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### **Abstract:**

To enhance DNA recovery with respect to yield, purity, and fair portrayal of microbial variety, two methods have been devised. Nevertheless, co-purified pollutants often impede DNA amplification from soil. DNA may also be amplified using polymerase chain reaction (PCR) with a variety of DNA targets. This review provides a synopsis of the approaches that have been considered in order to accomplish this difficult task. After lysing 100 g of soil with glass beads and SDS, we precipitated with potassium acetate, polyethylene glycol, phenol, and isopropanol to extract DNA.

Keywords: Soil, DNA, RNA, and Purification

# Introduction

Since their genes are so basic, genetically modified bacteria were the first creatures to undergo laboratory modifications.[1] These creatures have found new uses in modern medicine, where they play a crucial role in the production of huge quantities of pure human proteins. The two important requirements for metagenomic DNA extraction are efficient cell lysis and purification of DNA from the complex milieu of an environmental sample The impossibilityto culture most microorganisms from environmental samples is a fundamental obstacle to understanding microbial ecology and diversityThe year 2017 was covered by Denet and colleagues. The use of DNAbased techniques can overcome this limitation by allowing the fate of particular genes or organisms to be monitored directly in environmental samples. Historically, methods for DNA extraction from silt and

soil relied on massive 100g samples. These extracts were usually contaminated with humic acids which interfered with subsequent molecular biological manipulations. Extensive purification steps were then required to successfully amplify a PCR product, including CsCl-ethidium bromide density gradient centrifugationZhang et al., 2017, or the use of commercial reagents Borneman et al. 1996. These processes make the procedure more complicated and expensive. An approach to soil DNA extraction that requires little purification before PCR amplification is detailed in this work. We compare it to other approaches that are routinely used to extract DNA. A PCR product was obtained rapidly and inexpensively from large amounts of soil, even when contaminated with heavy metals.A rapid, inexpensive, large-scale DNA extraction method involving minimal





purification has been developed that is applicable to various soil typesZhang et al., 2017.

Upon considering the limitations of previous methods (variable efficiency, time consuming and high cost), the current study focused on developing a rapid inexpensive method for extraction of metagenomic DNA with sufficient quantity and purity to be broadly suitable for metagenomic applications. Since, cell lysis and purification are the key steps in metagenomic DNA extraction; this study

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### Vol-6 Issue-01 April 2017

includes a particular focus on these two factors. Cell lysis is accomplished by homogenizing with glass powder that is obtained from laboratory waste glasswareYamanouchi et al., 2018b. Silica, the major component of ground glass powder, has been widely used for DNA extraction from various sources including soils and sediments, tissues and blood of transgenic animals and plasmid from E.coli. Autoclaved silica-based sand has been reported for extraction of fungal DNA, and glass powder along with skim milk was used for detection of Phytophthora infestans Yamanouchi et al., 2018a.



#### Materials and methods

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Soil DNA Purification Protocol

#### A. Preparation the sample :

Spin Columns1. Add 550  $\mu$ l of Inhibitor Removal Resin to each empty Spin Column to be used. Centrifuge for 1 minute at 2000 x g to pack the column2. Decant flow-through and place the column in the same collection tube.3. Add another 550  $\mu$ l of Inhibitor Removal Resin to each packed column. Centrifuge for 2 minutes at 2000 x g.4. Move the column to a clean 1.5-ml collection tubeDahal et al., 2018.Pellet Wash Solution1. For 50 Extractions Kit: Add 45 ml of ethanol to the Pellet Wash Solution before first use. For 5 Extractions Kit: Add

4.5 ml of ethanol to the Pellet Wash Solution before first use.

#### B. Cell Lysis:

Weigh out 100 mg of the soil sample into a 1.5 ml tube.2. Add 250  $\mu$ l of soil DNA extraction buffer and 2  $\mu$ l of proteinase K; vortex briefly.3.



(Optional) To increase the yield of DNA, shake the tube at 37°C for 10 minutes or vortex for 2 minutes. Add 50 µl of Soil Lysis Buffer and vortex briefly.5. Incubate at 65°C for 10 minutes.6. Centrifuge for 2 minutes at 1000 x g.7. Transfer 180 µl of the supernatant to a new tube.8. Add 60 µl of Protein Precipitation Reagent, mix thoroughly by inverting the tube.9. Incubate on ice for 8 minutes. Centrifuge the tubefor 8 minutes at maximum speed.10. Carefully transfer 100-150 µl of the supernatant directly onto the prepared Spin Column .11. Centrifuge for 2 minutes at 2000 x g into the 1.5-ml tube. Discard the column.12. Add 6 µl of

DNA Precipitation Solution, vortex briefly. Incubate the tube at room temperature for 5 minutes.13. Centrifuge for 5 minutes at maximum speed. Carefully decant the supernatant.14. Wash the pellet with 500  $\mu$ l of Pellet Wash Solution. Invert to mix then spin for 3 minutes at maximum speed. Carefully decant the supernatant.15. Repeat the wash and spin.16. Resuspend the pellet in 300  $\mu$ l of TE BufferNan et al. 2014.

