Comparison of extraction methods of total microbial DNA from freshwater

P. Li, S.F. Yang, B.B. Lv, K. Zhao, M.F. Lin, S. Zhou, X. Song and X.M. Tang

1Biotech Research Institute of Shanghai Academy of Agricultural Sciences, Shanghai, China
2Shanghai Key Laboratory of Agricultural Genetics and Breeding, Shanghai, China
3Institute of ECO-Environment and Plant Protection, Shanghai Academy of Agricultural Sciences, Shanghai, China

Corresponding author: X.M. Tang
E-mail: sunsite@126.com

Received January 3, 2014
Accepted April 3, 2014
Published January 30, 2015
DOI http://dx.doi.org/10.4238/2015.January.30.16

ABSTRACT. The demand for molecular analysis of aquatic microbial communities in freshwater has highlighted the need for efficient methods of DNA extraction. The centrifugation method and filtration-membrane method are 2 widely used methods for extracting DNA. The objective of this study was to compare the extraction efficiency of 3 methods, including the centrifugation method, filtration-membrane method, and modified filtration-membrane method, by evaluating the quantity and purity of DNA extracts obtained from water. DNA extraction was analyzed by agarose gel electrophoresis, ultraviolet-spectroscopy, restriction enzyme digestion, and polymerase chain reaction. The results showed that the modified filtration-membrane method was the most efficient for extracting microbial DNA from freshwater with high integrity and purity and is suitable for molecular applications.

Key words: Extraction efficiency; Freshwater; Microbial diversity; Polymerase chain reaction; Total DNA
INTRODUCTION

As primary components of material and energy recycling pathways, microbes are essential to all life on Earth, playing a particularly important role in freshwater environments. Analysis of microbial community structures, the characteristics of microbial distribution, and the function in aquatic ecosystems is important for ecological studies (Wu et al., 2010).

The conventional method for analyzing water microbial populations involves isolating and traditional culturing of microbes. However, this method is limited because few microbes can be effectively cultured in water. It was previously estimated that approximately 0.25 and 0.001-0.1% microbes can be cultivated in freshwater and seawater, respectively (Amann et al., 1995). Because of the inherent limitations of culture-based methods, microbial ecologists have begun to develop culture-independent methods of community analysis. The use of culture independent methods circumvents the disadvantages of microbial isolation and culture, and the direct use of molecular biology methods increases the probability of identifying new microbes and analyzing the diversity present in the microbial community (Handelsman, 2004).

Therefore, extracting total DNA from water-dwelling microbes is very important. Currently, there are 2 types of methods for extracting total DNA from water microbes. First, microbe enrichment can be achieved using high-speed centrifugation (Zhang et al., 2012). Second, microbes can be absorbed and aggregated using a filtration membrane (Ye et al., 2009).

Before extracting DNA from freshwater organisms, a high concentration of microbes is required. The high-speed centrifugation method is the simplest and most cost-efficient method. However, the purity of extracted DNA is highest using the filtration membrane method. The compositions of freshwater environments are very complex and may contain chemical components, particularly humic acid, which inhibits the activity of enzymes such as restriction endonucleases and DNA polymerase (Mangiapan et al., 1996; Bürgmann et al., 2001; Dong et al., 2006). Further, the physico-chemical properties of humic acid and nucleic acids are similar, making it difficult to extract total DNA.

In this study, we compared 3 methods for extracting total microbial DNA, including the centrifugation method (Zhang et al., 2012), the filtration-membrane method (Ye et al., 2009), and the modified filtration-membrane method (developed in our laboratory), to determine the most suitable method for evaluating microbial community dynamics and the quantity and purity of DNA.

