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18S rRNA-Gene Metabarcoding Library Prep: Dual-PCR Method



In 1 collection

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Protocol status: Working We use this protocol and it's

working

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Keywords: 18S rRNA, eDNA, eukaryote diversity, Illumina Sequencing

Abstract

This protocol is used for eDNA metabarcoding of the 18S SSU rRNA Gene (Balzano et al 2015) using Pair-End Illumina Miseq. Sequencing. As part of the Hakai Institute Ocean Observing Program, biomolecular samples have been collected weekly, from 0 m to near bottom (260 m), to genetically characterize plankton communities in the Northern Salish Sea since 2015. This protocol is developed to provide taxonomic annotations of Eukaryote Nuclear DNA.





Guidelines

MIOP: Minimum Information about an Omics Protocol

MIOP Term	Value
analyses	Amplicon Sequencing, 18S
audience	scientists
broad-scale environmental context	marine biome ENVO_00000447
creator	Rute Carvalho
environmental medium	sea water [ENVO:00002149]
geographic location	North Pacific Ocean [GAZ:00002410]
hasVersion	1
issued	2024
language	en
license	CC BY 4.0
local environmental context	coastal sea water [ENVO: 00002150]
materials required	Sterile workbench, Thermo Cykler, MiSeq, Gel Electrophoresis syste, Qbit, Bioanalyzer
maturity level	Mature
methodology category	Omics Analysis
personnel required	1
project	Biomolecular surveys of marine biodiversity in the Northern Salish Sea, BC
publisher	Hakai Institute, Ocean Observing Program
purpose DNA metabarcoding	
skills required	sterile technique pipetting skills
target	Eukaryote Nuclear DNA
time required	3-5 days

AUTHORS

PREPARED BY All authors known to have contributed to the preparation of this protocol, including those who filled in the template	AFFILIATION	ORCID (visit https://orcid.org/ to register)	DAT
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RELATED PROTOCOLS

PROTOCOL NAME AND LINK	ISSUER / AUTHOR	RELEASE DATE This is the date corresponding to the version listed to the left
Content Cell	Content Cell	yyyy-mm-dd
Content Cell	Content Cell	yyyy-mm-dd

This is a list of other protocols which should be known to users of this protocol. Please include the link to each related protocol.

ACRONYMS AND ABBREVIATIONS

	ACRONYM / ABBREVIATION	DEFINITION	
Г	Content Cell	Content Cell	

GLOSSARY

	SPECIALISED TERM	DEFINITION	
Г	Content Cell	Content Cell	
	Content Cell	Content Cell	

BACKGROUND

This protocol is used for eDNA metabarcoding of the 18S SSU rRNA Gene (Balzano et al 2015) using Pair-End Illumina MiSeq Sequencing.

CITATION

Sergio Balzano, Elsa Abs, Sophie C. Leterme (2015). Protist diversity along a salinity gradient in a coastal lagoon. Aquatic Microbial Ecology.

https://doi.org/10.3354/ame01740

Method description and rationale

Assuming extracted DNA as starting material, this protocol includes the following steps:

- 1. First PCR: Triplicate locus-specific amplification of the 320 bp ling "Leray fragment" of the COI gene.
- 2. First PCR product purification (using magnetic beads)
- 3. Second PCR: Sample indexing using Nextera V2 indexing primers
- 4. Second PCR product purification (using magnetic beads)
- 5. Quantification and Pooling
- 6. Quality control
- 7. Pair End Sequencing on Illumina MiSeq V3 2*300 bp

Due to the risk of cross contamination, it is pivotal to separate work with amplified PCR products from pre-PCR steps. We preform pre-PCR steps (including DNA extractions) in separate clean rooms on surfaces steralized with hydrogen peroxide (PreEmpt) and UV.

Spatial coverage and environment(s) of relevance

As part of the Hakai Institute Ocean Observing Program, biomolecular samples have been collected weekly, from 0 to near bottom (260 m), to genetically characterize plankton communities in the Northern Salish Sea since 2015, developing a climatology from which we can begin uncover the physical, chemical and biological drivers of community and functional change in the dynamic coastal waters of coastal British Columbia. This protocol is developed to give a species-level resolution of **marine eukaryotes**.

