DIRECT LYSIS GLASS MILK METHOD OF GENOMIC DNA EXTRACTION REVEALS GREATER ARCHAEOAL DIVERSITY IN ANAEROBIC BIODIGESTER SLURRY AS ASSESSED THROUGH DENATURED GRADIENT GEL ELECTROPHORESIS

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ABSTRACT

DNA extraction from anaerobic biodigester slurry is a critical step in all phylogenetic and metagenomic approaches to characterize highly diverse biodigester ecosystem, but little is known about the efficiency of different extraction procedures and their impact on subsequent analyses of microbial communities. The assessment of performance differences, therefore, is a concrete step towards the determination of optimal DNA extraction method for biodigester slurry, which can provide a more reliable comparison of the meta-analysis results obtained in different conditions. Here, we report a highly efficient direct lysis genomic DNA extraction method (Glass milk method) from the slurry of an anaerobic biodigester. This method was compared with five commercially available DNA extraction kits and two different manual methods with regard to DNA extraction, purification efficiencies and representation of archaeal diversity through PCR-DGGE banding pattern of 16S rRNA gene, Shannon-Wiener and Simpson’s indices of diversity and multidimensional scaling suggested that manual methods of DNA isolation revealed far greater archaeal diversity as compared to the commercial kits. Further, in real-time PCR analysis, a methanogens-specific 16S rRNA gene concentration was observed more in manual methods than in commercial kits. The glass milk DNA extraction method proposed here is very useful in pursuing phylogenetic and metagenomic approaches to characterize the highly complex anaerobic biodigester ecosystem with greater reliability and precision.

KEYWORDS

Biodigester
PCR-DGGE
16S rRNA
DNA extraction
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1 Introduction

Microbial degradation of organic matter occurring in the absence of molecular oxygen, has been harnessed as one of the most effective methods of waste decomposition and renewable-energy production. During this process, bacteria degrade macromolecules such as carbohydrates, lipids and proteins into short chain fatty acids, hydrogen and carbon dioxide that are subsequently utilized by the archaea to produce methane, carbon dioxide and small amounts of other gases through a sequential process involving four main reactions: hydrolysis, acidogenesis, acetogenesis and methanogenesis. Methanogens are classified in Archaeal phylum Euryarchaeota (Solli et al., 2014), and methane is produced as the terminal step of the anaerobic process. The methanogenic microbes that usually dominate in biogas reactors belong to acetoclastic group (Zinder, 1993). Worldwide, a great deal of anaerobic digesters have been built with varying capacity ranging from large mechanized systems that operate at constant temperatures to smaller family and community digesters that have no provisions for temperature control (Nand, 1999). It is estimated that there are about 30 million anaerobic biodigesters of small-scale in Asia and Latin America that are of different type such as ambient temperature fixed-dome, floating dome type and Taiwanese model (Huttunen & Lampinen, 2005; Lansing et al., 2008). Defence Research & Development Establishment, Gwalior (India) has developed a technology for bioggradation of human waste using cold-active anaerobic microbial inoculum (AMI). The performance of biotoilets based on this biodigester technology has been demonstrated in different areas of high altitude, plains as well as on mobile platforms such as passenger coaches in Indian Railways.

Composition of the microbial community in a biogas digester directly determines its efficiency and biogas yield. The complex interactions between bacterial and archaeal microbiome dictate the performance of an anaerobic digester to a considerable extent, which are greatly influenced by environmental factors. The information on microbial community dynamics in anaerobic digester can provide newer insights in microbiology which may be useful for improving the design and performance of new generation biodigesters. Therefore, study of microbial community dynamics during the process of bioggradation of organic matter in biodigester is one of the crucial aspects concerning the monitoring of biodigester performance. In past, diverse culture-based as well as molecular methods have been recruited for gaining information on taxonomy and phylogeny of microbial communities of anaerobic biodigesters by various researchers (Vanwonerghem et al., 2014; Ziganshina et al., 2014). Similarly, culture-independent approaches such as Denaturing Gradient Gel Electrophoresis (DGGE) have also been used extensively in characterization of the complex microbial community. Although there has been a rapid development in high-throughput and inexpensive sequence analyses, nevertheless, DGGE still continues to serve as an important component of analyses of microbial communities (Green et al., 2010). According to Muyzer et al. (1993), DGGE profiles of the V3 region was simpler and relatively cost-effective for analysing the structural variation between different microbial systems or the spatio-temporal dynamics of the same system as compared to the sequencing analysis of 16S rDNA genes in clone libraries. It could also be useful in tracing the variation of dominant microbial organism in a biogas digester. PCR/DGGE strategies are in vogue for the purpose of characterization of the microbial communities, and evaluation of the time required for adaptation of the granular inoculum to the conditions in the anaerobic reactors fed with manure (Bergland et al., 2015).