MATERIAL AND METHODS

Collection of freshwater samples of rivers

Freshwater samples were collected from the Huangpu River (31° 14'N, 121° 29'E) in Shanghai on July 9, 2012. This river is used for irrigating rice and drinking water. The Huangpu River is a tributary of the largest river in China, the Yangtze River. The temperature, pH, and concentration of dissolved oxygen in the water column at the time of sampling were 18.2°C, 7.7, and 11.8 mg/L, respectively. Freshwater samples were collected from 0.5 m below the surface using a Ruttner Standard Water Sampler (KC-Denmark A/S Company, Silkeborg, Denmark) and immediately transferred to a sterile bottle. All samples were transferred to the laboratory and stored at 4°C.
Total microbial DNA extraction methods of freshwater organisms

Centrifugation method with slight modification

This method was described previously by Zhang et al. (2012) and used with some modifications. The 50-mL water sample was centrifuged at 5000 g for 20 min at room temperature. The precipitate was transferred to a 2-mL centrifuge tube and 1.5 mL 1% hexadecyltrimethylammonium bromide (CTAB) buffer was added and mixed thoroughly. The suspension was incubated at 65°C for 20 min. An equal volume of chloroform/isoamyl alcohol (24:1) was added and the sample was centrifuged for 10 min at 4°C at 13,000 g. Next, 2/3 volume isopropyl alcohol was added and the mixture was cooled at -20°C for 15 min. This sample was centrifuged for 10 min at 4°C at 13,000 g and the supernatant was discarded. Next, 700 µL 70% ethanol was added, followed by centrifugation for 10 min at 4°C at 13,000 g to precipitate the DNA. The supernatant was discarded. After this step, 500 µL 70% cold ethanol was added and the sample was centrifuged for 5 min at 4°C at 13,000 g and the precipitate was collected. The supernatant was discarded and the pellet was air-dried at 65°C. Sterile ddH₂O was added and the DNA solution was stored at -20°C.

Filtration-membrane method with slight modification

This method was described previously by Ye et al. (2009). The 50-mL water sample was filtered through a 0.22-µm pore filter (diameter, 45 mm; Millipore, New Bedford, MA, USA). The membrane was cut into 1-2 mm of the debris under sterile conditions, and debris was removed into a 2-mL microcentrifuge tube. Next, 1.5 mL STET buffer (8% sucrose, 50 mM Tris, pH 8.0, 50 mM EDTA, 0.1% Tween-20) was added to fill the tube and the sample was centrifuged for 5 min at 4°C at 10,000 g. The supernatant was removed and 1 mL STET buffer was added, followed by centrifugation for 2 min at 4°C at 10,000 g. The precipitate was collected. Next, 200 mL STET buffer and 4 µL 50 mg/mL lysozyme were added and the sample was mixed for 5 min at room temperature. Proteinase K was added to a 100-µg/mL concentration and the sample was mixed and incubated for 1 h at 37°C. After incubation, 0.5 M NaCl and 25 µL 5% CTAB were added and the sample was incubated for 10 min at 65°C.

To separate the DNA from the mixture, an equal volume of saturated phenol was added and the sample was centrifuged for 5 min at 4°C at 12,000 g; the supernatant was transferred to a 1.5-mL microcentrifuge tube. Next, an equal volume of phenol/chloroform/isoamyl alcohol was added (25:24:1, v/v) and the tube was centrifuged for 5 min at 4°C at 12,000 g; the supernatant was transferred to a 1.5-mL microcentrifuge tube. To the supernatant, an equal volume of chloroform was added, and then the sample was mixed and centrifuged for 5 min at 4°C at 12,000 g; the supernatant was transferred to a 1.5-mL microcentrifuge tube.

