Genetic Diversity of Bacteriophage Communities in Napahai Wetland

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Abstract—Many results have showed that the great diversity of bacteriophages in various aquatic environments, including marine water, freshwater and paddy water, but few studies was about the genetic diversity of bacteriophages in the wetland. To better understand the genetic diversity of bacteriophages communities of Napahai wetland, we selected viral capsid protein gene (g23) as a gene marker to reveal genetic diversity of bacteriophage in Napahai wetland. The g23 gene fragment of bacteriophage was amplified with the primers MZA1Bis and MIZA6. In this study, 21 different clones were found. Phylogenetic analysis demonstrated that some g23 genotypes were closely related to the T4-like phages, the others clustered two groups. Phylogenetic classification and genetic diversity of bacteriophages in the wetland. To better understand the genetic diversity of bacteriophage communities. In this study, fragments in g23 gene sequences were found. Phylogenetic analysis demonstrated that some g23 genotypes were closely related to the T4-like phages, the others clustered two groups. So the unknown sequences may represent the new particular g23 genotypes.

Keywords—T4-type phage, Napahai wetland; g23 gene; Genetic diversity.

I. INTRODUCTION

Viruses are general in various environments and have a great abundance in various environments. Viruses play an important role in regulating the structure and diversity of microbial populations. Because of great amount and biodiversity of viruses, they are considered as the largest genomic reservoir and among them the vast majority are bacteriophages in the natural environments[1].

The T4-type of phage as an important member of the Myoviridae family have caused widespread concern in the past few year, particularly from ecological scientists[2]. With the development of viral genomics, genetic diversity analysis is considered as a powerful improver on phylogenetic classification and genetic diversity of viruses in various environments[1]. The g23 as a capsid protein gene of T4-type phages will be the most frequently used marker gene for assessing gene-diversity of T4-type bacteriophage[1]. And in the past few years, most molecular reachers on phages in nature have mainly focused on the viruses of marine[3].

Wetlands are a natural genetic library, have characteristics of both land and water and play a number of roles in the environment. The Napahai wetland which located in Shangri-La is a unique seasonal plateau wetland with low latitude and high altitude in Yunnan, China. It is an isolated ecosystem that own distinct rainy and dry seasons. There are a large number of bacteriophages which are one of the most important factors to balance the wetland ecosystem, but the research on phage was still rare in Napahai wetland. In this study, the viral capsid protein gene (g23) was selected as a gene marker to reveal genetic diversity on phage communities of natural environments.

II. MATERIALS AND METHODS

A. Napahai wetland water sampling and ultrafiltration

The sample was collected from the Napahai wetland of Shangri-La (E99°37’22",N27°53’32",3266m, pH6.6) in 2013. Water samples were kept in dark at 4°C.

Centrifuged water samples at 8000g for 30 min at 4°C to remove soil particles, plankton etc. Then filtrated the water samples with a 0.45μm and 0.22μm cellulose membrane to remove other bacteria. Concentrates of virus were stored in the dark at 4°C[4].

B. DNA extraction and PCR amplification

The treated water samples were frozen in liquid nitrogen or -80°C frozen 8 h, nozzle after laminating. The lyophilized overnight freeze-dried into powder, DNA was extracted with the OMEGA virus genome extraction kit.

The viral capsid protein gene g23, was amplified using the primers MZA1Bis and MIZA6[5]. Fifty microliter of PCR mixture contained 1μL of forward and reverse primers (10pmol each), 1~2 μL of DNA template, 5μL of dNTP(2.5mM each), 0.5 μL of Ex-Taqpolymerase, and 5 μL of Ex-Taqbuffer and was filled up to the required volume (36.7–37.7 μL) with MillIQ water. PCR products were performed with the following cycle parameters: denaturation at 94°C for 4 min, 35 cycles of denaturation at 94°C for 45 s, annealing at 45°C for 45 s, ramping at 0.5°Cs-1, and extension at 72°C for 45 s, a final extension at 72°C for 10 min.