Personnel Required

1 Technician

Safety

Identify hazards associated with the procedure and specify protective equipment and safety training required to safely execute the procedure!

Training requirements

Serile work technique, pipetting skills, PCR, gel electrophoresis.

Time needed to execute the procedure

This protocol may take several days to complete depending on sample size.

Materials

Equipment

- pre-PCR and post-PCR separated workspaces
- Thermocycler (1 or 3)
- Gel electrophoresis equipment
- Qubit or plate reader
- Magnetic plate
- BioAnalyzer
- Real-Time PCR
- Illumina MiSeq

Protocol materials

PhiX Control v3 Illumina, Inc. Catalog #FC-110-3001 Step 33 MiSeq v3 (150 cycle) Kit Illumina, Inc. Catalog #MS-102-3001 Step 33 🔀 Qubit dsDNA Broad Range assay kit (500 assays) Invitrogen - Thermo Fisher Catalog #Q32853 | Step 28 🔯 Quant-iT dsDNA Pico Green assay kit (Invitrogen) Life Technologies Catalog #P7589 Step 28 XX NEBNext Library Quant Kit for Illumina - 100 rxns New England Biolabs Catalog #E7630S Step 32 Gel Red Nucleic Acid Gel Stain Biotium Catalog ##41003 In 2 steps Molecular Biology Grade Water Corning Catalog #46-000-CV Step 6

Before start

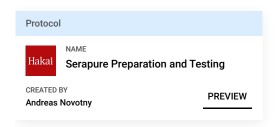
Read Minimum Information about an Omics Protocol (MIOP) and other recommendations under the "Guidelines" tab.

🔀 Bioanalyzer chips and reagents (DNA 1000) Agilent Technologies Catalog #5067-1504 Step 32



Preparations

- Ensure that the laboratory is appropriately configured and that staff has appropriate training. See "Guidelines" for more information. Pay attention to the separation of pre and post-PCR spaces and equipment.
- 2 Ensure that all reagents are aliquoted in appropriate amounts, and stored according to manufacturers' recommendations. Never pipet directly from reagent stocks.
- 3 Prepare the SPRI beads' working solution, and test their efficiency following this protocol.



- 4 Prepare primer working stocks (10μM) for both the first and second PCR steps. Here we use Nextera V2 Kit Sets A, B, C, and D. We advise preparing the indexing primers on 96-well plates according to this configuration:
 - Indexes_plate.xlsx 38KB
- We advise adding aliquots of the extracted DNA to a 96-Well PCR plate to facilitate the setup of the PCR reaction. This metadata template will help keep track of the samples, and if indexes are configured as described above, also the identity of sample indexes.

Triplicate PCR Amplification (1st PCR)

6 Preparations



Note

- Prepare PCR reactions in a clean working space (such as a biosafety cabinet) dedicated to pre-PCR tasks only.
- 2. Do not need to Qubit DNA samples before starting, only do it if the reaction does not work.
- 3. Use samples diluted 1:10 (1 μ l DNA in 9 μ l Nuclease-Free Water)
- 4. Test at least 8 samples before doing a batch/plate.
- 5. Include a negative control, an extraction blank (if you have it), and a positive control.
- 6. After testing, perform the PCR for all of the samples in triplicates.

Reagents:

- Molecular Biology Grade Water Corning Catalog #46-000-CV (Or equal)
- X Taq FroggaMix Froggabio Catalog #FBTAQM96
- 🔀 BSA-Molecular Biology Grade 12 mg New England Biolabs Catalog #B9000S
- Froggarose LE Froggabio Catalog #A87-500G
- ⊠ Gel Red Nucleic Acid Gel Stain Biotium Catalog ##41003
- Custom-designed primers (Balzano et al 2015) including:

	PCR Primer Name	Direction	Sequence (5' -> 3')
Γ	Balzano_565F_overhang	forward	TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGCCAGCASCYGCGGTAATTCC
	Balzano_981R_overhang	reverse	GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGACTTTCGTTCTTGATYRR



UV for 30 minutes the following:

- 96-well PCR plates (or 8-strip tubes)
- Sharpie
- Pipette tips
- Multichannel pipettes
- Pipettes
- Sterile Nuclease-Free Water

Thaw Taq, BSA, Primers, and nuclease-free water. Keep them in a cooling microcentrifuge tube

8 PCR reactions are carried out in triplicate 25µl reactions:

	Reagent	Volume (µI)
	Sterile Nuclease-Free water	7.3
Г	Forward primer (10µM)	0.6
Г	Reverse Primer (10µM)	0.6
Г	BSA (10mg/ml)	2
Г	2XTaq	12.5
	DNA (1-10 ng)	2
	TOTAL	25

30m 1

> 2h Z A

Seal the 96-well plates and transfer them to thermocyclers.

Amplified PCR products should never come in contact with equipment used for non-

From this point, no samples will reenter the pre-PCR working space.

	PCR step	Temperature	Duration	Repetition
Γ	denaturation	98°C	1 minutes	
	denaturation	98°C	30 seconds	
Г	annealing	53°C	30 seconds	
Γ	extension	72°C	45 seconds	
Γ	GO TO step 2			10 times
Г	denaturation	98°C	10 seconds	
Г	annealing	50°C	30 seconds	
Γ	extension	72°C	10 seconds	
Г	GO TO step 6			19 times
Г	final extension	72°C	2 minutes	
	HOLD	12°C	HOLD	

10 Run a subset of the PCR product (5 μ I) on a 1.5% agarose gel to check the size of the amplicons and the success of the amplification.



Expected result

If any additional bands appear that are not the desired product's size, increase the PCR's annealing temperature or perform additional purification steps.

Purification of first PCR product using SPRI beads

Preparations



Note

Prepare the purification in the post-PCR working space.

Size selection can be achieved using different ratios of magnetic beads to sample. A rate of bead to a sample of 0.8-1.5 will efficiently purify the amplicons away from primer dimers and allow the selection of fragments larger than 200 bp.

Materials

- Serapure SPRI beads. If not already prepared: <u>⇒ go to step #3</u>
- Magnetic 96-well plate stand
- Anhydrous Ethanol to make a fresh 80% ethanol solution
- Molecular grade water

UV for 30 minutes the following:

- 96-well PCR plates (or 8-strip tubes)
- Sharpie
- Pipette tips
- Multichannel pipettes
- Pipettes
- Sterile Nuclease-Free Water

Remove the magnetic beads from the fridge (allow 30 min to reach room temperature).

- Vortex the beads before use.
 - Add 16 μl beads to 20 μl of PCR product to obtain a ratio of 0.8.
 - Pipette up and down ten times (or until the solution is well mixed you will see that the color changes).
 - Spin tubes down to remove drops from the walls.
- 13 Incubate at room temperature without shaking for 5 min.

Then, place the plate on the magnetic stand until the supernatant has cleared (~ 3 min).

- 14 Remove the supernatant with a multichannel pipette, ensuring to not disturb the beads.
- 15 With the samples on the magnetic rack, wash the beads by adding 180 μ l of freshly prepared 80% ethanol and incubate for 30s. Carefully remove the supernatant without disturbing the
- 16 Repeat the washing step =5 go to step #15
- 17 Remove all residual ethanol using a pipette and air dry, leaving the samples on the magnetic stand (~ 5 min*).

Note

*This depends on the type of the magnetic rack – the O-ring magnet dries faster than the side magnet. Keep an eye on the beads and do not over-dry. Otherwise, you will not get an efficient DNA recovery.

- 18 Remove the plate from the magnetic stand and 40 µl of nuclease-free water for elution. Gently pipet up and down ten times to resuspend the beads. Incubate the plate at room temperature for 5 min.
- 19 Place the plate back on the magnetic rack for at least 5 min or until the supernatant is cleared.
- 20 Carefully transfer 30 µl of the clear supernatant to a new plate. Seal the plate.





5m

10m

10m

5m

A

5m

5m



Name the plate: Project, [Gene_name], PCR 1, Post-Purification Plate #, Date, Initials. Samples can be stored at -20°C for up to 7 days.



