

Turn Spring into success with our flexible voucher option, our all-in-one package and flexible service. At StarSEQ we know that every project is unique.

1-Step 16S and ITS Microbiome analysis promotion and voucher option



1-Step Bacterial microbiome analysis or 1-Step Fungal microbiome analysis:

16S V4 region, Primer combination: 515F – 806R¹

16S V4-V5 region, Primer combination: 515F – 909R¹

16S V3-V4 region, Primer combination: 341F – 806R⁴

ITS 1 region, Primer combination: ITS1F – ITS2²





Price expamles, all other sample sizes are possible. We charge per sample. The price is a combination of a fixed price for the run and a variable price for the libraries depending on the number of samples.

No shared runs, customer samples are sequenced exclusively on one MiSeq run!

Eukaryotes (microbial) analysis:

V8-V9 region, Primer combination: 18S-1422f (5'-ataacaggtctgtgatgccct-3')

– 18S_1797R (5'-ccttcygcaggttcacctac-3')⁵

V9 region, Primer combination: Illumina_Euk_1391f (5'-GTACACACCGCCCGTC-3')

– Illumina_EukBr_1510r (5'-TGATCCTTCTGCAGGTTCACCTAC-3'), blocking primer optional³;

V9 region samples could not be combined with ITS, V8-V9 or 16S samples and need to be sequenced in a seperate MiSeq run because of smaller amplicon size!

Send us your gDNA samples and make use of our "All in One" service:



- Quality control
- Single step amplicon generation with reduced bias
- Double indexing, quality check, quantification, normalization and pooling of amplicons
- Illumina MiSeq / NextSeq 2000™ sequencing package A-D: 2 x 300 nt paired-end sequencing with V3 chemistry
- Output package A-D: 20-30 M reads (including 25 % PhiX to balance the composition of bases)
- De-multiplexing of reads
- Data delivery via FTP server

1,2,3,4,5 Primer sequence references: Please see page 4!







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2-Step 16S and ITS Microbiome analysis

2-Step Bacterial and Archaea Microbiome analysis:

16S, V1-V2 region, Primer combination 27F (5'-AGAGTTTGATCMTGGCTCAG-3') – 338R (5'-TGCTGCCTCCGTAGGAGT-3')

2-Step Fungal (18S) microbiome analysis:

ITS, ITS 1/2 region, Primer combination: ITS-u2 (5'-GAAYCATCGARTCTTTGAACGC-3') – ITS-p4 (5'-CCGCTTAKTGATATGCTTAAA-3')

ITS, ITS 2 region, Primer combination: ITS3F (5'-GCATCGATGAAGAACGCAGC-3') – ITS4R (5'-TCCTCCGCTTATTGATATGC-3')

2-Step Barcoding and Custom Amplicon analysis

2-Step Barcoding:

cytochrome b, cytb1 (5'-ccatccaacatctcagcatgatgaaa-3') – cytb2 (5'-gccctcagaatgatatttgtcctca-3')

cytochrome oxidase I, CO I E (5'-ccagagattagagggaatcagtg-3') – CO 1 F (5'-cctgcaggaggaggaggaycc-3')

or 2-Step Custom Amplicons:

design of custom primers with Illumina specific overhangs

Request for offer: microbiome@starseq.com

Optional: gold standard bioinformatic analysis of 16S/18S/ITS sequences (QIIME2) The bioinformatics analysis pipeline consists of quality control, pre-processing of reads, taxonomy identification and visualization, calculation of alpha and beta-diversity metrics.

Request for offer: microbiome@starseq.com









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16S and ITS Microbiome analysis

Sequencing of your ready to load 16S/ITS/18S libraries

• Illumina MiSeq / NextSeq 2000™ sequencing: 2 x 300 nt paired-end sequencing with V3 chemistry



- Output 20-30 M reads (including 25 % PhiX to balance the composition of bases)
- De-multiplexing of reads
- Free 16S metagenomics analysis (Illumina App)
- Data delivery via FTP server

Only 3300 € for MiSeq sequencing



Optimize your research with our 16S/ITS workflow solutions







From sample prep to library prep to sequencing and bioinformatics. StarSEQ offers a complete and comprehensive workflow for various 16S/ITS applications. Sequencing of the hypervariable regions of the 16S rRNA gene is the gold-standard for identification and classification of bacterial communities within a given sample.

