap with both freshwater and marine environments.

Additionally, the complete set of contigs generated for each of the nine Lake Michigan viromes was assembled in a pair-wise manner using the assembler MetaVelvet. This cross-sample assembly strategy can identify similarities in taxa present within two samples, for example, whether representative of viral species or coding sequences present within both samples. The contigs produced by this assembly were used to compare the nine viromes to one-another. These analyses suggested a considerable level of diversity within the composite sample, alongside a core element (greens and yellow, associated with red) (Fig. 2).

Taxonomic classification of viromes

The BLAST results (against RefSeq) for the composite virome, as expected, largely comprised members of the Caudovirales comprising the families Myoviridae, Siphoviridae and...
Podoviridae (Fig. 3). Although most of the hits were to these dsDNA phages, similarities to ssDNA phages as well as eukaryotic-infecting viruses were also observed (represented by the group ‘other’; Fig. 3). The Krona tool (Ondov et al. 2011), available as part of Metavir, was used to visualise matches between the composite Lake Michigan sequences and viral proteins in the RefSeq database (Fig. 4). The composite virome shows 95% of the protein hits are to coding regions within dsDNA (no RNA stage) viruses: 65% of which were for Caudovirales.

Viral-associated proteins

Via Metavir, BLAST (against RefSeq) returned matches to proteins belonging to a variety of viral species. Some phage species (1182) were represented throughout the composite virome (i.e. all nine samples). The majority (80%) of these hits pertaining to these phages were to several in particular. Numerous hits to the Planktothrix phage PaV-LD (NC_016564) were observed throughout the nine viromes and the composite; Fig. 4 shows that this phage was determined to comprise 4% of the entire virome. Further investigation showed that nearly all of these hits; however, (over 7000; Fig. 5a) were to the gene PaVLd_ORF033R. This protein is annotated as an ABC-transporter protein (Gao et al. 2012) in RefSeq. Fig. 5a also highlights two other genes specific to this phage, present in comparatively low frequency: a DNA helicase and a peptidase.

Further investigation uncovered that all nine separate viromes generated numerous hits to phages infectious to Burkholderia: 50% of the 72 annotated protein-coding genes within the 47 399 bp genome of the Burkholderia phage BcepB1A (NC_005886) (Summer et al. 2006) were found within the Lake Michigan virome. Most of the hits were from three of our original samples (Fig. 5b); however, numerous matches to other phages infecting Burkholderia cepacia were also present throughout the composite. High coverage for species of phage known to possess larger genomes was also observed, for example, the 252 401 bp genome of Prochlorococcus phage P-SSM2 (NC_006883) shown in Fig. 5c. Unlike Planktothrix phage PaV-LD, a broader distribution of the genomes of the Burkholderia and Prochlorococcus phages was present throughout the sample set.

BLAST matches to eukaryote-infecting viruses were minimal, the exception being the genomes of the giant amoeba-infecting viruses. Each individual Lake Michigan virome exhibited similarities to hundreds of different coding regions within the genomes of the Mimiviridae, Moumouviridae, Megaviridae and Pandoraviridae. For instance, over 16% of the annotated genes in Acanthamoeba polyphaga mimivirus (NC_014649) were found in the composite virome (Fig. 5d).

Comparing sequence composition of contigs to existing annotated metagenomic information can also give us insight into the potential function of viruses. The BLAST hits to phage species produced by the Metavir analyses for the nine LM viromes were aggregated by function based upon their annotated protein product (Table 2). The number of BLAST hits per annotated function varied considerably between the nine viromes (a range of 91–635). Functions with the greatest number of hits were further investigated. As is typical of similar datasets, most of the generated ORFs were assigned to hypothetical proteins.
Discussion

Environmental phages are one of the most abundant, yet understudied elements of microbial ecology. Our level of knowledge regarding the total diversity and genetic composition of environmental phages is minimal. Evidence suggests that communities of phages found in freshwaters are likely to represent a more complex cohort than that of their marine counterparts (Kimura et al. 2013; Baker et al. 2006), in congruence with the more heterogeneous nature of freshwater. Broadly speaking, in comparison to marine communities, freshwaters are understudied regardless, and so the scope of this expected greater diversity is unknown. Samples were collected from two Chicago beaches of Lake Michigan, and the dsDNA-based viral diversity was examined, with whole genome sequencing. Nine samples were processed and assessed both separately and as a composite.

The nearshore waters of a large lake differ in composition, physicochemically, from offshore waters (Yurista et al. 2012; Pilcher et al. 2015), so it follows that bacterial and phage community composition will vary from position to position and over time: on both large and small scales. However, capturing and successfully interrogating the changes in phage diversity temporally and spatially would require an exhaustive sampling and deep-sequencing effort. Instead, we examined our samples as a composite, in order to generate a ‘baseline’ representation of the phage community present in our samples from Lake Michigan.