Nucleic acid extraction is the first step for any metagenomic study of microbial community dynamics. In anaerobic microbial system, microbial cells remain tightly bound to the substrate, and high levels of humic contamination poses particular challenges in terms of getting required quality and quantity of DNA that can be amplified through PCR for the purpose of analysis of microbial community (Qiao et al., 2013). During DNA isolation from activated sludge, it is difficult to achieve complete cell lysis owing to heavy encapsulation of microorganisms (McCllroy & Porter, 2010). Many researchers, therefore, have highlighted the importance of optimizations of extraction procedures, taking into account cell-lysis efficiency and recovery of DNA for getting information on size or structure of microbial community (Lipp et al, 2008; Morono et al., 2014). It has also been emphasized that suitable modifications in prevailing protocols are often imperative in case of the samples having low densities of microbial population or having high adsorptive properties, or complex matrices (Alain et al., 2011; Nielsen et al., 2014). Commercially available kits may be useful for nucleic acid extraction from conventional samples. Nevertheless, for meeting specific extraction requirements of complex samples like those of anaerobic digesters, such kits may not yield desired results. Another bottleneck is that kits may not be available at commercial level for unconventional uses for which only a limited number of scientists are interested (Ogram et al., 1987; Corinaldesi et al., 2005). According to Weaver (2013), susceptibility level of microbes to chemical lysis procedures tends to differ. The Gram-positive bacteria have cell walls with thicker layers of peptidoglycan in comparison to that of Gram-negative bacteria. Therefore, the procedures used in DNA extraction may be prejudiced toward isolating DNA from certain types of microbes only thereby making it a challenging aspect to isolate DNA that correctly represents the diversity of microbes in the sampled community (Weaver, 2013). These challenges underpin the relevance of tailor-made extraction methods to selected realms of
nucleic acid based research despite the extra effort on the part of preparing the appropriate reagents and more time-intensive protocols for extraction (Alain et al., 2011; Morono et al., 2014; Alawi et al., 2014).

Although considerable efforts have been made to study on DNA extraction from soil or composting, nevertheless, DNA extraction approaches for complex anaerobic microbial consortium are still limited. As a consequence of this lack of progress, DNA extraction remains a major bottleneck in the process of analyzing large number of samples from anaerobic digesters. Considering the above-mentioned challenges, the present study aimed to provide an improved direct lysis method (Glass milk method) for genomic DNA extraction of archaeal community from the slurry of an anaerobic biodigester, through optimization of extraction procedures and comparison with commercially available five DNA extraction kits and other two manual methods.

2 Materials and Methods

2.1 Operation of Experimental Biodigester

The experimental biodigester of 2 L working volume was seeded with 30 % of the cold active AMI that contained specially adapted consortium of bacteria belonging to four different groups namely hydrolytic, acidogenic, acetogenic and methanogenic bacteria. The biodigester was fed in semi-continuous mode at 10 days of hydraulic retention time (HRT) with a configuration of 100 g of night soil mixed with 100 ml of tap water and operated under mesophilic conditions. The bioreactor was constantly stirred before the evacuation of waste and after feeding. Slurry samples for microbial DNA extraction were taken in stabilized phase of biodigester. Methane content in biogas was measured every day using a Thermo ULTRA gas chromatography (Singh et al., 1995). The determination of chemical oxygen demand (COD) was performed according to the standard protocol (APHA, 2005).

