



Mar 29, 2024

Version 1

Metabarcoding Fecal Swabs or Stomach Contents for Fish and Crustaceans using 2-PCR protocol and Illumina MiSeq V.1

DOI

dx.doi.org/10.17504/protocols.io.ewov1qxokgr2/v1

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Protocol status: Working

We use this protocol and it's working

Created: November 18, 2023

Last Modified: March 29, 2024

Protocol Integer ID: 91145

Keywords: rrna gene of crustacean, stomach contents for fish, mitochondrial 12s rrna gene, rrna gene, metabarcoding fecal swab, mitochondrial16, 170bp region of the mitochondrial16, crustacean, fish, pcr protocol, pcr, resulting pcr product, metabarcoding, pcr product

Abstract

This protocol describes a method to metabarcode a 170bp region of the mitochondrial16S rRNA gene of crustaceans and a 163-185bp region of the mitochondrial 12S rRNA gene of fishes. These regions are subjected to PCR separately in multiple replicates and the resulting PCR products are pooled by sample and then indexed for sequencing on an Illumina MiSeq platform.

Image Attribution

Haley Capone

Guidelines

The PCR conditions described here are different from the PCR conditions described by Miya et al., and Berry et al. in their respective publications introducing the primers used here. This difference is due to the use of the Takara High Fidelity PCR EcoDry Premix in this protocol.

Materials

- 96-well PCR plates
- Adhesive foil PCR plate covers
- 1.5mL tubes
- Glenn et al. Adapterama I iNext indexing primers A-H and 1-12.
- PCR machine
- Equipment to run gels
- optionally: equipment for fluorometric quantification

Equipment	
96-well Magnetic Rack Separator	NAME
Magnetic Rack Separator	TYPE
Sergi Lab Supplies	BRAND
B08134P9RT	SKU
https://www.amazon.com/Magnetic-Separator-Protein-Purification-Format/dp/B08134P9RT/ref=asc_df_B08134P9RT/?tag=&linkCode=df0&hvadid=416872221972&hvpos=&hvnetw=g&hvrnd=12953200023550024012&hvpon e=&hvptwo=&hvqmt=&hvdev=c&hvdvcmdl=&hvlocint=&hvlocphy=903024	L I N K

Equipment

Magnetic Rack for for 1.5 mL Tubes	NAME
Magnetic Rack for DNA, RNA Purification; for 1.5 mL centrifuge Tubes	TYPE
Sergi Lab Supplies	BRAND
B0BZWXZMZ2	SKU
https://www.amazon.com/Magnetic-Rack-Purification-centrifuge-Tubes/dp/B0BZWXZMZ2/ref=asc_df_B0BZWXZMZ2/?tag=hyprod-20&linkCode=df0&hvadid=652498086131&hvpos=&hvnetw=g&hvrand=6716034042841103246&hvpone=&hvptwo=&hvmmt=&hvdev=c&hvdvcml=&hvlocint=&hvlocphy=9	LINK

Protocol materials

- ⊗ Nuclease-free water Integrated DNA Technologies, Inc. (IDT) Catalog #11-05-01-14
- ⊗ Nuclease-free water Integrated DNA Technologies, Inc. (IDT) Catalog #11-05-01-14
- ⊗ MiFish-F Integrated DNA Technologies, Inc. (IDT) Catalog #custom
- ⊗ MiFish-R Integrated DNA Technologies, Inc. (IDT) Catalog #custom
- ⊗ Crustacean16S-F Integrated DNA Technologies, Inc. (IDT) Catalog #custom
- ⊗ Crustacean16S-R Integrated DNA Technologies, Inc. (IDT) Catalog #custom
- ⊗ Crustacean16S-F Integrated DNA Technologies, Inc. (IDT) Catalog #custom
- ⊗ Crustacean16S-R Integrated DNA Technologies, Inc. (IDT) Catalog #custom
- ⊗ Nuclease-free water Integrated DNA Technologies, Inc. (IDT) Catalog #11-05-01-14
- ⊗ Takara High Fidelity PCR EcoDry Premix Takara Bio Inc. Catalog #639280
- ⊗ Takara High Fidelity PCR EcoDry Premix Takara Bio Inc. Catalog #639280
- ⊗ Agencourt AMPure XP Beckman Coulter Catalog #A63880
- ⊗ Buffer EB Qiagen Catalog #19086
- ⊗ 2x Kapa HiFi Hotstart Readymix Kapa Biosystems Catalog #KK2602
- ⊗ Nuclease-free water Integrated DNA Technologies, Inc. (IDT) Catalog #11-05-01-14
- ⊗ Buffer EB Qiagen Catalog #19086

