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IMR Sequencing Services - Current Pricing (Summer 2015)

16S/18S Amplicon Pricing (per sample) – Academic

Library Preparation + Sequencing (~50,000 seq./sample) DNA/RNA Extraction² Standard Bioinformatics Analysis Pipeline³

<50 samples	50-200 samples	>200 samples ¹
\$20	\$15	\$12
\$13	\$13	\$13
\$500 / project	\$750 / project	\$1000 / project

16S/18S Amplicon Pricing (per sample) – Commercial

Library Preparation + Sequencing (\sim 50,000 seq./sample) DNA/RNA Extraction²

Standard Bioinformatics Analysis Pipeline³

<50 samples	50-200 samples	>200 samples ¹
\$40	\$30	\$25
\$15	\$15	\$15
\$750 / project	\$1000 / project	\$1500 / project

Metagenome Pricing – MiSeq

Extraction + Library Preparation + Sequencing (~3-4 M seq./sample)
Standard Bioinformatics Analysis Pipeline⁴

	Academic		Commercial	
]	Per run of 6	Per sample	Per run of 6	Per sample
	\$3600	\$600	\$4800	\$800
	\$750 / project		\$1500 / project	

Metagenome Pricing – NextSeq (coming Spring 2016)

Extraction + Library Preparation + Sequencing (~3-4 M seq./sample)
Standard Bioinformatics Analysis Pipeline⁴

Academic		Commercial	
Per run of 96	Per sample	Per run of 96	Per sample
\$13440	\$140	\$19200	\$200
\$750 / project		\$1500 / project	

Metatranscriptome Pricing – NextSeq (coming Spring 2016)

Contact us – pricing is currently being developed and is dependent upon multiple experimental choices.

Notes/Details:

- 1. One control well is required per 96-well PCR library plate, hence a maximum of $4 \times 95 = 380$ samples can be done together on one MiSeq run.
- 2. Extra costs may be incurred if samples, due to their nature, require more specialized (length, cost, etc.) extraction procedures.
- 3. Details of our 16S pipeline are available at https://github.com/mlangill/microbiome_helper, but the following is a summary of the major deliverables clients will receive (analyses will require a "mapping file" from the clients containing any relevant metadata for the study):
 - Combined FASTA file of the quality-controlled sequences (formatted for use in QIIME)
 - Final OTU tables in text, BIOM and STAMP formats (from open-ref. picking at 97%)
 - FASTA file of representative sequences (one per OTU)
 - Taxonomic assignment files at various levels (ex: phylum, genus, etc.)
 - Alpha-diversity rarefaction plots
 - Beta-diversity UniFrac plots
 - Functional prediction files generated from PICRUSt
- 4. Details of our metagenomics pipeline are available at https://github.com/mlangill/microbiome-helper, but the following is a summary of the major deliverables clients will receive (analyses will require a "mapping file" from the clients containing any relevant metadata for the study):
 - FASTA files of the final sequences screened to remove human (or other) contaminants
 - Taxonomic composition of the samples from MetaPhlAn 2.0 (text and STAMP files)
 - Functional prediction files generated from HUMAnN (text and STAMP files) for individual KO numbers, KEGG modules and KEGG pathways

Custom Bioinformatics: Additional bioinformatic analyses can be requested at an hourly rate or through research collaboration. Please contact us for more details.

Customs primers/amplicons: Beyond the 16/18S amplicons offered "in-stock", essentially any PCR amplicon (max. size ~ 550 bp) can be adapted to our MiSeq system. Please consults us for primer design/protocol details.

Processing time: Time to completion may be variable depending the amount of other partial projects in the queue – this applies primarily to amplicon runs when providing less than the full capacity of 380 samples, but may also occur when submitting small numbers of samples for metagenomics runs (<6 for MiSeq or <96 for NextSeq).