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Data Analysis Methodology – Microbial Diversity

Visual Overview of the Data Analysis Process
Overview of the Data Analysis Process

Once sequencing of your data has completed, the data analysis pipeline will begin processing the data for amplicon sequencing. Library samples are not denoised and will not have a standard data analysis. The data analysis pipeline consists of two major stages, the denoising and chimera detection stage and the microbial diversity analysis stage. During the denoising and chimera detection stage, denoising is performed using various techniques to remove short sequences, singleton sequences, and noisy reads. With the bad reads removed, chimera detection is performed to aid in the removal of chimeric sequences. Lastly, remaining sequences are then corrected base by base to help remove noise from within each sequence. During the diversity analysis stage, each sample is run through our analysis pipeline to determine the taxonomic information for each constituent read and then this information is collected for each sample. This stage is performed for all customers whose data is sequenced using primers targeting the 16S, 18S, 23S, ITS or SSU regions. Analysis can performed on other regions but may require additional charges.

The data analysis pipeline is broken down into the following steps, each of which is discussed more thoroughly in the sections below:

- Denoising and Chimera Checking
  1. Denoising
  2. Chimera Checking
  3. FASTQ File Generation

- Microbial Diversity Analysis
  1. Quality Checking and FASTA Formatted Sequence/Quality File Generation
  2. Sequence Clustering
  3. Taxonomic Identification
  4. Data Analysis

Denoising and Chimera Checking

Denoising

The process of denoising is used to correct errors in reads from next-generation sequencing technologies. According to the paper “Accuracy and quality of massively parallel DNA pyrosequencing” by Susan Huse, et al. and “Removing noise from pyrosequenced amplicons” by Christopher Quince, et al. the per-base error rates from 454 pyrosequencing attain an accuracy rate of 99.5% [3] [4]. The paper “A tale of three next generation sequencing platforms: comparison of Ion Torrent, Pacific Biosciences and Illumina MiSeq sequencers” by Michael Quail, et al. states that the observed error rates generated by the Illumina MiSeq is less than .4% while the Ion Torrent PGM has an error rate of 1.78% [5]. Due to the
large number of reads and even higher number of base calls per sequencing run, the total number of noisy reads can be quite substantial. In order to determine true diversity it becomes critical to determine which reads are good and which reads contain noise introduced by the experimental procedure. The RTLGenomics analysis pipeline attempts to correct this issue by denoising entire regions of data prior to performing any other steps of the pipeline.

Our analysis pipeline performs denoising by performing the following steps on each region:

1. The forward and reverse reads are taken in FASTQ format and are merged together using the PEAR Illumina paired-end read merger [6].
2. The FASTQ formatted files are converted into FASTA formatted sequence and quality files.
3. Reads are run through an internally developed quality trimming algorithm. During this stage each read has a running average taken across the sequence and is trimmed back at the last base where the total average is greater than 25.
4. Sequence reads are then sorted by length from longest to shortest.
5. Prefix dereplication is performed using the USEARCH [7] algorithm. Prefix dereplication groups reads into clusters such that each sequence of equal or shorter length to the centroid sequence must be a 100% match to the centroid sequence for the length of the sequence. Each cluster is marked with the total number of member sequences. Sequences < 100bp in length are not written to the output file, however no minimum cluster size restriction is applied which will allow singleton clusters to exist in the output.
6. Clustering at a 4% divergence is performed using the USEARCH [7] clustering algorithm. The result of this stage is the consensus sequence from each new cluster, with each tagged to show their total number of member sequences (dereplicated + clustered). Clusters that contain <2 members (singleton clusters) are not added to the output file, thus removing them from the data set.
7. OTU Selection is performed using the UPARSE OTU selection algorithm [8] to classify the large number of clusters into OTUs.
8. Chimera checking, which is explained in more detail below in the section entitled “Chimera Checking”, is performed on the selected OTUs using the UCHIME chimera detection software executed in de novo mode [9].
9. Each clustered centroid from step 6 listed above is then mapped to their corresponding OTUs and then marked as either Chimeric or Non-Chimeric. All Chimeric sequences are then removed.
10. Each read from step 3 is then mapped to their corresponding nonchimeric cluster using the USEARCH global alignment algorithm [7].
11. Using the consensus sequence for each centroid as a guide, each sequence in a cluster is then aligned to the consensus sequence and each base is then corrected using the following rules where C is the consensus sequence and S if the aligned sequence:
   a. If the current base pair in S is marked to be deleted, then the base is removed from the sequence if the quality score for that base is less than 30.
   b. If the current position in S is marked to have a base from C inserted, then the base is inserted into the sequence if the mean quality score from all sequences that mark the base as existing is greater than 30.
   c. If the current position in S is marked as a match to C but the bases are different, then the base in S is changed if the quality score for that base is less than 30.
   d. If a base was inserted or changed, the quality score for that position is updated. If the base was deleted the quality score for that position is removed.
   e. Otherwise, leave the base in S alone and move to the next position.
12. The corrected sequences are then written to the output file.