To this supernatant, 1/10 volume 2 M sodium acetate and 2 volumes cold ethanol were added, the sample was mixed well, and then cooled at -20°C for 1 h. After 1 h, the sample was centrifuged for 10 min at 4°C at 13,000 g and the supernatant was discarded. Next, 500 µL 70% cold ethanol was added and the sample was centrifuged for 5 min at 4°C at 13,000 g and the precipitate was collected. The pellet was air-dried at 65°C; sterile ddH₂O was added and the DNA solution was stored at -20°C.
Modified filtration-membrane method

First, a 50-mL freshwater sample was filtered through a 0.22-μm hydrophobic polyvinylidene fluoride membrane (Sangon Biotech Co., Ltd., Shanghai, China). The membrane was cut into 1-2-mm sections under sterile conditions, and debris was removed to a 2-mL microcentrifuge tube. Next, 1.5 mL STET buffer (8% sucrose, 50 mM Tris, pH 8.0, 50 mM EDTA, 0.1% Tween-20) was added to fill the tube.

Second, 0.8 g sterilized zirconium beads (1.0-1.2 nm, Saint-Gobain Research Co., Ltd., Shanghai, China) was added and the sample was vortexed for 10 min by placing horizontally on a reciprocal platform shaker at high speed (approximately 100 oscillations/min) at room temperature. Next, 200 μL STET buffer and 4 μL lysozyme (50 mg/mL) were added and the sample was mixed for 5 min at room temperature. To the suspension, 6 μL proteinase K (20 mg/mL) and 30 μL SDS (10%) were added and the sample was incubated for 10 min at 37°C. After incubation, 1/6 volume 5 M NaCl and 1/9 volume CTAB (final concentration of 5%) were added, followed by incubation for 10 min at 65°C. After this step, 200 μL phenol/chloroform/isoamyl alcohol (25:24:1, v/v/v) was added, the sample was vortexed, and the tube was centrifuged for 5 min at 4°C at 15,000 g. The upper (aqueous) phase was transferred to a 2.0-mL microcentrifuge tube.

To this layer, 0.7 volumes isopropanol was added to precipitate the nucleic acids and the tube was incubated at room temperature for 10 min. The sample was centrifuged for 15 min at 12,000 g at room temperature and the isopropanol was carefully removed. The remaining pellet was resuspended in 50 μL TE buffer (10 mM Tris-HCl, pH 8.0), 5 volumes GES reagent (5 M guanidium thiocyanate, 100 mM EDTA, and 0.5% sarkosyl) was added, and the sample was vortexed briefly. Transfer the mixture to an adsorption column (EZ-500, Sangon Biotech Co., Ltd.). Next, 200 μL wash buffer I (5.5 M guanidium thiocyanate, 0.5 M Tris-HCl, pH 8.0) was added and the sample was centrifuged for 1 min at 12,000 g. Residual liquid in the adsorption column was removed and 200 μL washing buffer II (0.4 M NaCl, 50 mM Tris-HCl, pH 8.0, 50 mM MgSO4) was added; the sample was centrifuged for 1 min at 12,000 g and the residual liquid in the adsorption column was removed. To the center of the adsorption column, 50 μL sterile ddH2O was slowly added and the column was incubated for 15 min at 37°C, followed by centrifugation for 1 min at 12,000 g. The DNA solution was stored at -20°C.

Assessment of quality and quantification of DNA

Genomic DNA was analyzed by 2% agarose gel electrophoresis. The results were photographed after ethidium bromide staining. DNA concentration was determined using a NanoDrop® ND-1000 (NanoDrop, Wilmington, DE, USA) and DNA yield was calculated. To evaluate the purity of the extracted DNA, optical density values were measured at wavelengths of 230, 260, and 280 nm (A230, A260, and A280, respectively) and the A260/280 and A260/230 ratios were calculated. Absorption values of the DNA solutions were measured by full-spectrum wavelength scanning over a range of 220-320 nm. The average ratio of A260/280 was calculated for each set of triplicate samples and was used to estimate the purity of extracted nucleic acids: samples with a mean A260/280 of 1.8-2.0 were considered to be free of contamination, those with A260/280 < 1.8 were considered to contain protein or other contaminants, and those with A260/280 > 2.0 were considered to be due to the presence of RNA.
Restriction enzyme digestion analysis

λDNA (Sangon Biotech) was added to DNA samples that had been diluted with wa-
ter (1-, 50-, 100-, 200-fold dilution) following purification using the 3 methods. The samples
were digested using the restriction enzyme HindIII and then the inhibition of restriction
enzyme digestion was assessed. The 20-μL reaction contained different dilution of DNA,
10 U HindIII, and 0.5 mg λDNA. Digestion was carried out for 1.5 h at 37°C and then for
2 min at 65°C.