For the purposes of assessment of diversity in the samples collected, the current study was designed to assess the presence of DNA viruses, in congruence with most of the other metaviromics studies performed in the environment. Furthermore, the web server we used to process our data (Metavir) has
previously been used with marine, brackish and freshwater sample sets (Roux et al. 2012). Owing to the nature of sample preparation (for example, multiple displacement amplification is known to produce a bias for ssDNA viruses (Kim and Bae 2011)), we theorised that most of BLAST hits returned by our sequencing effort would likely belong to dsDNA viruses. As anticipated, dsDNA phages were found to mostly comprise viruses in all datasets. Direct comparison of the Lake Michigan virome generated in the current study with similar research presents inherent challenges, owing to the innate levels of heterogeneity, which each habitat is likely to exhibit, as well as differences in sample preparation. Despite these challenges, we were able to compare the LM virome with other freshwater viromes using BLAST-based analyses using Metavir. The comparison revealed that the dataset generated from the current study appears to be divergent from previous studies (Fig. 1). In light of known vacillations in community composition in freshwaters this is to be expected, and suggests that disparity between freshwater samples from different locations, as well as between freshwater and marine samples, is also likely. This said, rarefaction curves produced as part of Metavir’s analysis (Fig. S1) suggest that out of the nine viromes produced, only one captured the entire representative diversity. Metavir restricts inputs to 300 000 reads, and Roux et al. (2012) described similar findings for two freshwater lake samples: viral diversity was not entirely captured according to rarefaction curves. In addition, the potential for bias in sampling, sample preparation and sequencing is considerable (Schoenfeld et al. 2010), and therefore our low diversity virome may have fallen subject to these issues. However, quality of this sample was assessed before sequencing (as part of standard QC protocols before Illumina sequencing) and was judged to be congruent with the others. Owing to the general paucity of knowledge in this area, it is difficult to interpret why one sample may have been so markedly different to others taken from the same environment, albeit at different times. However, in freshwaters such differences in diversity, on a temporal and spatial basis, are possible,

![Graphs](image-url)
if not expected. These discrepancies are countered to some extent by our decision to analyse our samples as a composite. Overall, however, these results suggest a considerable reservoir of unexplored viral ecology.

Furthermore, the BLAST-based comparisons highlighted only a small fraction of the genomic content within the Lake Michigan viromes: this is largely unavoidable owing to the paucity of viral data in RefSeq, for example, which means most of the hits are of unknown origin. Regardless of the environmental niche investigated, most of the sequence reads generated from projects examining metaviromes in particular, demonstrate infrequent homology to previously annotated function or taxonomy (Yin and Fischer 2008). This is owing to the low frequency of entire phage genomes in the RefSeq database and, in turn, an even lower incidence of entirely and accurately annotated phage genomes. This remains a key confounding issue with regard to examining phage diversity in the environment and further highlights the requirement for continued culture-based investigations based on physical isolation of phages.

Taxonomically, most of the viruses in our virome belong to the order Caudovirales. Caudovirales, dsDNA viruses comprising the families Myoviridae, Podoviridae and Siphoviridae, and represent the low-hanging fruit of the phage world. They are easy to isolate from environmental samples and maintain in the laboratory, and therefore they are easy to find in metaviromic datasets. For this reason they have substantial presence in the RefSeq database and the International Committee on Taxonomy of Viruses database (http:// ICTVonline.org/virustaxonomy.asp, accessed 8 August 2015). We identified several phage taxa of interest to examine further. Usually, phages are named for their bacterial hosts; however, this can represent a challenge when classifying diversity representatively. With a BLAST e-value cut-off of 1e-5, all nine viromes showed numerous hits to the *Planktothrix* phage PaV-LD (NC_016564). Further investigation showed that nearly all of these matches (Fig. 5a) were to the gene PaVLD ORF033R, which is annotated as an ABC-transporter protein (Gao et al. 2012). The presumptive abundance of this gene in our samples is extremely unlikely to represent a single ‘species’ of phage only, and is more likely owing to the high presence of non-species or virus-specific ABC-transporter proteins (ubiquitous throughout all extant prokaryotic and eukaryotic phyla alike) in the sample. Owing to previous identification of a high incidence of *Burkholderia* in samples taken from the lake (data not shown), *Burkholderia* phage BeepBIA1 was highlighted for further examination.