Chimera Checking

As discussed in the paper “Chimeric 16S rRNA sequence formation and detection in Sanger and 454-pyrosequenced PCR amplicons” by Brian Haas, et al. the formation of chimeric sequences occurs when an aborted sequence extension is misidentified as a primer and is extended upon incorrectly in subsequent PCR cycles [10]. This can be seen in Figure 1, shown below.
Because amplification produces chimeric sequences that stem from the combination of two or more original sequences [9], we will perform chimera detection using the de novo method built into UCHIME.

The RTLGenomics analysis pipeline performs chimera detection and removal by executing UCHIME [9] in de novo mode on the clustered data that was output by our denoising methods. By using this method, we can determine chimeras across entire region of data even after accounting for noise and removing low quality sequences.

**Microbial Diversity Analysis**

In order to determine the identity of each remaining sequence, the sequences must first be quality checked and demultiplexed using the denoised data generated previously. These sequences are then clustered into OTUs using the UPARSE [8] algorithm. The centroid sequence from each cluster is then run against either the USEARCH global alignment algorithm or the RDP Classifier against a database of high-quality sequences derived from the NCBI database. The output is then analyzed using an internally developed python program that assigns taxonomic information to each sequence and then computes and writes the final analysis files.

**Quality Checking and FASTA Formatted Sequence/Quality File Generation**

The denoised and chimera checked reads generated during sequencing are condensed into a single FASTA formatted file such that each read contains a one line descriptor and one to many lines of sequence/quality scores. The RTLGenomics analysis pipeline takes the FASTA formatted sequence and quality files and removes any sequence which fails to meet the following quality control requirements:

1. Sequences must be at least \( \frac{1}{2} \) the expected length given the primer sets used.
2. Sequences must be assigned to an expected sample on the run by barcode.

Sequences that pass the quality control screening are condensed into a single FASTA formatted sequence and quality file such that each read has a one line descriptor followed by a single line of sequence/quality data. The descriptor line in both files has been altered to contain the samples name followed by the original descriptor line, separated with a unique delimiter (::).

This stage of the pipeline creates the FASTA reads archive which contains the following files:

1. The sequence reads from all samples concatenated into a single sequence file. The original tags have been removed from each sequence and an “artificial tag” has been added in its place. The title of the file will be `<name>_<order ID>.fas`.
2. The quality scores from all samples concatenated into a single quality file. The scores are labeled with the corresponding sample name and will have a matching line in the .fas file.
the original tags were removed from the sequence and an “artificial tag” was put into its place, the quality scores have been similarly altered such that the original scores for the tag have been removed and an “artificial quality tag” has been added in its place. The artificial quality tag consists of Q30s for the length of the tag. This file will be labeled <name>_<order ID>.qual.

3. A mapping file consisting of samples included in the analysis. This file contains the information for each sample such that each line has the sample name, tag and primer used for the sample. This file will be labeled as: <name>_<order ID>.txt, and is formatted for use with QIIME/QIITA.

4. A second mapping file formatted for use with mothur, containing the primer information and barcodes used for each sample.

**OTU Selection**

OTU selection clusters sequences into clusters using an OTU selection program. By default, the OTU selection method is used to determine OTUs and uses the centroid sequence for each OTU to determine taxonomic information. OTU selection is performed using the guidelines discussed in the paper “UPARSE: Highly accurate OTU sequences from microbial amplicon reads” by Robert Edgar [8]. In that paper, the following methodology is laid out in order to select OTUs:

1. Perform dereplication on the sequences.
2. Remove all singleton clusters from the data set and sort the data by abundance.
3. Trim all sequences to the same length.
4. Perform OTU clustering using UPARSE.
5. Map original reads to the OTUs.

Dereplication of sequences is performed using the USEARCH prefix dereplication method [7]. Once complete we removed all singleton clusters and sorted the remaining sequences by cluster size from largest to smallest. The sequences are then run through a trimming algorithm that trims each sequence down to the same size. It should be noted that the sequences are only trimmed for UPARSE and the final taxonomic analysis is based upon the full length sequences. Next we use the UPARSE algorithm to select OTUs [8]. Using the USEARCH global alignment algorithm [7] we then assign each of the original reads back to their OTUs and write the mapping data to an OTU map and OTU table file.

**Tree Building**

Once OTU selection has been performed, a phylogenetic tree in Newick format is constructed. In order to construct the phylogenetic tree, multiple sequence alignment must be done on the OTU sequences in order to generate equal length aligned sequences. The multiple sequence aligner MUSCLE [11] [12], developed by Robert Edgar, is used with a maximum of 2 iterations in order to perform the alignment of
the OTU data. The finished multiple sequence alignment is then passed into FastTree [13] [14], developed by Morgan Price at the Lawrence Berkeley National Lab, a program used to infer approximately-maximum-likelihood phylogenetic trees from aligned sequence data. If you would like to learn more about how FastTree works, please visit the following link: http://www.microbesonline.org/fasttree/#How.