Troubleshooting


Before start

Work in a pre-PCR lab, as separated as possible from post-PCR products.


Clean work area with 10% bleach solution before beginning work for the day, then change gloves so that no bleach carryover to your samples or reactions occurs.

Prepare Primers


- 1 Order metabarcoding primers with diversity spacers and Illumina overhang sequences (Illumina, 2013):

 MiFish-F Integrated DNA Technologies, Inc. (IDT) Catalog #custom (Miya et al., 2015):

TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGNNNGTCGGTAAACTCGTGCCAGC

 MiFish-R Integrated DNA Technologies, Inc. (IDT) Catalog #custom (Miya et al., 2015):

GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGNNNNCATAGTGGGGTATCTAATCCCAGTTTG

 Crustacean16S-F Integrated DNA Technologies, Inc. (IDT) Catalog #custom Berry et al., 2017):



TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGNNNGGACGATAAGACCCTATA


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

(Berry et al., 2017):


GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGNNNATTACGCTGTTATCCCTAAAG



We got ours from <https://www.idtdna.com/> as custom oligos at 25nm scale, with standard desalting.

- 2 Reconstitute primers to  100 micromolar (μM) stock solutions by adding  40 μL of

 Nuclease-free water Integrated DNA Technologies, Inc. (IDT) Catalog #11-05-01-14

- 3 Make  5 micromolar (μM) working solutions of each primer by adding  95 μL of









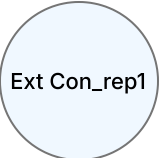































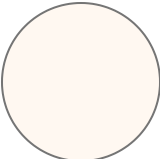
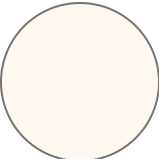