Qualitative PCR analysis

Primers used in this study included those specific for bacterial 16S rDNA.
The following bacterial primers were used: 27F: 5'-AGAGTTTGATCCTGGCTACCTTGTTACGACTT-3', and 1492R: 5'-GGCTACCTTGTTACGACTT-3'. Amplification was carried out in
a 50-μL reaction volume containing 1X buffer (50 mM KCl, 10 mM Tris-HCl, pH 8.3, 1.5 mM
MgCl₂), 2 μL template DNA, 200 μM dNTPs, 0.6 μM of each primer, and 1.5 U Taq DNA
polymerase (Takara Biotechnology Co., Ltd., Dalian, China). All PCR amplifications were
performed in a GeneAmp PCR 9700 system (Applied Biosystems, Foster City, CA, USA)
using the following temperature program: 94°C for 5 min, followed by 35 cycles at 94°C for
1 min, 61°C for 1 min, 72°C for 1 min, and a final extension at 72°C for 7 min. Amplified
products were electrophoresed on a 2% agarose gel for approximately 20 min at 100 V and
stained with ethidium bromide for visualization.

RESULTS

Total DNA extraction and detection

In an initial experiment, 50-mL freshwater samples were extracted using 3 methods
according to the protocols outlined above. DNA extraction procedures for these methods were
carried out in triplicate. Figure 1 shows that only the filtration-membrane method and modi-
fied filtration-membrane method were useful for extracting sufficient amounts of DNA to be
detected by agarose gel electrophoresis, while no DNA was detected using the centrifugation
method. The DNA yield using the centrifugation method was below detection range, which
may have been because microbial communities could not be aggregated to a sufficient extent
for DNA extraction, and this could only be achieved through centrifugation (5000 g, 20 min).
Compared to the modified filtration-membrane method, the extracted DNA bands from the
filtration-membrane method were brighter, indicating a higher concentration, and degraded.
Degradation may have been the result of nuclease contamination. Lack of SDS lysis using the
filtration-membrane method may have been because SDS inhibited nucleases by preventing
their interaction with hydrophilic nucleic acids.

DNA extraction yield and purity

To examine the purity of the DNA extract, the UV absorption at 230, 260, and 280
nm were measured and the A₂₆₀/₂₃₀ and A₂₈₀/₂₆₀ ratios were calculated (Table 1). The filtration-
membrane method provided higher final DNA yield than the other methods. The DNA yield
using the modified filtration-membrane method may have been lower than that of the filtration-membrane method because of the consecutive application of 2 steps of adsorption to the column during DNA purification. The amount of eluted DNA was reduced after purification on the adsorption column, and thus a 2nd application may have resulted in a higher loss of DNA (Knauth et al., 2013).

The $A_{260/230}$ value of the DNA extract following the modified filtration-membrane method was 1.91, while the $A_{260/230}$ ratios of the DNA extract obtained using the other methods were lower, indicating the presence of co-purified contaminants, as described in the NanoDrop manual. The DNA extracted using the modified filtration-membrane method showed the highest purities based on the $A_{260/230}$ and $A_{260/280}$ ratios. The reason may be that this method involved a purification step with 5.5 M guanidine thiocyanate, which was necessary to prevent inhibition of PCR.

