# **BIOC3301: Protocol for Metagenomics Analysis**

Team 8: Kan Yan Chloe Li, Siân Ashk O'Shea, and Sin Rachel Lau

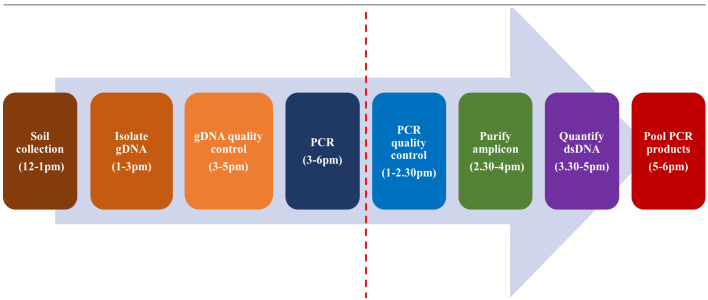


Figure 1: Overview of experiment. Line break at the point at which the PCR is running; this can be done overnight.

#### 1: Collect soil from Gordon Square Garden

#### Precautions and factors to consider:

- Location: Gordon Square Garden, London, UK
- Ensure sampling GPS coordinates, date, time and conditions (temperature, weather) are noted
- Always wear gloves to avoid contaminating soil samples

#### **Equipment and materials:**

- Small shovel
- Bulb planter
- Rubber gloves
- Falcon tubes (50 mL)

- 1. Generate map overview of Gordon Square and divide into 5 equal regions (see map overview in Appendix 1). Note: Systematic sampling is representative of entire park to avoid bias and increase chances of sampling within different conditions (e.g. different microbiomes, different chemical and physical soil conditions).
- 2. Generate axes for each region of the map.
- 3. Generate 2 random numbers (ensures unbiased samples) within the axes using a calculator and use as coordinates to locate the sampling site within the region of the map.
- 4. Use Global Positioning System to locate coordinate location. Location must be on soil and not on for example, a path. Otherwise repeat random generation of coordinates.
- 5. Use a bulb planter to take a soil sample at a depth of 10 cm. Note: Depth selected to avoid detritus present on or near the surface but to maintain a high bacterial presence, as this decreases with increased depth. Ensure to take samples at a consistent depth.
- 6. Transfer ~2 cm<sup>3</sup> of the sample to a 50 mL Falcon tube (keep each sample separated in a different container).
- 7. Label the sample with its region and number.
- 8. Note observable conditions for each sample site, including: sunlight/shading, moisture of soil, compactness of soil, vegetation, litter, animal remains, etc.
- 9. Use pH test strip to measure the pH of the soil sample.
- 10. Take a further 2 samples to act as controls.
- 11. Autoclave one control to act as the negative controls.
- 12. Add bacterial genomic DNA to the other control, to act as the positive control.
- 13. Store samples in a refrigerator.

# 2: Soil gDNA isolation using PowerSoil® DNA Isolation Kit (MO BIO Laboratories, CA, USA)

#### **Precautions:**

- Always wear gloves
- Solution C5 contains ethanol, which is flammable. Keep away from naked flames.

## **Equipment and materials:**

- Microcentrifuge
- Pipettors and pipette tips
- Vortex and vortex adaptor
- PowerSoil® DNA Isolation Kit, which contains:
  - o PowerSoil® Bead Tubes
  - o PowerSoil® Solution C1
  - o PowerSoil® Solution C2 Inhibitor Removal Technology®
  - o PowerSoil® Solution C3
  - o PowerSoil® Solution C4
  - o PowerSoil® Solution C5 ethanol wash solution
  - o PowerSoil® Solution C6 10 mM Tris
  - o PowerSoil® Spin Filters units in 2 mL tubes
  - o PowerSoil® Collection tubes 2 mL