2.2 DNA extraction using commercial kits

To study the community structure and population dynamics during degradation of night soil in anaerobic biodigester, DNA recovery using five commercial kits and three manual methods was compared. The kits evaluated in this study were selected to represent commercially available DNA extraction methodologies, the use of which has been reported in literature. Five commercially available DNA extraction kits, i.e., ZR Soil Microbe DNA Kit (ZR), Power Soil DNA Isolation Kit (PS), Ultra Clean Fecal DNA Kit (UCF), Ultra Clean Soil DNA Isolation Kit (UCS), Qiagen Stool DNA Kit (QS) were evaluated in this study. To compare the five extraction kits for their efficiency to recover DNA from biodigester slurry, extractions were performed in triplicate according to the manufacturers’ instructions. Comparison of various extraction methods and approach followed is provided in Table 1.

**Table 1 Comparison of various DNA extraction methods**

| DNA extraction method | Approach followed* |
|-----------------------|--------------------|
| ZR                    | Bead beating purification by spin filter columns |
| PS                    | Bead beating purification by spin filter columns; Kit intended for use with samples containing high humic acid |
| UCF                   | Bead beating purification by spin filter columns; Kit intended for DNA isolation from feces |
| UCS                   | Bead beating purification by spin filter columns |
| QS                    | Bead beating purification by spin filter columns; Kit intended for DNA isolation from feces |
| MM1                   | Freeze thawing in liquid nitrogen in presence of SDS; CTAB purification |
| MM2                   | Freeze thawing in liquid nitrogen in presence of SDS; PEG 4000 purification |
| MM3                   | Freeze thawing in liquid N2 in presence of SDS; Adsorption to silica followed by DNA precipitation with Isopropanol |

*For consistency, each resultant DNA extract sample was suspended in 100 µl of the eluant solution.
ZR, Zymo Research Kit; PS, Power Soil Kit; UCF, Ultra Clean Fecal DNA Kit; UCS, Ultra Clean Soil DNA Kit; QS, Qiagen Stool Kit; MM1, Manual Method 1; MM2, Manual Method 2; MM3, Manual Method 3.

2.3 DNA extraction using Manual methods

2.3.1 Direct lysis and purification method

Direct lysis of microbial cells was carried out employing combination of physical and chemical methods (freeze thawing in liquid nitrogen in presence of anionic detergent, Sodium Dodecyl Sulphate). DNA was extracted using the method suggested by Yeates et al. (1997) and purified subsequently with CTAB (Manual Method 1, MM1) and PEG 4000 (Manual Method 2, MM2).
2.3.2 Glass milk method (Manual Method 3, MM3)

2.3.2 Desorption of microbial cells

Biodigester slurry (1 ml) was washed twice with 0.85 % NaCl and pellet was resuspended in phosphate buffered saline solution containing 0.15 % Tween 80. The suspension was vortexed for 15 min and the feed material was pelleted using low speed centrifugation (500 x g for 30 seconds at room temperature). Supernatant fractions were recovered and transferred to sterile tubes followed by centrifugation at 10000 x g for 20 min at room temperature. The pelleted material was used for further DNA extraction.

2.3.2.2 Glass milk preparation

Silicon dioxide (5 g), of 0.5-10 μm diameter obtained from Sigma, USA, was suspended in 50 ml of water and centrifuged at 2000 x g for 5 min. The supernatant was discarded and resuspended in 50 ml of water. The pH was adjusted to 6.8 with conc. HCl. The sediment was left for 2 h allowing precipitation of silica and then supernatant was discarded. The sedimentation procedure was repeated twice followed by centrifugation at 3000 x g for 5 min. Residual water was removed with pipette and the silica pellet was resuspended in 30 ml of 3 M guanidine isothiocyanate. Using a wide bore or cut-off tip, 1.5 ml of slurry was taken in a 2-ml eppendorf vial containing the Zirconium glass beads (Biospec Products Inc) and spun at 14000 x g for 5 min in a micro-centrifuge. The supernatant was discarded and the pellet was resuspended in 1 ml cell lysis buffer, 100 μl of 3 M potassium acetate solution and 100 μl H2O. The sample was frozen in liquid nitrogen and thawed in water at room temperature followed by vigorous vortexing for 2 min. This step was repeated twice. It was spun at 4 C for 15 min at 14000 x g. Supernatant (300 μl) was transferred to fresh tube along with 600 μl of glass milk and mixed by continuous shaking on rotating table for 5 min followed by centrifugation at 14000 x g for 1 min. Supernatant was discarded and the pellet was washed with 500 μl of ice-chilled ethanol (70%) followed by centrifugation at 10000 x g for 1 min. To remove any residual ethanol, final spinning was done for 20 seconds. DNA was eluted from matrix by adding 110 μl of nuclease free H2O to the pellet, vortexing and final spinning at 12000 x g for 1 min. Supernatant (100 μl) containing genomic DNA was collected in a sterilized tube (IAEA, 2004). The genomic DNA, so isolated, was checked for its quality, purity and concentration as described below.