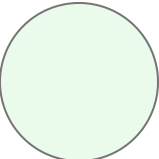





 Nuclease-free water Integrated DNA Technologies, Inc. (IDT) Catalog #11-05-01-14

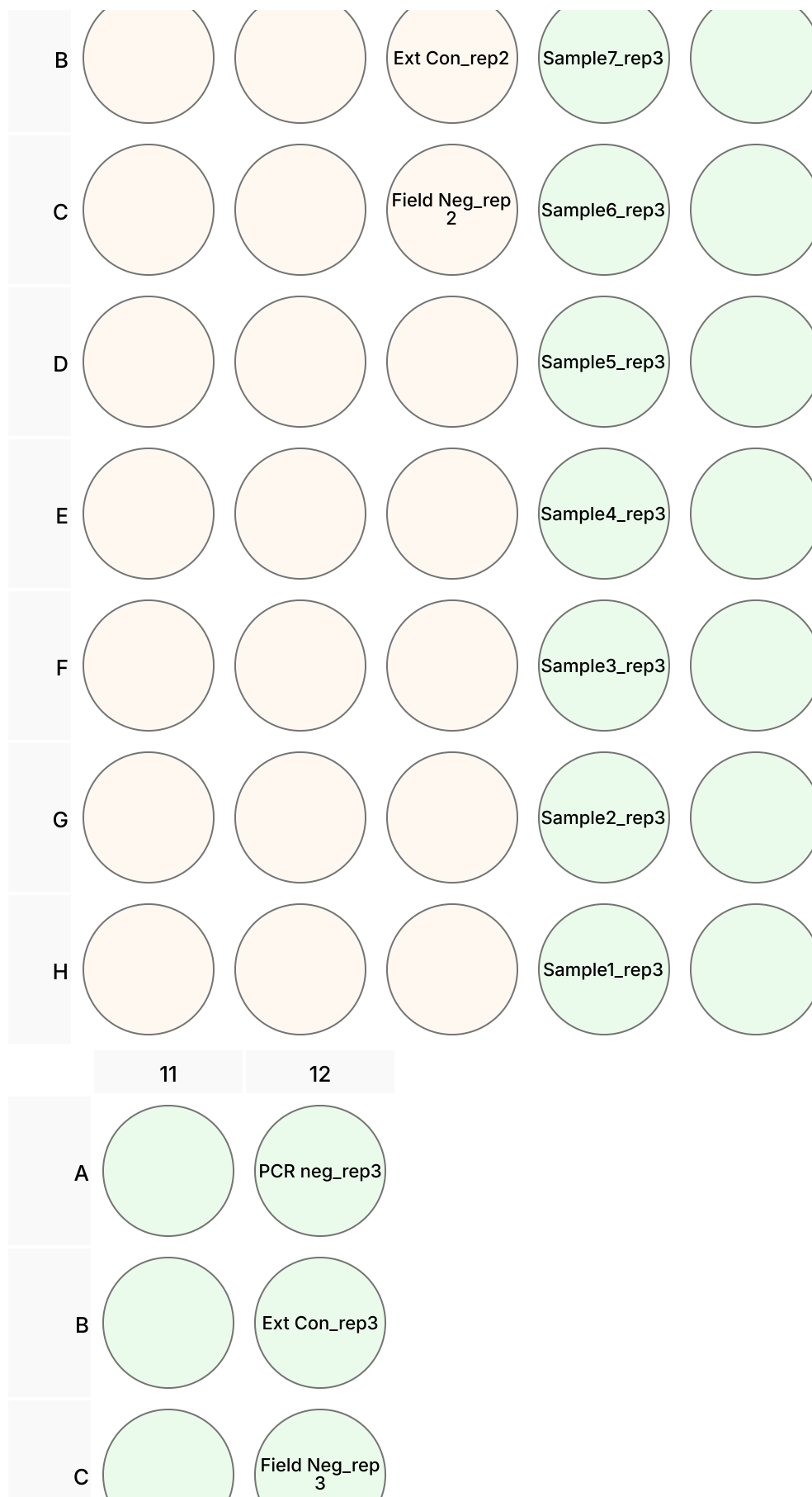
and  5 μL of primer stock solution for each  100 μL of primer that you intend to use within the next week or so.



Create Plate Map

- 4 Determine which sample will go into each well. This should be the same for each primer set and each replicate. Include at least one extraction control (you can combine aliquots of the extraction controls from each round of DNA extraction into one tube, and use that as your single extraction control), and include a PCR negative control for each plate of PCR. See example below of 21 samples, a field negative sample, a combined extraction control, and a PCR negative.




	1	2	3	4	5
A					
B					
C					
D					
E					
F					
G					
H					
	6	7	8	9	10
A					
					














- 4.1 Do not mix sample types between invasively sampled methods (fecal swabs, or stomach contents) and non-invasively sampled methods (eDNA from water or sediment) in the same PCR procedure. And don't plan to sequence both types in the same sequencing run with the combinatorial indexing scheme used here. The potential for contamination of the lower quantity eDNA samples by the higher quantity fDNA samples is too high.

MiFish Takara PCR Recipe

- 5 Add  24 μL of your MiFish metabarcoding mastermix to each well of  Takara High Fidelity PCR EcoDry Premix **Takara Bio Inc. Catalog #639280**
- 5.1 Add  1 μL DNA extracted from stomach contents or fecal swabs.
- 5.2 Mix and stir together with pipette tip, swirling to make sure the liquid is in the bottom, and bringing any bubbles to the surface of each reaction.
- 5.3 Cap each row of reaction tightly before beginning any other PCR reaction in the same room.

