Census of the Viral Metagenome within an Activated Sludge Microbial Assemblage


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Received 16 October 2009/Accepted 3 February 2010

The viral metagenome within an activated sludge microbial assemblage was sampled using culture-dependent and culture-independent methods and compared to the diversity of activated sludge bacterial taxa. A total of 70 unique cultured bacterial isolates, 24 cultured bacteriophages, 829 bacterial metagenomic clones of 16S rRNA genes, and 1,161 viral metagenomic clones were subjected to a phylogenetic analysis.

Bacteriophages play an active role in the ecology of natural environments, influencing prokaryotic population dynamics (5, 15) and mediating lateral gene transfer between diverse bacterial species, for example. Activated sludge (AS) microbial assemblages in wastewater treatment plants have been shown to harbor great numbers of viruses with a wide range of genome sizes (7, 9, 10, 16). Historically, the focus of wastewater viral studies has been on specific host-virus interactions, the application of phages as tools in microbial source tracking, or the use of phages to improve the efficiency of the wastewater treatment process (e.g., foam and pathogen control) (2, 4, 8, 12, 17). Despite the interest in the wastewater viral community, a census of the activated sludge total viral community has not, to our knowledge, been investigated using both culture-based and metagenomic approaches.

Taxonomic assignment of viral metagenomic clone sequences. Samples were collected from the AS aeration basin at the H. C. Morgan Water Pollution Control Facility in Auburn, AL, treated with 10% beef extract buffer (to separate viruses from aggregates), precipitated with polyethylene glycol, and subjected to cesium chloride gradient centrifugation. Viral DNA was then extracted by benzonase, proteinase K, and sodium dodecyl sulfate treatments and purified by phenol-chloroform extraction and isopropanol precipitation. An AS viral metagenomic library was constructed at the Lucigen Corporation (Middleton, WI) using a linker-amplified shotgun library approach (3, 18), and 1,161 cloned insert DNA sequences were determined using a single-vector primer. Trimmmed viral insert sequences were classified as known (E value < 0.001; n = 694), unknown (n = 97), or novel (n = 370) as described previously (18). The known sequences were assigned to their taxonomic affiliations by BLASTx comparisons to the GenBank nonredundant nucleotide (nr/nt) database and selection of the top BLASTx hits for phylogenetic affiliation (Fig. 1).

The dominance of viral metagenomic sequences with significant homology to bacterial DNA (nearly 60% of the known sequences) is likely the consequence of the high frequency of prophage sequences within bacterial genomes deposited in GenBank databases (1). Considerable care was taken to prevent inclusion of prokaryotic (or eukaryotic) chromosomal DNA within the viral metagenomic library (i.e., CsCl purification and nuclease treatment were performed), and the high percentage of viral metagenomic sequences with homology to bacterial genomes has been observed in other environments (3). Of the 191 known sequences with homology to viral genomes, 95% were homologous to bacteriophage sequences and were within the viral families Myoviridae (40.3%), Siphoviridae (31.9%), or Podoviridae (25.6%) or considered unclassified phages (2.2%). The remaining sequences were classified as mobile genetic elements (9%), eukaryotic (8%), or archaeal (<1%) in nature. The taxonomic heterogeneity of sequences obtained in the viral metagenomic library underscores the diversity of the AS system, which receives an influx of microbiota from both human and environmental sources. In addition, this survey revealed a high percentage of novel predicted gene products (37%), illustrating the relative scarcity of wastewater viral DNA sequences in the GenBank nr/nt or environmental (env) databases.

Comparison of bacterial and viral phylogenetic affiliations. Because bacterial community structure is expected to be the primary determinant of bacteriophage diversity within an environment, the bacterial taxa from AS were surveyed using culture-dependent and -independent analyses. A survey of AS 16S rRNA genes was conducted with AS bacterial cultures (70 unique bacterial cultures) isolated on LB or a synthetic wastewater medium (14) using aerobic and anaerobic growth conditions. In addition, a bacterial culture-independent survey (n = 829) was conducted using as the template AS metagenomic DNA extracted from the same sample used for viral studies and PCR amplified using universal bacterial primers 27F (5’-AGAGTTTGATCMTGGCTCAG-3’) and 1492R (5’-

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† Published ahead of print on 12 February 2010.
GYTACCTTGTTACGACTT-3' (11). The 16S rRNA gene amplicons from the metagenomic DNA template were cloned into the TOPO-TA pCR2.1 vector (Invitrogen, Carlsbad, CA) and transformed into *Escherichia coli* cells. Each 16S rRNA gene clone sequence was trimmed for quality, compared to the GenBank nr/nt database using the BLASTn search algorithm, and classified according to bacterial division affiliation of the top BLASTn hit (Fig. 2).