2.3.3 DNA quantification and Agarose gel electrophoresis

The genomic DNA was quantified by ND- 1000 spectrophotometer (Nanodrop, Thermoscientific, USA). The purity of DNA was checked by the absorption ratio at 260/280 nm and at 260 nm/230 nm. The integrity of the extracted genomic DNA was tested by 0.8 % agarose gel electrophoresis.

2.4. PCR-DGGE Analysis

The PCR mixture for each sample contained 25 ng of template DNA (final concentration of 1 ng/μl), 200 μM of each dNTP, 2.5 mM MgCl2, 1X PCR buffer, and 1U of Taq DNA polymerase (Fermentas) as well as 0.5 μM of primers designed to amplify the V3 region of rrs gene from either bacteria or from archaea. Primers used in the study were: 16S1 (5’ GAGTTTGATCCTGGTCA 3’) and 16S2 (5’ ACGGCTACCTTGATACAGCTT 3’) as suggested by Lane (1991) and Arc344F (5’ ACG GGG YGC AGC AGG CGC GA 3’) and Arc519R (5’ GWA TTA CCG CCG CKG CTG 3’) as suggested by Yu and coworkers (2008). A touch down PCR protocol was employed, consisting of an initial denaturation step at 95ºC for 5 min, followed by 20 cycles of 95ºC for 60 seconds, 65ºC to 55ºC for 30 seconds (Decrease of 0.5ºC per cycle) then consistent annealing temperature of 55ºC for 15 cycles and 72ºC for 60 seconds; and a final extension step at 72ºC for 10 min. PCR product (25 μl) was subjected to DGGE using an 8 % (w/v) polyacrylamide gel with a 30 – 65 % denaturing gradient for 5 h at 150 V in a DCode Universal Mutation Detection System (Bio-Rad). Gels were stained with 1x SYBR Green solution (Invitrogen) and visualized using Fluorchem 20, Alpha Innotech.

2.5. DGGE banding pattern analysis

DGGE banding patterns were analyzed using GelCompar® II v6.5 software (Applied Maths, Belgium). Bands were normalized following the conversion of scanned gels. A reference position was defined to align the banding patterns after the bands were associated with the standard for proper comparison. A Kolmogorov–Smirnov nonparametric test of the equality of continuous, one-dimensional probability distributions was performed to compare the given sample with a reference probability distribution. Similarity values among the banding patterns were calculated on the basis of comparison of the corresponding bands using the Dice similarity coefficient. The band matching tolerance and optimization were set at 0.50%. Dendrograms were calculated using UPGMA (unweighted-pair group method using arithmetic averages) clustering algorithm with branch quality set at cophenetic correlation. Further, the statistical analysis of the data on relative archaeal diversity (as estimated through DGGE profiling of various DNA samples obtained through different extraction procedures) was conducted using GelCompar® II v6.5 software employing following methods:

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2.5.1. Shannon-Wiener and Simpson’s indices of diversity

Using GelCompar® II v6.5 software, the Shannon-Wiener and Simpson’s indices of diversity were calculated, taking into account the diversity (i.e., number of categories present in the sampled population), as well as the equitability (i.e., the evenness of the distribution of entries over the different categories (Anonymous, 2010). For a sampled population of \(N\) entries belonging to \(K\) categories, the Shannon-Wiener index (\(H\)) of diversity was calculated as follows:

\[
H = -\sum_{i=1}^{K} \frac{n_i}{N} \cdot \ln \left( \frac{n_i}{N} \right)
\]

where \(n_i\) is the number of entries in category \(i\).