(IF this is the cleanup of the second PCR product = go to step #28)

Indexing PCR amplification (2nd PCR)

22 Preparations

Reagents:

- 🔀 Taq FroggaMix Froggabio Catalog #FBTAQM96
- Molecular Biology Grade Water Corning Catalog #46-000-CV
- X Froggarose LE Froggabio Catalog #A87-500G
- ⊠ Gel Red Nucleic Acid Gel Stain Biotium Catalog ##41003
- i5 and i7 index plates (10 μM) If not already prepared:

 go to step #4

PCR Primer Name	Direction	Sequence (5' -> 3')
Nextera V2 Index1	forward	CAAGCAGAAGACGGCATACGAGAT[i7]GTCTCGTGGGCTCGG
Nextera V2 Index 2	reverse	AATGATACGGCGACCACCGAGATCTACAC[i5]TCGTCGGCAGCGTC

UV for 30 minutes the following:

- 96-well PCR plates (or 8-strip tubes)
- Sharpie
- Pipette tips
- Multichannel pipettes
- Pipettes
- Sterile Nuclease-Free Water

Thaw Taq, i5 and i7 indexes, and nuclease-free water. Keep them in the IsoFreeze microcentrifuge tube rack.

23 Dilute the cleaned-up PCR (1:10) with sterile nuclease-free water.



24 Prepare PCR reaction in 25µl reactions:

	Reagent	Volume (µI)
Г	Sterile Nuclease-Free water	5
Г	Forward primer (10µM)	2.5
Г	Reverse Primer (10µM)	2.5
Г	2XTaq	12.5
Г	DNA (1-10 ng)	2.5
	TOTAL	25

Seal the 96-well plates and transfer them to thermocyclers.

	PCR step	Temperature	Duration	Repetition
Г	denaturation	95°C	3 minutes	
Г	denaturation	95°C	30 seconds	
Г	annealing	55°C	30 seconds	
Г	extension	72°C	30 seconds	
Г	GO TO step 2			7X
Г	final extension	72°C	5 minutes	
	HOLD	12°C	HOLD	



6 Run a subset of the PCR product (5µI) on a 1.5% agarose gel to check the size of the amplicons and the success of the amplification.



Note

If any additional bands appear that are not the size of the desired product, additional purification steps need to be carried out.

Purification of indexed libraries (Second bead cleanup)

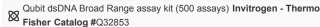
27 Repeat the Ampure XP bead cleanup for all the indexed libraries.





Quantification and pooling, and quality control

Use a fluorometric quantification method that uses dsDNA dyes to measure the concentration of your libraries (Qubit or plate reader). If using Qubit, give preference to the broad range kit if you visualize a strong band in the gel:



OR

🔯 Quant-iT dsDNA Pico Green assay kit (Invitrogen) Life Technologies Catalog #P7589

Expected result

Samples will have approximately similar concentrations (usually). Re-check samples that showed very high or low concentrations on Qubit/plate reader quantification.

29 Calculate sample volume to have a final amount of 10-40 ng. This amount may vary depending on the overall quantification. For example, if on average the concentration of your samples is about 3 ng/μl and you have 20 μl of product, you can calculate the volume to make up to 60 ng per sample.

Note

Check the final volume that you will get after pooling – sometimes you will end up with 2 mL or more. Then use the proper Eppendorf tube for pooling (1.5, 2.0, or 5 mL).

30 Measure the final library pool concentration on Qubit using

X Qubit dsDNA HS Assay kit Thermo Fisher Scientific Catalog #Q32854

31 Label tube: [Gene_name], [Project_Name], Pooled Amplicons. Date, Initials, pool concentration.

Sequencing parameters

32 Library fragment size (BP) is determined using

Bioanalyzer chips and reagents (DNA 1000) **Agilent Technologies Catalog #**5067-1504

Molarity of thefinal pool is assessed using

 \bowtie NEBNext Library Quant Kit for Illumina - 100 rxns New England Biolabs Catalog #E7630S

33 COI libraries are sequenced an a MiSeq instrument using:



Citations

Sergio Balzano, Elsa Abs, Sophie C. Leterme. Protist diversity along a salinity gradient in a coastal lagoon $\underline{\text{https://doi.org/10.3354/ame01740}}$