- 1. Add 0.25 g of soil sample to **PowerSoil® Bead Tube**. If sample is of high water content, centrifuge at room temperature for 30 s at 10,000 g and remove water using pipette.
- 2. Gently vortex to mix the sample.
- 3. Check that **Solution C1** has not precipitated. If precipitated, heat solution to 60°C using a water bath until the precipitate has dissolved before use.
- 4. Pipette 60 μL of **Solution C1** to the sample and invert several times or vortex briefly.
- 5. Secure **PowerBead tubes** horizontally using the MoBio Vortex Adapter tube holder for the vortex. Vortex at maximum speed for 10 min. **Note:** If you are using the 24 place Vortex Adapter for more than 12 preps, increase the vortex time by 5-10 min. If cells are difficult to lyse, a 10 min incubation at 70°C, after adding solution C1, can be performed.
- 6. Centrifuge **PowerTubes** at 10,000 g (~13,000 rpm) for 30 s at room temperature. **CAUTION**: Do not exceed 10,000 g or tubes may break.
- 7. Transfer supernatant to a clean 2 mL **Collection Tube**. Expect 400-500 μL of supernatant. **Note:** supernatant may have a dark appearance and still contain some soil particles.
- 8. Pipette 250 µL of **Solution C2** and vortex for 5 s.
- 9. Incubate tubes at 4°C for 5 min.
- 10. Centrifuge tubes at 10,000 g for 1 min at room temperature.
- 11. Avoiding the pellet, pipette up to 600 μL of supernatant to a clean 2 mL Collection Tube.
- 12. Add 200  $\mu$ L of **Solution C3** to the tubes and vortex briefly.
- 13. Incubate at 4°C for 5 min.
- 14. Centrifuge the tubes at room temperature for 1 min at 10,000 g.
- 15. Avoiding the pellet, transfer up to (but no more than) 750 μL of supernatant into a clean 2 mL collection tube.
- 16. Shake to mix **Solution C4**. Add 1.2 mL of **Solution C4** to the supernatant and vortex for 5 s. **CAUTION:** ensure solution does not exceed rim of tube.
- 17. Load ~675 µL of the collection tube solution onto a labelled **Spin Filter**.
- 18. Centrifuge at 10,000 g for 1 min at room temperature.
- 19. Discard the flow through and add an additional 675 μL of supernatant to the **Spin Filter**.
- 20. Centrifuge at 10,000 g for 1 min at room temperature.
- 21. Load the remaining supernatant onto the **Spin Filter**.
- 22. Centrifuge at 10,000 g for 1 min at room temperature.
- 23. Add 500 µL of Solution C5.
- 24. Centrifuge at room temperature for 30 s at 10,000 g.
- 25. Discard the flow through from the 2 mL Collection Tube.
- 26. Centrifuge at room temperature for 1 min at 10,000 g.
- 27. Carefully place **Spin Filter** in a clean 2 mL **Collection Tube**. **CAUTION**: Avoid splashing of **Solution C5** onto the **Spin Filter**.
- 28. Add 100 µL of **Solution C6** to the centre of the white filter membrane.

- 29. Centrifuge at room temperature for 30 s.
- 30. Discard the **Spin Filter**. The gDNA is in the tube.
- 31. Store DNA at -20 to -80°C.

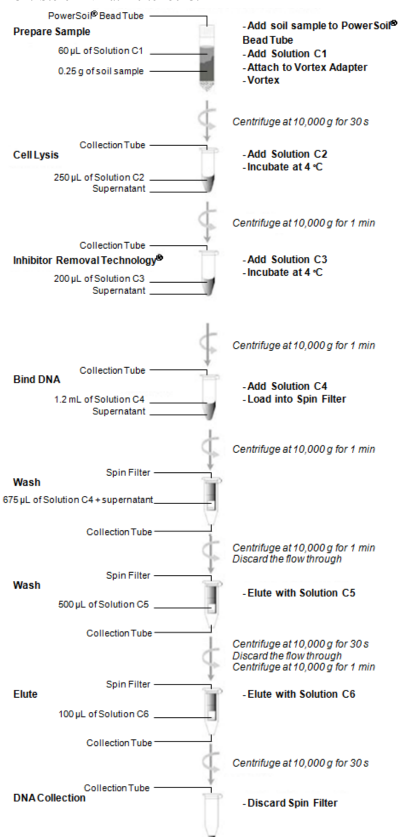


Figure 2: Workflow summary of the soil gDNA isolation process using the PowerSoil® DNA Isolation Kit.

## 3: Check quality of gDNA by gel electrophoresis

#### **Precautions:**

## **Equipment and materials:**

• Always wear gloves

- Pipettors and pipette tips
- Eppendorf tubes
- Agarose powder
- 12 x14 cm horizontal gel electrophoresis device
- 1% ethidium bromide
- 5X loading dye