The culture-independent 16S rRNA gene clone library revealed more than a 3-fold increase in the number of represented bacterial divisions (*n* = 10) compared to that observed from the culture collection (*n* = 3), and a significant shift in relative dominance of each phylum was observed between the two surveys. Although *Proteobacteria* taxa dominated the culture-dependent survey (58% of analyzed sequences), the percentage of sequences representing this division in the culture-independent survey dropped to 25%. Similarly, *Firmicutes* representation was almost 10-fold lower in the culture-independent survey than in the culture-dependent study (4% versus 36%, respectively). In contrast, the *Bacteroidetes* taxa, which were only minimally represented in the culture-dependent survey, comprised more than half (51%) of the bacterial taxa identified in the 16S rRNA clone library. The minimal overlap between data obtained from these culture-dependent and -independent analyses reflects the strong cultivation bias that is common to virtually all environmental studies.

Viral enrichments performed on all of the bacterial isolates identified 24 bacterial hosts from which phages could be isolated. Seventy-five percent of the phages were isolated from *Firmicutes* (8%) or *Actinobacteria* (17%) hosts. In the culture-independent viral metagenomic survey, the 389 sequences from the viral metagenomic library that were most similar to bacterial genomes were included in this taxonomic classification (i.e., viral, culture independent [top bacterial BLASTx hit]), as were the predicted hosts of the 187 viral sequences with significant homology to bacteriophage DNA (i.e., viral, culture independent [top phage BLASTx hit]) (Fig. 2). Although the two viral surveys demonstrate similar phylogenetic distributions, there are some striking differences observed between the analyses of the viral metagenomic library and the bacterial surveys. For example, only two sequences from the viral metagenomic library (0.5% of bacterial BLASTx hits and 3.1% of phage BLASTx hits) were highly similar to *Bacteroidetes* DNA despite the high relative abundance of *Bacteroidetes* taxa in the bacterial culture-independent survey (Fig. 2). *Proteobacteria* was the dominant phylum in every survey except for the bacterial culture-independent study, in which the *Bacteroidetes* division was the most highly represented (51%). In addition, four of the bacterial phyla predicted to be present within the viral metagenomic library (*Deinococcus, Cyanobacteria, Chloroflexi, and Spirochetes*) were absent from both of the bacterial phylogenetic surveys. Both of the Gram-positive phyla (*Actinobacteria* and *Firmicutes*) had slightly higher representation within the viral metagenomic survey than in the bacterial culture-independent survey. The phylogenetic distributions of the phage metagenomic sequences with the top BLASTx hits as bacterial versus the top BLASTx hits as viral were similar in every case, with the *Proteobacteria, Firmicutes*, and *Actinobacteria* taxa dominating the viral met-
agenomic survey. This study highlights the relative paucity of information within the GenBank databases concerning bacteriophages infecting Bacteroidetes taxa; presumably a very high percentage of the phage metagenomic sequences annotated as unknown or novel could be affiliated with the Bacteroidetes phylum.

Comparison of abundant viral metagenomic loci and cultured phage DNA. Contig assembly of viral metagenomic sequences was performed using assembly criteria of 80% identity with at least a 20-bp overlap in order to assemble sequences from the most abundant viral types (3). The resulting contigs were compared to the nr/nt and env GenBank databases by BLASTx, and PCR primers were designed to target contigs with significant homology to bacterial hosts in the AS culture collection. Purified DNA from cultured phages was used as the template in a PCR with the metagenomic contig-specific primers, yielding three amplicons from the cultured phages. Predicted phylogenetic relationships between these amplicons, the metagenomic contigs, and their nearest neighbors in the nr/nt database were inferred using the maximum parsimony method. The resulting cladogram (>50% bootstrap support) indicated that every metagenomic contig was more closely related to a cultured phage amplicon than to any other sequence in GenBank databases (data not shown).

Metabolic profiling of viral metagenomic DNA sequences. Sequences from the viral metagenomic library clones were analyzed by the MetaGene annotator, which identified 1,815 open reading frames (ORFs) within the library. The ORF sequences were then compared to those found in the SEED platform (http://www.theseed.org) and organized into predicted functional categories based on the subsystems approach described by Overbeek et al. (13).

Metabolic profiles were predicted for the viral metagenomic sequences using the SEED platform (Fig. 3). The most dom-
inant subsystems were those involved with macromolecule metabolism, although ORFs encoding virulence factors and pigment production collectively comprised approximately 20% of the sequences from the viral library. Several common subsystems evident in the AS viral metagenome (e.g., carbohydrate and amino acid metabolism) are highly represented in viral metagenomic surveys in other environments (6). Considering that this was not an exhaustive survey of AS viral metagenomes, no conclusions regarding changes in the relative dominance of each subsystem may be made, but overall the distribution of metabolic systems in an AS viral metagenome is similar to that of viral metagenomes from other environments (6). This is the first study to explore the functional genetic diversity present within an activated sludge viral community. Future studies will further this work by employing next-generation sequencing methods and expanding upon the viral sequence repository available from multiple natural environments.

Nucleotide sequence accession numbers. All 16S rRNA gene sequences from the cultured isolates and bacterial 16s rRNA clones were submitted to GenBank under accession numbers GU002706 to GU003868. The viral metagenomic DNA sequences were registered as part of the Wastewater Viral Metagenome project at GenBank (Trace Archives) and are under TI reference numbers 225120377 to 2251204802.

REFERENCES