Simpson’s index of diversity is defined as the probability that two consecutive entries will belong to different categories. Given \(K\) categories present in a sampled population of \(N\) entries, the probability of sampling category \(i\) twice consecutively is as follows:

\[
P_i = \frac{n_i(n_i - 1)}{N(N - 1)}
\]

where \(n_i\) is the number of entries in category \(i\).

Here, the probability (\(P\)) of any two samples of the same category is given by:

\[
P = \sum_{i=1}^{K} P_i
\]

Hence, Simpson’s index of diversity (\(D\)) will be \(D = 1 - P\).

2.5.2. Multi-dimensional scaling (MDS)

Using GelCompar® II v6.5 software, MDS was carried out based on the Euclidean distance between two entry points according to the following formula:

\[
d_{ij} = \sqrt{\sum_{k=1}^{p} (x_{ik} - x_{jk})^2}
\]

where \(p\) is the number of dimensions, \(d_{ij}\) is the distance, and \(x_{ik}\) is the data value of the \(i^{th}\) row and \(k^{th}\) column.

2.6. Statistical Analysis

Means and standard deviations of the data pertaining to DNA concentration and purity ratios (i.e., 260/280 and 260/230) were calculated using Microsoft Excel 2007 (Microsoft Corporation, Redmond, WA, USA). Differences between mean values of various parameters viz., DNA concentration and absorbance ratios were compared through Analysis of Variance (ANOVA) followed by Least Significant Difference (LSD) Post-Hoc test. \(P\) value<0.05 was considered statistically significant. SPSS Statistics 17.0 program was used for calculating ANOVA and LSD. Accordingly, statistically significant differences among mean values have been shown with different letters on the columns in Figures 1 and 2.

3 Results and Discussion

3.1. Biodigester performance

Lab-scale experimental Biodigester (2L) operated in fed-batch mode at 10 day-HRT showed the pH value varying between 7 and 8 during the operational period. The COD removal efficiency was recorded as 78 %, and methane content in biogas was found 62 % during the operational period. The effluent concentrations of

Figure 1 DNA yield (ng /µl) in samples obtained through various methods. Error bar shows standard deviation of mean (n=3). Columns without common letter differ significantly at LSD \(P<0.05\). For acronyms, refer Table 1.

Figure 2 DNA purity in terms of absorbance ratios (\(A_{260}/A_{280}\) and \(A_{260}/A_{230}\)) of DNA samples obtained through various methods. Error bar shows standard deviation of mean (n=3). Columns without common letter differ significantly at LSD \(P<0.05\). For acronyms, refer Table 1.
COD, and associated methane yield represented the efficient organic removal and stable process operation (Data not shown) as reported earlier (Singh et al., 1993).

3.2. DNA yield and purity

In current study, the DNA yield (Figure 1) and purity (Figure 2) varied with different extraction methods. Significantly higher DNA yield was obtained in the case of glass milk method (MM3) followed by CTAB purification (MM1) and PEG 4000 purification (MM2) methods (Figure 1) with observed values being 1116.16 ng/µl, 755.83 ng/µl and 262.30 ng/µl respectively. On the contrary, commercially available kits provided very poor yield with no statistically significant difference in DNA concentration amongst them (Figure 1). The DNA yield obtained using commercial kits ranged from 14.43 ng/µl to 80.43 ng/µl.

Several studies have compared extraction methods and reported that they differ in their ability to recover DNA, indicating that no single DNA extraction method is optimal for all kind of samples (Whitehouse & Hottel, 2007; Queipo-Ortuno et al., 2008; Vesty et al., 2017) thereby necessitating further efforts to develop more efficient methods catering to specific requirements. Each method prescribed has advantages and disadvantages rendering the selection of a suitable method a daunting task. The extraction of DNA from sludge samples in anaerobic digesters has rarely been studied or optimized, so there is a lack of standardized methods (Dong et al., 2013). Our findings also unequivocally strengthen the view that the overall quality and yield of genomic DNA is influenced by DNA extraction approach. According to Yeates and coworkers (1997), absorbance ratio of $A_{260}$ to $A_{280}$ (>1.7) and $A_{260}$ to $A_{230}$ (>2.0) is indicative of pure DNA. In the present study, purity of DNA obtained using the glass milk method (MM3) was found to be the best among all the methods as evident from maximum values of 1.83 and 1.96 for ratios of $A_{260}$ to $A_{280}$ and $A_{260}$ to $A_{230}$, respectively (Figure 2). Purity of extracted DNA using other kits (ZR, UCS and QS) was not appreciable as evident from low absorbance ratio ($A_{260}$ to $A_{280}$) ranging from 1.19 to 1.70 indicating presence of absorbing contaminants, possibly the humic acids (Li et al., 2004; Shamia et al., 2017), in the preparations. Purity of DNA in terms of ratio of $A_{260}$ to $A_{230}$ was also found to be significantly high for the glass milk method (MM3) in comparison to other methods.