**Effect of contaminants in extraction DNA on the restriction enzyme digestions**

$\lambda$DNA digestion of DNA that had been diluted with water was accomplished using the restriction enzyme HindIII. Contaminants, such as acid, are known to interfere with restriction enzyme digestion and PCR amplification (Mangiapan et al., 1996; Bürgmann et al., 2001; Dong et al., 2006). Restriction digestion of $\lambda$DNA produced clear bands, indicating that the restriction reaction could be carried out in diluted DNA samples. As shown in Figure 2, when $\lambda$DNA was added to extracted DNA (1-fold dilution) obtained using the 3 described methods,
only the modified filtration-membrane method sample was digested to produce clear bands, indicating that the least amount of contaminant was present in this sample. Meanwhile, for extracted DNA (50-, 100-, 200-fold dilution) using all 3 methods, the λDNA could be digested to produce clear bands. The reason may be that the contaminants in the extracted DNA were diluted by equal multiples, which did not inhibit restriction enzyme digestion.

**Figure 2.** Restriction enzyme digestion patterns of λDNA adding dilutions of water DNA. λDNA enzyme digestion of diluted DNA samples (1-, 50-, 100-, 200-fold dilution) by centrifugation method (lanes 1–4); filtration-membrane method (lanes 5–8); modified filtration-membrane method (lanes 9–12); lane M = λDNA/HindIII DNA marker.

**PCR analysis**

After evaluating the quality and quantity of DNA, PCR was used to amplify the 16S rDNA of bacteria to produce a fragment of approximately 1450 bp. These samples were used to analyze the DNA obtained using the 3 methods. The results are shown in Figure 3.

**Figure 3.** Agarose gel results showing PCR product amplification. Centrifugation method (lanes 1 and 2); filtration-membrane method (lanes 3 and 4); modified filtration-membrane method (lanes 5 and 6); lane M = DL2000; no template control: lanes 7 and 8.
The PCR amplicons showed nearly equal results regarding DNA extraction regardless of which of the 3 methods was used. As shown in Table 1, the filtration-membrane method showed a higher final DNA yield, while PCR product amplification was lower than that using the modified filtration-membrane method. This may be because the DNA extracted using the filtration-membrane method contained more contaminating substances such as proteins, polysaccharides, and humic acid, which severely inhibited PCR amplification. The efficiency of PCR using DNA purified using the modified filtration-membrane method was the highest, indicating that this sample is the most suitable for use in molecular applications.

**DISCUSSION**

Acquiring quality genomic DNA samples is a key step in performing molecular applications involving genomic studies (Chacon-Cortes et al., 2012). Moreover, the components present in freshwater are very complex, containing humic acid, which is similar to nucleic acids in its physical and chemical characteristics. Therefore, humic acid and nucleic acids compete for binding the DNA precipitation or absorption sites of the purification column (Harry et al., 1999).

An ideal extraction method should fulfill the following 2 requirements: the amount of extracted DNA should be high and the extracted nucleic acids should be pure and intact, allowing for molecular biology studies to be performed. In the modified filtration-membrane method, the following techniques enhanced and optimized the extraction efficiency and purity. Using a 0.22-µm filtration membrane in combination with zirconium beads is advantageous because the cells were disrupted not only in a chemical manner but also mechanically. The silica membrane of the absorption column was very important for DNA purification, while contaminant substances such as proteins and polysaccharides were efficiently removed. Therefore, these purification steps may be indispensable when the DNA will be used for PCR.

**CONCLUSIONS**

In this study, we compared 3 methods of extracting DNA from total freshwater microbial communities. The results showed that the modified filtration-membrane method was the most suitable for use in molecular applications.

**ACKNOWLEDGMENTS**

Research supported by the the Fund of Key Project of Science and Technology of the Shanghai Committee of Agriculture (#2011-6-2), the Shanghai Postdoctoral Sustentation Fund (Grant #13R21421000), the China Postdoctoral Science Foundation (Grant #2013M531197), and the National Science and Technology Major Project of the Ministry of Science and Technology of China (#2012ZX07101-009)

**REFERENCES**