### ISSN: 2320-3730

## Vol-6 Issue-01 April 2017

DNA does not amplify by PCR. 1) Optimize cycling conditions. Decrease the annealing temperature of the cycling profile by 2 degrees or more. Some primer pairs require a lower annealing temperature (less stringent conditions) when amplifying soil DNA.2) Use less starting material. Some environmental samples contain significantly larger amounts of enzymatic inhibitors. When using these samples, begin the extraction with less starting material (50 mg).3) Load less extract onto the column.

5) Rewash the pellet with the Pellet Wash Solution. This step is important in removing residual inhibitors of DNA amplification. Eliminate the vortex mixing step. Eliminate the 2 minute vortex mixing step when extracting the DNA. Shake at 37°C instead or simply skip this step entirely.

Soil (loamy sand) was collected on campus at SemnanUniversity in Iran. The SokanSemnan National Park Station samples represent the extremes of pristine vs polluted soils and were compared by further soil testing(Table11).

Troubleshooting DNA Extractions

Ph	3.90
Organic matter %	5
Field capacity 0.33 bar	7.05
CEC (cmol)	1.1
As (mg/kg)	<3
Hg(mg/kg)	<0.7
Zn(mg/kg)	5
Cr(mg/kg)	3.3
Cd(mg/kg)	<0.4
Ni(mg/kg)	1.7
Pb(mg/kg)	15
Cu(mg/kg)	9.5
Mn(mg/Kg)	13

Table 1. Physicochemical	analysis of soil samples
	analysis of som samples

DNA extraction from soil using bead beating Extraction buffer (100 ml of 100 mMTris-HCl [pH 8.0], 100 mM sodium EDTA [pH 8.0], 1.5 M NaCl) was mixed with 100g (wet weight) of soil. Glass beads (100g, Bio-Spec Products, Bartesville,U.S.) were added and the sample



blended in a Bead-Beater (Bio-Spec Products) for 2 minutes. Sodium dodecyl sulphate (SDS) was added (10 ml; 20 %) and blending continued for a further 5 sec. The sample was incubated at 65°C for 1 hr, transferred to centrifuge bottles (250 ml) and centrifuged at 6000g for 10 min. The supernatant was collected, and the soil pellet reextracted with further extraction buffer (100 ml), incubation at 65°C for 10 minutes and centrifugation as described above. Supernatants were transferred to centrifuge tubes (50 ml) containing a half- volume of polyethylene glycol (30%)/sodium chloride (1.6 M), and incubated at room temperature for 2 h. Samples were centrifuged (10,000g for 20 min) and the partially purified nucleic acid pellet wasresuspended in 20 ml of TE (10 mMTris-HCl, 1 mM sodium EDTA, pH 8.0). Potassium acetate (7.5 M) was added to a final concentration of 0.5 M. Samples were transferred to ice for 5 min then centrifuged (16,000 g, 30 min) at 4°C to

precipitate proteins and polysaccharidesShi-Ying et al., 2018. The aqueous phase was extracted with phenol/chloroform and chloroform/isoamyl alcohol and DNA was precipitated by adding 0.6 volume isopropanol. After 2 hat room temperature, DNA was pelleted by centrifugation (16,000g for 30 min) and resuspended in TE (1 ml).

#### DNA extraction using sonication

Extraction buffer (100 ml) was mixed with soil (50g) on ice. The mixture was sonicated using a High Intensity Ultrasonic Processor (Vibra Cell) with a standard 13mm horn solid probe for 150 seconds. The sample was cooled in ice andthe sonication repeated. SDS was added (10 ml; 20%) and the sample incubated at 65°C for 1

h. The sample was transferred to centrifuge bottles (250 ml) and centrifuged at 6000g for 10 min. The supernatant was collected, and the soil pellet reextracted with further extraction buffer (50 ml).

### ISSN: 2320-3730

## Vol-6 Issue-01 April 2017

incubation at 65°C for 10 minutes and centrifuged were adopted as above. Extraction was then continued as per bead beating methodShokri et al., 2016.