MiFish Takara PCR Conditions

- 6  95 °C for  00:01:00
- 35 cycles of:
-  95 °C for  00:00:30
-  66 °C for  00:01:00
- followed by:
-  68 °C for  00:01:00
- Hold at  4 °C

3m 30s

Crustacean_16S Takara PCR Recipe

- 7 Make your Crustacean_16S Mastermix:

For each **PCR replicate of each sample** you intend to process (+10% overage), mix:



2 µL

[M] 5 micromolar (µM)



Crustacean16S-F Integrated DNA Technologies, Inc. (IDT) Catalog #custom

2 µL

[M] 5 micromolar (µM)



Crustacean16S-R Integrated DNA Technologies, Inc. (IDT) Catalog #custom

20 µL



Nuclease-free water Integrated DNA Technologies, Inc. (IDT) Catalog #11-05-01-14

For a full plate of 96 reactions, multiply 105.6*the per-sample volumes in the recipe to make the mastermix.

8 Add 24 µL of your Crustacean_16S metabarcoding mastermix to each well of



Takara High Fidelity PCR EcoDry Premix Takara Bio Inc. Catalog #639280

8.1 Add 1 µL DNA extract

8.2 Mix and stir together with pipette tip, swirling to make sure the liquid is in the bottom, and bringing any bubbles to the surface of each reaction.

8.3 Cap each row of reaction tightly before beginning any other PCR reaction in the same room.

Crustacean_16S Takara PCR Conditions

4m

9 95 °C for 00:01:00

35 cycles of:



95 °C for 00:00:30



50 °C for 00:01:00



68 °C for 00:00:30

followed by:



68 °C for 00:01:00

then hold at 4 °C

4m




Visualize PCR Products

- 10 Make a 1.7% to 2% agarose gel and run a representative sample of reactions on it to make sure the PCRs worked, producing bands in the 250-300bp range. Check some PCR negatives to see that they don't have bands. Be very careful opening the PCR plate wells at this point to avoid cross-contamination.

Prepare EtOH for bead cleanup, and bring beads to room temperature

12m 30s

- 11 Get AmpureXP beads out of the refrigerator, and bring to room temp, swirl to mix occasionally, or use a rocking platform.
- 12 Make fresh 80% EtOH so that you will have at least  200 μL of EtOH per well of the combined plate.

- 13 Get 2 sterile DNAase/RNAse free 96-well PCR plates out of their packaging and immediately cover with adhesive foil.

15m




UV clean the plates for  00:15:00

One plate will be for the bead-cleanup steps, and the other will be for the final, cleaned reactions.

Perform a 1.5x bead cleanup with Ampure XP beads.

12m 30s

- 14 in the bead-cleanup plate, do the following steps for one 8-sample row of the plate at a time, pulling back the foil cover for each row after the previous one has been completed.

- 14.1 pipette mix  10 μL combined PCR product with  15 μL  Agencourt AMPure XP **Beckman Coulter Catalog #A63880** .

5m

Incubate  00:05:00 at room temperature.

- 15 After the  00:05:00 incubation, place 96-well plate on a

7m



Equipment

96-well Magnetic Rack Separator

Magnetic Rack Separator

Sergi Lab Supplies

B08134P9RT

<https://www.amazon.com/Magnetic-Separator-Protein-Purification-Format/dp/B08134P9RT?tag=&linkCode=df0&hvadid=416872221972&hvpos=&hvnetw=g&hvrnd=12953200023>

for 00:02:00 or until liquid is clear.

16 remove and discard liquid from the row, being careful not to touch the beads with the pipette or to let the beads dry for more than 30 seconds.


16.1 Add 100 μL of 80%EtOH to each well of beads. Incubate at Room temperature for 00:00:30 30s

16.2 Remove the EtOH, then immediately add another 100 μL of 80% EtOH to the wells, incubate for 00:00:30 Room temperature . 30s


16.3 Remove ALL EtOH, and let the row of beads dry just enough to lose some shine but not enough to start cracking. This should be approximately 00:00:30 to 00:01:00 . 1m 30s


16.4 Remove the plate with cleaned beads from the magnetic plate, and add 30 μL of Buffer EB Qiagen Catalog #19086 to each well of beads, pipette mixing each well thoroughly. Incubate 00:05:00 at Room temperature 5m