In addition to hits to the ABC-transporter coding region (YP_004957 306) for the *Planktothrix* phage PaV-LD genome (Fig. 5a), hits to other phage species, including *Bacillus* phage SPbeta and *Bacillus* phage G, were also identified. Further investigation into these hits revealed that sequence similarity between the Lake Michigan virome contigs and the RefSeq database was localised to the ABCB_MRP_Like domain (CDD ID: cd03228). Expanding our analysis to all hits, not just to phage species, there were a significant number that were analogous to coding regions annotated as ABC transporters within the genomes of amoeba-infecting viruses. As the Metavir analysis is limited to the RefSeq annotated viral-genome collection, we checked to see if other viral metagenomic studies had also observed this function in their sample. Selecting the protein sequence for this domain from the *Bacillus* phage-SPbeta genome, BLASTp searches were conducted against the non-redundant protein-sequences database (limiting the search to sequences taxonomically classified as viruses) as well as the metagenomics database (env_nr). In the case of the former, numerous hits to sequences of the taxon *Phycodnaviridae* were identified. This suggests that there is incongruity between the initial classification of *Bacillus* phage and the second search, which employed more stringent parameters. Similarly, the BLASTp search to the env_nr database identified statistically significant hits (E-value = 8e-37) to sequences generated from global-ocean sampling studies. Although *Prochlorococcus* is a marine cyanobacterium, it has been determined previously that P-SSM2, a myovirus, is likely to contain several core genes, both belonging to T4-like groups and those which are cyanobacterial in origin (Sullivan et al. 2005). *Prochlorococcus* is a dominant genus of cyanobacteria in the marine environment, but a vast catalogue of freshwater cyanobacteria will be indigenous and present in association with freshwater cyanophages. Cyanobacteria, and by definition, cyanophages, contain highly mobile genetic elements and therefore both groups, both marine and freshwater may possess the genes identified in our sample set (Ivanikova et al. 2007). This explains why genes that are, ostensibly, marine in origin, may have validity in a freshwater dataset. Our dataset contained hits against P-SSM2 (Fig. 5c): >50% of its annotated coding regions. This phage has a significantly larger genome than that of the *Burkholderia* phage, that of the mimivirus is larger still: based on the distribution of hits to these genomes, we hypothesise that relatives of these phages are actually present in the lake, as opposed to PaV-LD. Hits to the *Planktothrix* phage, in this case, likely reveal only the presence of ABC-transporter genes. These analyses reconfirm that in a dataset as complex as an aquatic virome, where most of BLAST hits are unknown, the presence of specific genes is what may give insight into community composition, as opposed to the species assigned to that gene specifically.

Those ORFs that returned more descriptive hits (as opposed to hypothetical, and including DNA helicases, DNA integrases, DNA polymerases, portal proteins, structural proteins and the large subunit of terminases), all belonged to proteins most prevalent in single-annotated phage genomes (Kristensen et al. 2013) as opposed to those mined from metagenomic data. Integrases are highly ubiquitous, and can be shared by phages, whereas previous bioinformatic analysis (Kristensen et al. 2013), found that many other functions are phage-specific (i.e. pertaining to phages, and not bacteria), including the large subunit of terminases, portal proteins, and structural proteins. Other phage-specific ORFs, which are frequently found in annotated viral genomes, such as those belonging to the cro or cl-repressor system (important in lytic and lysogenic development (Folkmanis et al. 1977)) and homing HNH endonucleases (important in lateral transfer (Friedrich et al. 2007)), were also identified in the Metavir hits, albeit far fewer in number (1315 and 3853 respectively). This suggests the presence of lysogenic or potentially lysogenic phages in the Lake Michigan viromes. Phages are able to propagate only by the successful infection of a host. There are two major mechanisms for infection: (1) lysis, in which viral progeny are released from the host cell as a
fatal event, and (2) lysogeny, during which genomic materials from a phage are integrated with that of a host cell. Over time, this may result in genotypic and phenotypic changes to the host cell, or eventual cell death. Therefore, most genomic information available for phages pertains to lytic phages. Of course, phages may also change lifestyle over time, a factor which may only be determined in culture (Watkins et al. 2014). Clearly, the two lifestyles may confound metagenomic survey-based examinations of total viral diversity, as sample preparation methods are largely designed to capture ‘free’ viruses. In addition, one phage may infect more than one host, and in some cases, this broad host-range may extend across bacterial genera. Phages previously examined and labelled as infectious to one genus, may ultimately have a different preferred host. This phenomenon, may, similarly, be changeable according to resource limitation or abundance, physical factors, or the proliferation of a specific taxon of bacteria over another. These factors, among others, should be taken into consideration while examining metagenomics datasets.

The nearshore waters of Lake Michigan represent a viral population, which is diverse from other publicly available datasets and which demonstrates phylogenetic divergence from previously studied viral communities. Although the viromes analysed in the current study are composed largely of groups of dsDNA phages commonly present in other such datasets (as expected), they are a potentially rich source of further information regarding the genetic heterogeneity of freshwater systems in general. The data obtained from the current study, from a composite sampling regime, has been used to assess the diversity of Lake Michigan by examining the presence of commonly seen viral-associated genes and proteins. Not only will this work enable continued investigations into the viral ecology of freshwater systems, including studies designed to infer impact and interaction of phage communities on and with their hosts, but it has laid a foundation for the mammot task of examining the entire community of Lake Michigan.

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References


Supplementary material

Assessment of a metaviromic dataset generated from nearshore Lake Michigan

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\begin{figure}
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\includegraphics[width=\textwidth]{figs/figS1.png}
\caption{Rarefaction curves for the Lake Michigan viromes, as generated in Metavir.}
\end{figure}