C. Sequence analysis

The g23 positive clones were analyst by using the BLAST search tool at NCBI (http://www.ncbi.nlm.nih.gov/). A neighbor-joining tree was founded by Molecular Evolutionary Genetic Analysis software (MEGA 3.0) [6] with 1000-fold bootstrap support.

III. RESULTS

A. DNA extraction

The treated water samples were extracted with the PCI solution (phenol:chloroform:isoamylalcohol=25:24:1,v/v) and once with CIA solution (chloroform:isoamylalcohol=24:1,v/v), and DNA was contaminated by the other proteins. So the OMEGA virus genome extraction kit was used to extract DNA.
Figure 1. Concentrated phage metagenome of NPHW-2 in May 2013. M: DNA markers; 1: phage metagenome (extracted with phenol/chloroform method); 2: phage metagenome (extracted with OMEGA virus genome extraction kit).

B. PCR amplification

The primer was designed to detect the T4-type phage in Napahai wetland, then the viral capsid protein gene, g23, was amplified with the primers M2IAIBis and MIZA6. About 500-bp PCR products were obtained from all samples, and the best annealing temperature was 50°C (Fig. 2).

Figure 2. Temperature gradient amplification of concentrated phage of Napahai wetland with primers M2IAIBis/MIZA6

C. Phylogenetic diversity of T4-type phages isolates

In this study, as shown in Fig. 3, many g23 clones formed several clusters separate from those of floodwater origins. This finding clearly showed that many g23 genes in this study differed from those obtained from marine and freshwater environments [7], which was consistent with our assumptions that g23 sequences in wetland were unique and different from those of marine and freshwater origins. The tree demonstrated that the homology similarity among T4-type phages and clone12, clone17, and clone22 were 81%, 85%, and 86%, respectively. About 11 g23 fragments from Napahai wetland formed clusters which were different from those of marine and freshwater environments. And the remainders were similar with the Uncultured phage clone from Antarctic Lake and Kotokel Lake.

Figure 3. Phylogenetic relationships among the clones amplification of concentrated phage of Napahai wetland with primers M2IAIBis/MIZA6

IV. DISCUSSION

Viruses are extremely abundant in lakes and oceans, yet little is known about the composition of these viral communities. In particular, the distribution of genotypes is virtually unexplored for bacteriophages [8]. This study investigated the genetic diversity of bacteriophage communities in Napahai wetland. In this study, sequence analyses of the viral g23 gene have been used to analyze the diversity of natural assemblages of wetland bacteriophages. First File[e] [8] gained gene fragments ranging from 380 to 600 bp from marine samples. In addition, Jia et al. [2] and Fujii et al. obtained PCR products in a range from 415 to 614 bp in the Anjo paddy field and from 422 to 644 bp in the Omagari paddy field [7]. Guanghua Wang et al. [7] obtained PCR products ranging from 350 to 599 bp. The length was about 500 bp in this study. These findings indicate that g23 genes are widely variable in length in marine and soil environments, probably reflecting the great diversity of phage communities in those environments [7].

In this paper, some of the clones obtained in this study formed several small clusters with clones that were ungrouped in previous studies [7] by HUANG Hui-Zhen et al. [9], there were also some had some homology with the previous studies by Tatyana V et al. [10] and Alberto López-Bueno et al. [11]. We speculated that Napahai wetland were similar with Lake Kotokel and Antarctic Lake, which all located in high latitudes and they were
had the similar low temperature environments. Finally, there were about half of clones had no homology with the sequences obtained in the database. They are the unique phages communities in Napahai wetland.

In conclusion, this study indicated that rich diversity of bacteriophages existed in Napahai wetland. And there were many unique phages in Napahai wetland which were different from those in marine, freshwater environments. Further studies on the covariation of the genetic diversity of virus and host populations with changing environmental variables would provide new insights into the ecological roles of wetland viruses.

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REFERENCES