The majority of nucleic acid extraction recipes are prepared keeping in view the complete lysis of cells and subsequent extraction of intracellular nucleic acids into aqueous solution, the elimination of organic and inorganic molecules (non-nucleic acid) from the aqueous extracts, and the minimization of product losses during the course of purification (Lever et al., 2015). The yield and quality of cellular and bacterial DNA extracts re variably affected by the cell lysis methodology (Sohrabi et al., 2016). The protocol of glass milk method was also designed keeping these aspects in view. Here, the efficient cell lysis was achieved through optimal degree of freeze-thawing and bead beating. Following cell disruption, the nucleolytic degradation was effectively inhibited using guanidine isothiocyanate as reported earlier (Lippke et al., 1987). This was followed by precipitation of SDS-bound proteins using potassium acetate to allow their removal from DNA. In a similar study, highly efficient cell lysis from the samples of anaerobic digestion sludge employed for the evaluation of effectiveness of DNA extraction provided DNA recovery efficiency as high as 86.16% (Dong et al., 2013). SDS is known to aid in lysing cells and solubilizing proteins. In addition, K+-SDS interactions have been utilized in past for protein and DNA analysis studies (Li et al., 1989; Hejazi et al., 2013). Taken together, the extraction yield of PCR-ready DNA was greater in glass milk method (MM3) as compared to other two manual methods and five commercial kits indicating its potential utility and advantage for downstream molecular biological processes for studying microbial communities of anaerobic digester.

3.3. PCR-DGGE profiling

PCR product of the 1492 bp size was obtained for universal 16S primers following thermal amplification. In order to reveal the metagenomic attributes of the extracted DNA, nested PCR was performed on amplicons obtained with universal 16S primers. Many studies have employed DGGE profiles of the V3 regions in order to reflect the dominant community structure and to analyse the structural variation between different systems or the dynamics of the same system (Yu et al., 2008; Liu et al., 2009; Bergland et al., 2015). In the current study, the primers for V3 hyper variable regions of 16S tRNA gene generated the band of ~176 bp in agarose gel electrophoresis. Furthermore, the PCR-DGGE profile indicated wide variation in archaeal diversity in the samples obtained through various approaches of DNA extraction with maximum diversity being observed with respect to glass milk method (Figure 3). Quantitative measurement of diversity in a community can be reflected by calculating Shannon-Wiener and Simpson’s indices of diversity which are among the most commonly used indices. These indices represent as to how evenly the basic entities (such as archeal species) are distributed in the sampled community. In the current study, the pattern of diversity was further confirmed based on comparative values of Shannon-Wiener and Simpson’s indices of diversity (Figure 4A & 4B) with maximum diversity observed for the glass milk method. In addition, optimized three-dimensional representation of similarity matrix calculated based on Euclidean distance for various samples employing MDS technique also indicated greater archeal diversity in respect of glass milk method (Figure 5). Consistent with present findings, recent studies have also brought to light the
instances wherein the choice of cell lysis method used during DNA extraction was found to impact the recovery of specific bacterial phyla. Mechanical lysis was reported to increase the number of different bacterial phyla recovered from saliva (Lazarevic et al., 2013), while the addition of lysozyme to mechanical lysis improves overall bacterial DNA yield from saliva (Sohrabi et al., 2013).

Overall, the glass milk method described here provided high quality DNA even without the use of column purification that can be successfully amplified by PCR. In addition, greater recovery of archaea from the test samples as established through PCR-DGGE profiling indicates the suitability of this method for studying the phylogenetic community composition and functionality in complex communities present in anaerobic biodigester slurry.
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