DNA extraction using enzymatic lysis

Extraction buffer (100 ml) containing proteinase K (5 mg) was mixed with soil (50g) in 250 ml centrifuge tubes. The sample was incubated at 37°C for 30 minutes with shaking at 180 rpm. SDS was added (10 ml; 20%) and the sample incubated at 65°C for 90 min. The supernatant was collected after centrifugation at 6000g for 10 min at room temperature. Extraction was continued as per bead beating method.

DNA extraction from bacterial cells isolated from soil .

The bacterial fraction of soil was separated from the inorganic or humic layer by a differential centrifugation technique. Bacterial cells were lysed using lysozyme and the DNA purified using ammonium acetate precipitation and ethanol precipitation. DNA was resuspended in TE.

### **Test forCo-Extraction of Contaminants**

Co-extracted humic acids are the major contaminant when DNA is extracted from soil. These compounds absorb at 230 nm whereas DNA absorbs at 260 nm and protein at 280 nm. To evaluate the purity of the extracted DNA, absorbance ratios at 260 nm/230 nm (DNA / humic acids) and 260 nm/280 nm (DNA / protein) were determined.

<u>Table 2</u>Comparison of DNA extraction methods using a single soil

with further extraction burlet (50 mil),						
Method*	:	Number of samples	A260230	A260280		
Bacterial	l cells	4	0.83=0.03	1.10=0.003		
Chemica	l lysis	10	1.06=0.03	1.31=0.03		
Sonicatio	on	4	1.20=0.10	1.41=0.07		
Bead bea	ating	6	1.82=0.05	1.69=0.02		

#### ISSN: 2320-3730



Vol-6 Issue-01 April 2017

DNA diluted 1:100 \*

Table 3: Crude DNA ratios for different soil samples extracted using bead beating.

Sample*	Soil type	A260230	A260280
Western	Clay loam	1.22	1.42
university	Clay loam	1.83	1.71
Sokan Ku-Ring Gai	Loamy sand	1.03	1.3
Balmain power station	Loamy sand	1.33	1.53

DNA diluted 1:100\*

#### **Polymerase Chain Reaction (PCR)**

DNA (1 ml of 1:50 dilution) was mixed with 9 ml of Genereleaser<sup>TM</sup> (Bioventures Inc. Murfreesboro, Tennessee, USA) in a 0.5 ml tube and overlaid with 2 drops of sterile mineral oil. Genereleaser<sup>TM</sup> is a proprietary agent that sequesters inhibitors of PCR. Negative controls containing water only, and Genereleaser<sup>TM</sup> only, were included in each set of reactions. Reaction tubes were heated on the high setting of a 650 Watt microwave oven for 7 min (4550 W/min) in a microwave transparent rack (Bioventures Inc.). An Erlenmeyer flask containing 100 ml of water was included as a microwave sink. Tubes were incubated for at least 10 min at 80°C in an Omn-E PCR machine (Hybaid). PCR master mix (40 µl) was then added to each tube.Final concentrations of reagents were as follows: 20 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 75 mMTris-HCl (pH 9.0), 0.01% (w/v) Tween 20, 2 mM MgCl<sub>2</sub>, 0.5 mM of each primer, 0.2 mM of each deoxyribonucleotide triphosphate, and 1 U Red Hot DNA Polymerase (Advanced Biotechnologies, Surrey, UK). The following thermal cycle was performed : 94°C 3 min (1 cycle), 94°C 1 min, 55°C 1min, 72°C 2 min (35 cycles), 72°C 5 min (1 cycle) Burgmann et al. 2001.

### **Gel Electrophoresis**

An aliquot (7  $\mu$ l) of each amplification reaction was analysed on 2% w/v agarose gels cast and run

in TBE buffer (pH 8.3) (12). Gels were stained with ethidium bromide and photographed using transmitted U.V. light and Polaroid film. A 100 base pair marker (Pharmacia, LKB) was included on every gel.

### **Results and Discussion**

DNA extraction from soil has three requirements: extraction of high molecular weight DNA; extraction of DNA free from inhibitors for subsequent molecular biological manipulations to be performed; and representative lysis of microorganisms within the sample. In this paper, we tested a number of DNA extraction methods for their ability to fulfill these requirementsZhang et al. 2004.

DNA extracted using sonication was more degraded than the oneobtained withthe other tested methods. The size of extracted DNA ranged from less than 500 bp to greater than 20 kb. Methods that shear DNA, such as sonication, generally result in DNA of 100-500 bp. Higher molecular weight DNA is desirable for PCR since the greater the size of the DNA, the less likely is the formation of chimeras during PCR. The bead beating method used here performed better than those previously reported which usually extract DNA of less than 10 kb in size. The DNA extraction methods that did not use sonication all produced DNA of greater than 20 kb.

