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earthEMP 16S Illumina Amplicon Protocol V.2

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Protocol status: Working

Index sequencing primer added to this version

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Abstract

The 16S protocol detailed here is designed to amplify prokaryotes (bacteria and archaea) using paired-end 16S community sequencing on the Illumina platform. Primers 515F-806R target the V4 region of the 16S SSU rRNA.

For running these libraries on the MiSeq and HiSeq, please make sure you read the supplementary methods of Caporaso et al. (2012). You will need to make your sample more complex by adding 5-10% PhiX to your run.

Guidelines

16S amplification primers

The current primers have been modified from the original 515F-806R primer pair (Caporaso et al., 2011) in the following ways:

1. Barcodes are now on the forward primer of the new 515FB-806RB primer pair. This enables the usage of various reverse primer constructs to obtain longer amplicons (tested on 806RB and 926R).
2. Degeneracy was added to both the forward and reverse primers to remove known biases against Crenarchaeota/Thaumarchaeota (515FB, also called 515F-Y, Parada et al., 2016) and the marine and freshwater Alphaproteobacterial clade SAR11 (806RB, Apprill et al., 2015).

The primer sequences without linker, pad, barcode, or adapter are as follows:

- Current, 2015-present (fwd-barcoded: 515FB-806RB):
 -

FWD:GTGYCAGCMGCCGCGTAA; REV:GGACTACNVGGGTWTCTAAT

- Original, pre-2015 (rev-barcoded: 515F-806R):
 -

FWD:GTGCCAGCMGCCGCGTAA; REV:GGACTACHVGGGTWTCTAAT

Note: Studies in the [Qiita](#) database will have

library_construction_protocol

as

515f/806rbc

if original primers or

515fb/806r

if current primers ("bc" stands for barcode).

Ordering primers:

The primer sequences in this protocol are always listed in the 5' -> 3' orientation. This is the orientation that should be used for ordering. See the page [Primer Ordering and Resuspension](#) for more information. Primers and primer constructs were designed by Greg Caporaso (2011, 2012). Modifications to primer degeneracy were done by the labs of Jed Furuham (Parada et al., 2016) and Amy Apprill (Apprill et al., 2015). Forward-barcoded constructs were redesigned by Walters et al. (2016) based upon the original constructs generated by Caporaso et al. (2012).

- [Illumina 16S Primer Constructs \(shorter: 515FB–806RB\)](#)
- [Illumina 16S Primer Constructs \(longer: 515FB–926R\)](#)

515FB forward primer, barcoded:

Field descriptions (space-delimited):

1. 5' Illumina adapter
2. Golay barcode
3. Forward primer pad
4. Forward primer linker
5. Forward primer (515FB)

```
AATGATAACGGCGACCACCGAGATCTACACGCT XXXXXXXXXXXX TATGGTAATT GT GTGYCAGCMGCCGCGTAA
```

806RB reverse primer:

Field descriptions (space-delimited):

1. Reverse complement of 3' Illumina adapter
2. Reverse primer pad
3. Reverse primer linker
4. Reverse primer (806RB)

```
CAAGCAGAAGACGGCATACGAGAT AGTCAGCCAG CC GGACTACNVGGGTWTCTAAT
```

PCR reaction mixture

Reagent	Volume
PCR-grade water	13.0 µL
PCR master mix (2x)	10.0 µL
Forward primer (10 µM)	0.5 µL
Reverse primer (10 µM)	0.5 µL
Template DNA	1.0 µL
Total reaction volume	25.0 µL

Notes:

- PCR-grade water from Sigma (cat. no. W3500) or MoBio (cat. no. 17000-11)
- Platinum Hot Start PCR Master Mix (2x) from ThermoFisher (cat. no. 13000014)
- Final master mix concentration in 1x reaction: 0.8x
- Final primer concentration in 1x reaction: 0.2 µM

Thermocycler conditions

- Primers: 16S V4 515f-806rB
- Amplicon size: ~390 bp
- Cycle times are longer for 384-well thermocyclers.

Temperature	Time, 96-well	Time, 384-well	R
94 °C	3 min	3 min	
94 °C	45 s	60 s	
50 °C	60 s	60 s	
72 °C	90 s	105 s	
72 °C	10 min	10 min	
4 °C	hold	hold	

16S sequencing primers

Read 1 sequencing primer:

Field descriptions (space-delimited):

1. Forward primer pad
2. Forward primer linker
3. Forward primer

TATGGTAATT GT GTGYCAGCMGCCGCGGTAA

Read 2 sequencing primer:

Field descriptions (space-delimited):

1. Reverse primer pad
2. Reverse primer linker
3. Reverse primer

AGTCAGCCAG CC GGACTACNVGGGTCTAAT

Index sequencing primer:

Note: The 5' adapter sequence/index sequencing primer has an extra GCT at its 3' end compared to Illumina's usual index primer sequences. These bases were added to the 3' end of the Illumina 5' adapter sequence to increase the Tm for read 1 during sequencing.