16.5 Place back on the magnetic rack for  00:01:00 until liquid is clear again.






1m






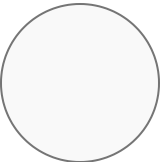
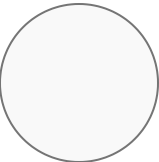
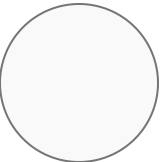
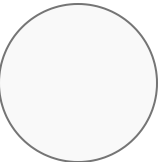
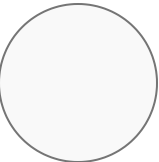
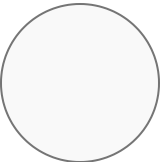
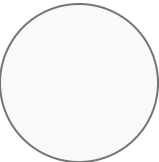
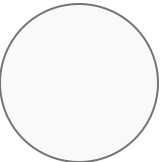
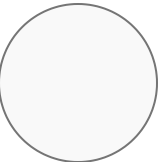
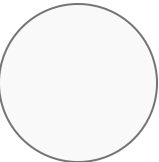
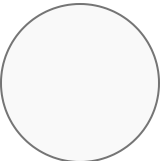
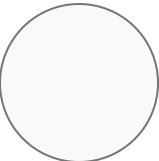
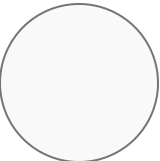
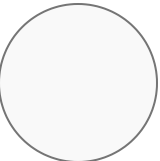
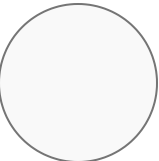
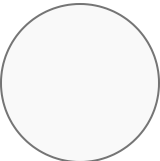
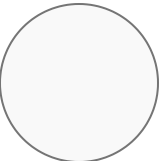
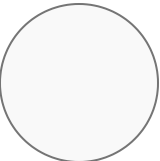
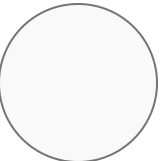
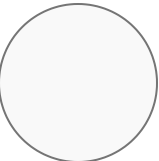
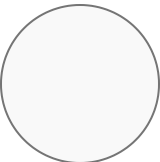
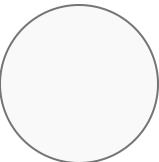
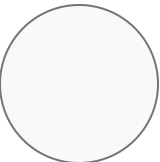
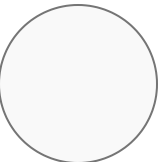
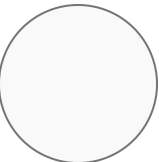
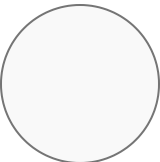
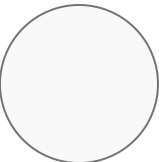
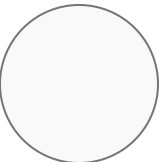
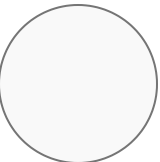
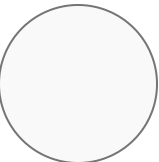
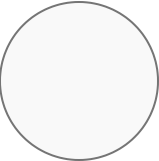
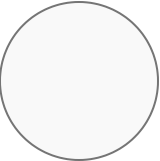
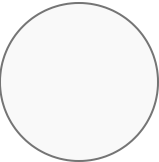
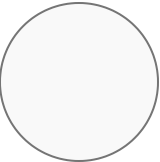


16.6 Roll back the foil on the final cleaned reactions plate for the appropriate row. Remove the  30 μ L clear eluate from the bead-cleanup plate, and place in the appropriate wells of the final cleaned reactions plate. Immediately cover this cleaned PCR product with either 8-strip caps.

16.7 uncover the next row of samples for cleaning and  go to step #14 until all rows are cleaned.

Prepare Indexing PCR

17 Create an indexing plate map and make sure your chosen indexes (iNext indexes) are color balanced if you aren't doing full 96-well plates at one time.