- 1. Measure 0.8 g of agarose powder.
- 2. Cast a ~100 mL **0.5% agarose gel** by mixing 0.8 g of agarose powder with 2 mL of 1X TAE buffer and 98 mL of distilled water in a microwavable flask.
- 3. Microwave flask in pulses with occasional swirling, for 1-3 min until the agarose is completely dissolved. **CAUTION:** do not over boil the solution.
- 4. Allow agarose solution to cool to  $\sim 50^{\circ}$ C (hands can comfortably hold flask).
- 5. Add 0.5  $\mu$ L of stock ethidium bromide to a final concentration of ~0.5  $\mu$ g/mL.
- 6. Pour the agarose into a tray with the narrow well comb in its fixed position. Allow it to set at room temperature for 20-30 min (until the gel has completely solidified).
- 7. Once solidified, place the agarose gel into the electrophoresis unit/gel box.
- 8. Make 50 mL of 1X TAE buffer.
- 9. Carefully pour the buffer solution over the gel so that it covers the entire gel.
- 10. Transfer 1  $\mu$ L of gDNA sample(s) (concentration between 50 ng 500 ng) into clean, labelled tube(s). Bring the total volume up to 4  $\mu$ L with 1X TE Buffer, pH 8.0.
- 11. Add 1  $\mu$ L of 5X loading dye. The amount of loading dye should remain consistent among all samples to maintain even DNA migration on the gel.
- 12. Mix by pipetting up and down a few times.
- 13. Load onto gel according to the format listed below (where n is number of samples to be run):
  - > Well 1: 10 μl of 1kB + ladder (Invitrogen), 100 ng/μL.
  - ➤ Well 2 to n: gDNA samples.
  - ➤ If DNA floats out of the well, some ethanol from Solution C5 may remain in the sample. Remove residual ethanol in a speed vacuum, desiccator, or air dry.
- 14. Run gel at 120V until the dye line is  $\sim$ 70-75% of the way down the gel ( $\sim$ 90 min).
- 15. Switch off power and disconnect the electrodes from the power source.
- 16. Carefully remove the gel from gel box.
- 17. Using a device that has UV light, take an image and visualise the DNA. **Note:** Expect the bands to appear streaked due to 'ungentle' bead treatment. Look for presence of smearing as this can indicate: (i) RNA contamination, (ii) DNA degradation, and (iii) impurities such as proteins and polysaccharides; these factors can inhibit chemical reactions during library construction.

# 4: Amplify 16S rDNA V4 region by PCR using BioMix<sup>TM</sup> (BioLine Reagents Limited, London, UK)

#### **Procedure:**

- 1. Set up one large 72 µL master-mix and distribute into three tubes, according to table 1, as follows:
  - A. Add 24  $\mu$ L of the master-mix (sequences for primers in Appendix 2).
  - B. Add 1  $\mu$ L of isolated gDNA.

Table 1: Volumes for PCR Mixture			
Reagents	Volume for 1X25 μL (μL)	Volume for 75 μL master-mix (μL)	
2X BioMix (contains reaction buffer,	12.5	37.5	
Mg <sup>2+</sup> , dNTPs, <i>Taq</i> polymerase and			
additives)			
515fB forward primer (5 pmol)	0.5	1.5	
806rB reverse primer (5 pmol)	0.5	1.5	
Template DNA	1.0	N/A	
PCR grade water	10.5	31.5	
Total	25.0	72.0	

- 2. Add to thermocycler on set cycle. Thermal cycle programme:
  - A. 94°C, 1 min
  - B. 94°C, 15 s
  - C. 50°C, 45 s
  - D. 72°C, 30 s
  - E. Repeat stages A-D 35 times
  - F. 72°C, 5 min
  - G. 4°C HOLD

**IMPORTANT:** Pool 3 samples (25 μL) into one tube prior to PCR quality control and purification.

## 5: Check quality of amplicons by gel electrophoresis

#### **Precautions and factors to consider:**

- Always wear gloves
- If gel has been prepared earlier, wrap in clingfilm and store in fridge

#### **Equipment and materials:**

- Pipettors and pipette tips
- Eppendorf tubes
- Agarose powder
- Tris-acetate-EDTA buffer
- Ethidium bromide
- 5X loading dye
- 12 x 14 cm horizontal gel electrophoresis device

- 1. Measure 1.5 g of agarose powder.
- 2. Cast a ~100 mL **1.5% agarose gel** by mixing 1.5 g of agarose powder with 2 mL of 50X TAE buffer and 98 mL of distilled water in a microwavable flask.
- 3. Microwave flask in pulses with occasional swirling, for 1-3 min until the agarose is completely dissolved. **CAUTION:** do not over boil the solution.
- 4. Allow agarose solution to cool to  $\sim 50^{\circ}$ C (hands can comfortably hold flask).
- 5. Add 0.5 µL of 1% stock ethidium bromide to a final concentration of ~0.5 µg/mL.
- 6. Pour the agarose into a tray with the narrow well comb in its fixed position. Allow it to set at room temperature for 20-30 min (until the gel has completely solidified).
- 7. Once solidified, place the agarose gel into the electrophoresis unit/gel box.
- 8. Make 50 mL of 1X TAE buffer.
- 9. Carefully pour the buffer solution over the gel so that it covers the entire gel.
- 10. Transfer 1  $\mu$ l of amplicon sample(s) into clean labelled tube(s) and bring the total volume up to 4  $\mu$ L with 1X TE Buffer, pH 8.0.
- 11. Add 1 µL of 5X loading dye. The amount of loading dye should remain consistent among all samples to

maintain even DNA migration on the gel.