AATGATAACGGCGACCACCGAGATCTACACGCT

References

- Apprill, A., McNally, S., Parsons, R. & Weber, L. Minor revision to V4 region SSU rRNA 806R gene primer greatly increases detection of SAR11 bacterioplankton. *Aquat Microb Ecol* 75, 129–137 (2015). [doi:10.3354/ame01753](https://doi.org/10.3354/ame01753)
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- Walters, W. et al. Improved Bacterial 16S rRNA Gene (V4 and V4-5) and Fungal Internal Transcribed Spacer Marker Gene Primers for Microbial Community Surveys. *mSystems* 1, 915 (2016). [doi:10.1128/mSystems.00009-15](https://doi.org/10.1128/mSystems.00009-15)

Materials

STEP MATERIALS

 Quant-iT PicoGreen dsDNA Assay Kit [Thermo Fisher Scientific Catalog #P11496](#)

 UltraClean PCR Clean-Up Kit (follow manufacturer's instructions) [Mobio Catalog #12500](#)

 Quant-iT PicoGreen dsDNA Assay Kit [Thermo Fisher Scientific Catalog #P11496](#)

 UltraClean PCR Clean-Up Kit (follow manufacturer's instructions) [Mobio Catalog #12500](#)

Protocol materials

 UltraClean PCR Clean-Up Kit (follow manufacturer's instructions) [Mobio Catalog #12500](#)

In Materials, Materials, Step 6

 Quant-iT PicoGreen dsDNA Assay Kit [Thermo Fisher Scientific Catalog #P11496](#)

In Materials, Materials, Step 4

Safety warnings

- ⚠ Please refer to the SDS (Safety Data Sheet) for hazard information.

- 1 Amplify samples in triplicate, meaning each sample will be amplified in 3 replicate 25- μ L PCR reactions.
- 2 Pool triplicate PCR reactions for each sample into a single volume (75 μ L).

Note

Do not combine amplicons from different samples at this point.

- 3 Run amplicons from each sample on an agarose gel.

Note

Expected band size for 515f-806r is ~300-350 bp. Low-biomass samples may yield faint or no visible bands; alternative methods such as a Bioanalyzer could be used to verify presence of PCR product.

- 4 Quantify amplicons with Quant-iT PicoGreen dsDNA Assay Kit (follow manufacturer's instructions).

 Quant-iT PicoGreen dsDNA Assay Kit **Thermo Fisher Scientific Catalog #P11496**

Protocol



NAME

Quanti-iT™ Pico Green dsDNA Assay (Invitrogen P7589)

CREATED BY

Bonnie Poulos

PREVIEW

- 4.1 Warm Quant-iT PicoGreen reagent to room temp in the dark.

Note

PicoGreen reagent is diluted in dimethylsulfoxide (DMSO) which solidifies at refrigerator temperatures. It must be completely liquified before use by allowing it to come to room temperature. Vortex solution briefly to mix well and centrifuge for 5 sec to bring liquid to bottom of tube; then dispense for use in the assay. PicoGreen reagent is also light-sensitive, so reagent should be protected from light.

 Quant-iT PicoGreen dsDNA kit **Thermo Scientific Catalog #P7589**

- 4.2 Prepare 1XTE buffer from 20X stock solution using nuclease-free water: will need 200 µl/well (for diluting standards, samples and PicoGreen).

Note

Prepare 1X TE by pipetting 2.5 mL of 20X stock TE into a sterile 50 mL centrifuge tube and filling to 50 mL mark with molecular biology grade water. Invert tube to mix.

- 4.3 Dilute DNA standard to either “High” 2 µg/mL (1:50 of λ DNA stock) or “Low” 50 ng/mL (1:1000 of λ DNA stock).

Note

It is best to run standards in duplicate, and if amount of DNA in samples is unknown or varies widely, it is also best to run both the high and low DNA standards.

- 4.4 Determine amount of sample to assay (eg, 2µl sample in total of 100µl TE buffer). Add correct amount of TE buffer to all wells. Add standards to wells. Then add samples to wells.

Note

See Guidelines for amount of DNA standards to add to standard wells.

- 4.5 Dilute PicoGreen 1:200 in TE buffer and protect from light until ready to add to plate.

Note

A 1:200 dilution of PicoGreen reagent is prepared by adding 10 µl of PicoGreen per 2 mL of 1X TE buffer. You will need 100 ul diluted PicoGreen per well containing 100 ul sample.

- 4.6 Add equivalent volume (100 µl) of diluted PicoGreen to every well (keeping plate in the dark as much as possible).

- 4.7 Tap plate to mix.

- 4.8 Incubate 5 minutes at room temperature keeping plate in the dark.

 00:05:00

- 4.9 Take fluorescent readings using 485nm excitation and 535nm emission filters.

- 4.10 Determine standard curve and calculate concentration of DNA in samples (see table in the guidelines).
- 5 Combine an equal amount of amplicon from each sample (240 ng) into a single, sterile tube. Higher amounts can be used if the final pool will be gel-isolated or when working with low-biomass samples.

Note

When working with multiple plates of samples, it is typical to produce a single tube of amplicons for each plate of samples.

- 6 Clean amplicon pool using MoBio UltraClean PCR Clean-Up Kit ([follow manufacturer's instructions](#)).

Note

If working with more than 96 samples, the pool may need to be split evenly for cleaning and then recombined. Optional: If spurious bands were present on gel (in step 3), one-half of the final pool can be run on a gel and then gel extracted to select only the target bands.



UltraClean PCR Clean-Up Kit (follow manufacturer's instructions) **Mobio Catalog #12500**

- 7 Measure concentration and A260/A280 ratio of final pool that has been cleaned.

Note

For best results the A260/A280 ratio should be between 1.8-2.0.

- 8 Send an aliquot for sequencing along with sequencing primers listed in Guidelines.