Organic matter is the major source of inhibitors that may be co-extracted from soil with the



microbial DNA. In particular, humic acids pose a considerable problem and will interfere in enzymatic manipulations of

DNAHolben et al. 1985. DNA polymerases have been found to be inhibited by as little as  $1 \mu l$  of undiluted humic-acid-like extract, regardless of the amount of DNA presentSaburi et al., 2017.

The humic materials in soil have similar size and charge characteristics to DNA resulting in their copurification

, evident by the extractions being brown in colour. Humic contaminants also interfere in DNA quantitation since they exhibit absorbance at both 230nm and at 260nm, the later used to quantitate DNA. This characteristic can be used to determine the level of contamination of humic material by examining absorbance ratios. A high 260/230 ratio (>2) is indicative of pure DNA, while a low ratio is indicative of humic acid contamination and a high 260/280 ratio (>1.7) is indicative of pure DNA, while a low ratio is indicative of protein contamination. When the DNA extraction methods were compared ((Table2),, the bead beating method consistently extracted DNA with higher 260/230 and 260/280 ratios. This indicated that the

#### ISSN: 2320-3730

## Vol-6 Issue-01 April 2017

DNA was contaminated with fewer humic acidlike compounds. Although the extracts were still brown in colour, dilution of the DNA to 1:50 from all methods was suitable to produce a PCR product. Heavy metal ions, such as are present in the Balmain soil ((Table1), also contribute to inhibitory effectsHolben et al. 1985. Here we have demonstrated that a PCR product from soil DNA contaminated with humic acids and heavy metals can be obtained without the use of expensive purification products.

To determine the diversity of microorganisms from which DNA had been extracted, different primer sets were tested ((Table4), including both multi- and single-copy genes. The multi-copy targets included the prokaryotic small subunit rRNA, prokaryotic rRNA intergenic spacer region, the eukaryotic rRNA internal transcribed spacer (ITS) region, the ITS region for lichen fungi, and the HSP70 family of proteins while the low abundance targets included fungal β-tubulin, and nifH genes. With dilution of DNA from each PCR extraction technique, successful amplification was achieved with all primers tested (Fig.1)..



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Vol-6 Issue-01 April 2017



### Fig. 1:

Example of PCR amplification products using various DNA targets with soil extracted by enzymatic lysis or bead beating. Lane 1: 100 bp marker; lane 2: enzymatic lysis DNA with 16S rRNA primers; lane 3: bead eating DNA



#### ISSN: 2320-3730

### Vol-6 Issue-01 April 2017



with 16S rRNA primers;

**Fig 2**In this simple and rapid process, the soil sample are homogenized and lysed by the buffer containing glass beads, Proteinase K and detergents. Provided special buffer will remove debris, proteins, and polysaccharides by precipitation and other contaminants are washed away by alcohol containing wash buffer. Finally, the purified DNA is eluted by low-salt elution buffer or water.

Due to ease of the method, the reduced coextraction of inhibitors (Tables2 and <u>3</u>) and the greater confidence that bead beating would lyse all microbial cells in the soil, this was the method of choice and concentrated on for further analysis . Bead beating has been found to have a lysis efficiency of greater than 90% . The PCR results reported here provide further evidence to support this with products from both bacterial and fungal elements of the soil microbiota being obtained. The bead beating direct lysis method described here extracts between 1.5 and 2.35 mgml<sup>-1</sup> of DNA from 100g of soil or 15-23.5  $\mu$ g DNAg<sup>-1</sup> soil. Extraction methods using small soil samples ranging from 5g to 100 mg of soil have extracted 9-25 µg DNAg<sup>-1</sup> soil, 12 µgg<sup>-1</sup>, 1-100 µgg<sup>-1</sup>, and 2.5-26.9  $\mu$ gg<sup>-1</sup>. The method described here is therefore at least as efficient as the above methods. Various methods are available for metagenomic DNA extraction based on chemical ormechanical lysis of microbial cellspresent in the soil. Among these methods, glass bead beating is considered to be an effective technique for metagenomic DNA extraction. This method has also been modified in previous reports to be suitable for different soil types. Commercial kits such as Fast DNA SPIN kit for soil, MP Biomedicals, Santa Ana, CA) and Ultra Clean Mo Bio Soil DNA isolation kit are also based on the method of bead beatingYamanouchi et al., 2018b.

The focus of DNA extraction methods has moved to rapid performance of molecular techniques, avoiding extensive purification steps. Using the bead beating DNA extraction method described here, crude microbial DNA could be extracted from a variety of soil types and dilution of this DNA was sufficient for successful PCR from both



high- and low-copy number genes.

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### ISSN: 2320-3730

## Vol-6 Issue-01 April 2017

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