- 12. Mix by pipetting up and down a few times.
- 13. Load onto gel according to the format listed below (where n is number of samples to be run):
  - > Well 1: 10 μl of 1kB + ladder (Invitrogen), 100 ng/μL.
  - ➤ Well 2 to n: Amplicon samples.
- 14. Run gel for **30 min** at **120V** in 1X TAE buffer. **CAUTION**: Do not run for a long duration to prevent DNA running off the gel.
- 15. Switch off power and disconnect the electrodes from the power source.
- 16. Carefully remove the gel from gel box.
- 17. Using a device that has UV lights, take an image and visualise the DNA.

**Note:** Analyse the gel photo for amplicon presence and its quality; the expected size is 384 bp. Using the standard molecular marker, make an educated estimate of the amount of product in the amplicon band, from this, establish a rough concentration of the amplicon.

## 6: Purify amplicon using QIAquick PCR Purification Kit (Qiagen, Hilden, Germany)

#### **Precautions and factors to consider:**

- Always wear gloves
- Label all tubes and filters
- SpinClean Buffer contains alcohol, so be careful not to wipe off the labels accidentally

## **Equipment and materials:**

- PCR product tubes 25 μL
- SpinBind 125 μL per prep
- SpinClean Buffer 10 mM Tris, 50 μL per prep
- Spin Filter unit
- 2 mL collection tube
- Microcentrifuge at 10,000 g
- Pipettors (volumes of  $50 \mu L 500 \mu L$  required)

- 1. Shake to mix the **SpinBind** before use.
- 2. Add 5X volumes of **SpinBind** to the PCR reaction mixture. For example, add 500  $\mu$ L to 100  $\mu$ L PCR reaction. Or, pool PCR products and then add the binding buffer.
- 3. Mix well by pipetting.
- 4. Label all the tubes and filters.
- 5. Transfer PCR/SpinBind mixture to a **Spin Filter unit**.
- 6. Centrifuge for 10-30 s at a minimum of 10,000 g (~13,000 rpm) in a table-top microcentrifuge.
- 7. Remove **Spin Filter basket** and discard the liquid flow-through from the tube by decanting.
- 8. Replace the Spin Filter basket in the same tube.
- 9. Add 300 µL **SpinClean Buffer** to the labelled Spin Filter.
- 10. Centrifuge for 10-30 s at a minimum of 10,000 g.
- 11. Remove **Spin Filter basket** and discard liquid flow-through by decanting.
- 12. Replace the Spin Filter basket back into the same tube.
- 13. Centrifuge for 10-30 s at a minimum of 10,000 g.
- 14. Transfer **Spin Filter** to a clean 2 mL collection tube.
- 15. Add 50 μL of **Elution Buffer** (10 mM Tris) solution provided or sterile water directly onto the centre of the white **Spin Filter** membrane.
- 16. Centrifuge 10-30 s at a minimum of 10,000 g.
- 17. Remove **Spin Filter basket** from the 2 mL collection tube.
- 18. Seal tube and store DNA at -20°C.

# 7: Quantify dsDNA using SpectraMax® Quant<sup>TM</sup> AccuClear<sup>TM</sup> Nano dsDNA Assay – Explorer Kit (Molecular Devices, LLC, California, US)

## **Equipment and materials:**

- Fluorescence microplate reader
- SoftMax Pro Software
- 96-well microplate (solid black)
- Vortex
- De-ionised water (buffer solution)
- dsDNA standard
- 100X AccuClear Nano dye

#### Workflow:

- ➤ Add sample and reagent
- > Cover and mix
- > Fluorescence measurement
- Data analysis