	1	2	3	4	5
A	S8 iNextA-F + iNext1-R	S16 iNextA-F + iNext2-R	etc.	MiFishPCRneg iNextAF+4R	Crust..PCRneg iNextAF+5R
B	S7 iNextB-F + iNext1-R	S15 iNextB-F + iNext2-R			
C	S6 iNextC-F + iNext1-R	S14 iNextC-F + iNext2-R			
D		etc.			
E					
F					
G					
H					
	6	7	8	9	10
A					
					

B					
C					
D					
E					
F					
G					
H					
	11	12			
A					
B					
C					



enn T. C., Ro Nilsen, Troy J. Kieran, Jon G. Sanders, Natalia J. Bayona-
z, ng Pierson, et al. 2019. "Adapterama I: Universal Stubs
D n Un indexed or 147,456 Combinatorially-Indexed Illumina
s (n ext)." 2019 (10). <https://doi.org/10.7717/peerj.7755>.

mental file S10

E v tic micromolar (μM) of each indexing primer you intend

18 Indexing PCR Mastermix Recipe:

6 μL 2x Kapa HiFi Hotstart Readymix **Kapa Biosystems Catalog #KK2602**

2.1 μL

Nuclease-free water **Integrated DNA Technologies, Inc.**
(IDT) Catalog #11-05-01-14

per sample.

Multiply by number of wells *10% as explained above, to create master mix.

19 In a new, clean 96-well plate (UV before use if possible and prepare in a pre-PCR space):

Add 8.1 μL Indexing Mastermix to each well that will be used and add 0.7 μL of
the 5 micromolar (μM) iNext forward indexed primer for each horizontal row of the
plate (8 letters), and 0.7 μL 5 micromolar (μM) of the iNext reverse indexed
primer for each vertical column of the plate (12 numbers) according to the indexing plate
map.

Take the prepared indexing reactions to the post-PCR space to add the cleaned PCR
product.

20 In the post-PCR area, add 2.5uL of cleaned PCR 1 product to their associated wells from the indexing plate map.

Indexing PCR Conditions

21 95 $^{\circ}\text{C}$ 00:03:00

8 cycles of:

98 $^{\circ}\text{C}$ 00:00:20

4m 35s



65 °C

00:00:15

72 °C

00:01:00

then hold 4 °C

Optional gel to check Indexing PCR

- 22 Optional: visualize PCR products in a 1.7-2% gel. Bands should be around 350-400bp.

Combine and Clean all indexed samples from each plate

- 23 Combine 10uL of up to 70 indexed samples (library) into a single 1.5mL tube. If there are more than 70 samples, you will need another tube.
- 24 Multiply the volume of the pooled libraries in each tube by 0.9 to get the volume of Ampure XP beads needed to clean up the reactions.

Perform a 0.9x bead cleanup with Ampure XP beads

28m

- 25 In the 1.5mL tube of pooled libraries, add 0.9x volume of Ampure XP beads and pipette mix well. incubate Room temperature for 00:10:00
- 26 Make enough fresh 80% EtOH to have 2x the total volume of the beads+library pool plus a bit extra.
- 27 Place 1.5mL tube into a magnetic rack

10m

5m



Equipment

Magnetic Rack for for 1.5 mL Tubes

Magnetic Rack for DNA, RNA Purification; for 1.5 mL centrifuge Tubes

Sergi Lab Supplies

BOBZWXZMZ2

<https://www.amazon.com/Magnetic-Rack-Purification-centrifuge-Tubes/dp/B0BZWXZI20&linkCode=df0&hvadid=652498086131&hvpos=&hvnetw=g&hvrnd=671603404284>

and incubate Room temperature for 00:05:00

28 Discard liquid and add an equal or greater volume of 80% EtOH. Incubate

Room temperature for 00:01:00

1m

29 Repeat the ethanol wash a second time [go to step #28](#) , then after the second 80% EtOH wash, remove all EtOH and dry the beads slightly (just until no longer wet-looking but not cracking either).

30 Resuspend beads with 100 μ L Buffer EB [Qiagen Catalog #19086](#) by pipette mixing thoroughly. Incubate Room temperature 00:10:00

10m

31 Place 1.5 mL tube back on magnet rack and wait until liquid is clear, approximately

00:02:00

2m

32 remove 100uL of the clear eluate from the tube with beads while on the magnet and place in a new 1.5mL tube.

33 Quantify with Qubit Broad range and visualize in a gel, then send for sequencing on a lane of MiSeq.



Protocol references

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- 16S Metagenomic Sequencing Library Preparation." 2013. Illumina.
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