StarSEQ is using established gold-standard primer sequences to target regions covering the hypervariable regions V3-V4, V4 and V4-V5. In contrast to common used methods, StarSEQ has designed its own single-step technique for simultaneous amplification, indexing and adaptoring of samples to reduce the bias caused during PCR.





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16S and ITS Microbiome analysis

Applications:

- Environmental Metagenomics (soil, water, air, biofilms, complex organic communities)
- Human or Animal Microbiome (skin, stool, gut, blood, swab)
- Monitoring of animal health (e.g. alternatively or supplementarily to FELASA-Test)
- Sterility Monitoring
- Detection of Contamination
- Monitoring of Biogas Plant
- Biosafety Monitoring
- Food Quality
- Clinical Samples . . .



As an additional service we offer also the purification of gDNA starting from 18 €.

Primer sequence references:

- ¹ Apprill A, McNally S, Parsons R, Weber L. 2015. Minor revision to V4 region SSU rRNA 806R gene primer greatly increases detection of SAR11 bacterioplankton. Aquat Microb Ecol 75:129–137. Parada AE, Needham DM, Fuhrman JA. Every base matters: assessing small subunit rRNA primers for marine microbiomes with mock communities, time series and global field samples. Environ Microbiol. 2016;18: 1403-1414.
- Parada, A. E., Needham, D. M., & Fuhrman, J. A. (2016). Every base matters: assessing small subunit rRNA primers for marine microbiomes with mock communities, time series and global field samples. Environmental Microbiology, 18(5), 1403–1414. https://doi.org/10.1111/1462-2920.13023
- ¹ Walters, W., Hyde, E. R., Berg-Lyons, D., Ackermann, G., Humphrey, G., Parada, A., ... Knight, R. (2016). Improved Bacterial 16S rRNA Gene (V4 and V4-5) and Fungal Internal Transcribed Spacer Marker Gene Primers for Microbial Community Surveys. mSystems, 1(1), e00009-15. https://doi.org/10.1128/mSystems.00009-15
- ² White, T. J., T. Bruns, S. Lee, and J. W. Taylor. 1990. Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. Pp. 315-322 In: PCR Protocols: A Guide to Methods and Applications. Academic Press, New York, NY. Gardes, M., and T. D. Bruns. 1993. ITS primers with enhanced specificity for basidiomycetes application to the identification of mycorrhizae and rusts. Mol. Ecol. 2: 113-118.
- ³ Amaral-Zettler, L. A., McCliment, E. A., Ducklow, H. W., & Huse, S. M. (2009). A Method for Studying Protistan Diversity Using Massively Parallel Sequencing of V9 Hypervariable Regions of Small-Subunit Ribosomal RNA Genes. *PLOS ONE, 4*(7), e6372. Retrieved from https://doi.org/10.1371/journal.pone.0006372
- ⁴ Klindworth A., Pruesse E., Schweer T., Peplies J., Quast C., Horn M., et al. . (2013). Evaluation of general 16S ribosomal RNA gene PCR primers for classical and next-generation sequencing-based diversity studies. Nucleic Acids Res. 41, 1–11. 10.1093/nar/gks808 see also https://support.illumina.com/downloads/16s_metagenomic_sequencing_library_preparation.html
- Enadley, I. M., Pinto, A. J., & Guest, J. S. (2016). Design and Evaluation of Illumina MiSeq-Compatible, 18S rRNA Gene-Specific Primers for Improved Characterization of Mixed Phototrophic Communities. Applied and Environmental Microbiology, 82(19), 5878 LP-5891. https://doi.org/10.1128/AEM.01630-16

 see also https://support.illumina.com/downloads/16s_metagenomic_sequencing_library_preparation.html