- 1. Warm reagents to room temperature before use.
- 2. Vortex reagents to minimise reagent loss in the cap, then centrifuge vials briefly.
- 3. Dilute 20X AccuClear Nano Buffer to 1X with  $dH_2O$  (3388  $\mu L$  required, calculating for 3400  $\mu L$ : 170  $\mu L$  20X buffer in 3230  $\mu L$   $dH_2O$ ).
- 4. Dilute 100X dye (1:100) in 1X **AccuClear Nano Buffer** in a plastic container (**do not use glass**), and vortex to mix well (26 μL dye in 2600μL buffer).
- 5. Pipette 200 μL of working solution into each microplate well (total required is 2600 μL).
- 6. Prepare a set of dsDNA standards by diluting the 25 ng/μL dsDNA standard in the 1X AccuClear Nano Buffer as indicated in Table 2 below.
- 7. Pipette 10 µL of SpectraMax Quant AccuClear Nano dsDNA standards into separate microplate wells and mix.
- 8. Incubate microplate at room temperature for 5-10 min in the dark.
- 9. Measure fluorescence using a fluorescence microplate reader (SoftMax® Pro Software; excitation = 468 nm; emission = 507 nm).
- 10. Use standard curve (linear regression automatically computed by the plate-reader; a Molecular Devices SpectraMax i3) to calculate the concentrations of our DNA samples by using the RFU value from the reader. Note: If fluorescence of unknown sample exceeds the linear range, further dilute the sample and use 10  $\mu$ L of diluted sample to perform the assay.

Table 2: Preparation of dsDNA Standards			
Standard	Final Concentration	Standard Volume	1X AccuClear Nano Buffer Volume
A	25 ng/μL	100 μL of 25 ng/μL AccuClear Nano Standard	None
В	10 ng/μL	40 μL of 25 ng/μL AccuClear Nano Standard	60 μL
С	3 ng/μL	12 μL of 25 ng/μL AccuClear Nano Standard	88 μL
D	1 ng/μL	10 μL of 10 ng/μL AccuClear Nano Standard	90 μL
Е	1.3 ng/μL	10 μL of 3 ng/μL AccuClear Nano Standard	90 μL

## 8: Pool PCR products for sequencing and final analysis

## 8 a) Pooling for a final DNA concentration of 10 nM

Use concentration results from dsDNA assay to adjust the **final DNA concentration to 10 nM** using the following calculation: (where: MW = number of bases [384] x average Mr of bases [660 gmol<sup>-1</sup> per base pair] = 253,440 gmol<sup>-1</sup>; variables x, y, and z are unknown):

nM = ng/
$$\mu$$
L  
10 nM=  $\frac{ng/\mu L}{253,440} \times (1 \times 10^6)$ 

$$\therefore$$
 ng/ $\mu$ L =  $x$  ng/ $\mu$ L

$$n = \frac{m}{Mr} = \frac{ng/\mu L}{253,440} = \left(\frac{ng/\mu L}{\frac{253,440gmol^{-1}}{10^{-6}\mu L}}\right) = y \ mol^{-1} = z \ nM$$

## 8 b) DNA sent for sequencing using Illumina MiSeq

The Illumina MiSeq Reagent Kits v2 for 500 cycles (catalog ID: MS-102-2003) was used.

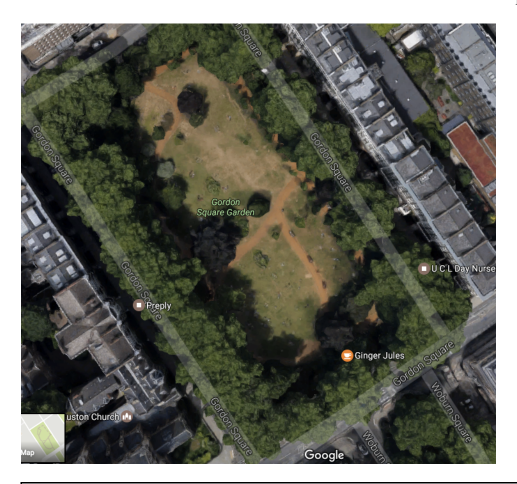
This gives 250 cycles per read (each cycle is a nucleotide).

### 8 c) Analysis by computational methods

Use software, for example QIIME v1.9.1 (or current version).

#### **Appendix**

Appendix 1: Map overview of Gordon Square Garden generated using Google Maps.



#### **Appendix 2: Sequences Of Primers Used**

#### Forward:

515fB PCR Primer Sequence:

AATGATACGGCGACCACCGAGATCTACACGCT XXXXXXXXXXXX TATGGTAATT GT GTGYCAGCMGCCGCGGTAA

(where: Y is C/T; M is A/C)

Description: 5' Illumina adaptor, GoLay barcode, Forward primer pad, Forward primer linker, Forward primer

### Reverse:

806rB PCR Primer Sequence:

CAAGCAGAAGACGCATACGAGAT AGTCAGTCAG CC GGACTACNVGGGTWTCTAAT

(where: N is any; W is A/T)

Description: Reverse Complement of 3' Illumina adaptor, Reverse primer pad, Reverse primer linker, Reverse primer (806rB)