**Taxonomic Identification**

In order to determine the taxonomic information for each remaining sequence, the sequences must be run through either the USEARCH global alignment program or the RDP classifier. By default the USEARCH based method is employed however the RDP classifier can be substituted if a customer has requested that we use the RDP classifier instead. In either case the data is identified using a database of high quality sequences derived from NCBI that is maintained in house. If a customer would prefer we classify their data using a different database such as GreenGenes, then we can substitute that database in place of our own. If a non-standard database is requested that requires RTLGenomics to spend time converting or creating, then a small fee may be charged. Taxonomic identification may not be carried out for every assay depending on the availability of sequences for a gene. Accuracy may also be affected if the number of sequences available is small or not representative of the sampled group.

**USEARCH Global Search (Default)**

The global search method uses a mixture of the USEARCH global search algorithm along with a python program to then determine the actual taxonomic assignment that is assigned to each read. This method is described in the paper “An extensible framework for optimizing classification enhances short-amplicon taxonomic assignments” by Nicholas Bokulich, et al. [15]. The paper describes a methodology in which a high quality database is used in pair with USEARCH to rapidly find the top 6 matches in the database for a given sequence. From these 6 sequences you then assign a confidence value to each taxonomic level (kingdom, phylum, class, order, family, genus and species) by taking the number of taxonomic matches that agree with the top match and then divide by the number of total matches, e.g. If Bacteria is the top kingdom match with 5 matches showing Bacteria and 1 match showing Archaea, our algorithm would assign the kingdom Bacteria a confidence of 5/6 = .83.

**RDP Classifier**

The RDP Classifier is naïve Bayesian classifier than can rapidly determine taxonomic information for sequences while automatically determining the confidence it has at each taxonomic level [16]. The RDP classifier is run against an internally maintained database or a customer requested database along with a taxonomic file to help determine confidence values by giving the classifier a taxonomic tree.
Diversity Analysis

Regardless of the classifier that was used, the data next enters the diversity analysis program. This program takes the OTU/Derep table output from sequence clustering along with the output generated during taxonomic identification and begins the process of generating a new OTU table with the taxonomic information tied to each cluster. This updated OTU table is then written to the output analysis folder with both the trimmed and full taxonomic information for each cluster. For each taxonomic level (kingdom, phylum, class, order, family, genus and species) four files are generated which contain the number of sequences per full taxonomic match per sample, the percentage per full taxonomic match per sample, the number of sequences per trimmed taxonomic match per sample and the percentage per trimmed taxonomic match per sample. These files are all described in more detail below.

FAQ (Frequently Asked Questions)

What does R1 and R2 mean on my raw FASTQ files?

Most sequencing performed on our Illumina sequencers was run using paired-end sequencing. This generates two FASTQ files where the first (R1) file contains reads that start at the forward primer and go towards the reverse and the second (R2) file contains the corresponding reverse reads that are oriented as starting from the reverse primer and going towards the forward. For more information please see the section titled “Paired End FASTQ Files” found below on page 19.

Can I use QIIME or MOTHUR with RTLGenomics Data?

QIIME [1] and MOTHUR [2] are open-source bioinformatics pipelines for performing microbiome analysis from raw DNA sequencing data. RTLGenomics provides clients that receive our analysis a copy of their data already quality checked and ready for use in both QIIME and MOTHUR. You can find the sequence data for QIIME and MOTHUR in the FASTA zip archive discussed in the section “FASTA Archive File Descriptions” found on page 21.

You can find documentation for or download QIIME at http://qiime.org/.

You can find documentation for or download MOTHUR at http://www.mothur.org.

How do I use QIIME with RTLGenomics Data?

QIIME’s split_libraries.py workflow is used to demultiplex sequence data and format it to work with the downstream analysis tools. This particular workflow requires three files to work, a FASTA formatted sequence file, a FASTA formatted quality file and a mapping file that tells QIIME what the sample’s
names, barcodes and primers are. We strongly recommend you use the following options when running split_libraries.py using our FASTA data:

split_libraries.py –m <Mapping TXT file> -f <Fasta FNA File> -q <Qual File> -o <Output Folder> -H 1000 –p

The –m, -f, -q and –o options are all required and tell the workflow where the mapping, FASTA sequence, FASTA quality and output files are respectively. However, we also recommend you add the following options:

- –p
  o The –p options instructs QIIME to ignore the primer because otherwise it will require the primer be present in order for the sequence to be valid.
  o The forward primer was present on most platforms, but the reverse primer was often not sequenced on the 454 and Ion Torrent.
  o Our pipeline removes the primer from each sequence during our QC steps, so even if the primer exists on the raw data it will not exist in the data that has been prepared for QIIME.

- –H 1000
  o The –H option sets the maximum number of homopolymers the sequence can contain and still be considered valid.
  o We recommend using this option with Illumina data because the Illumina platform does not suffer from issues regarding homopolymers (at least not the point that the other platforms do). As such you can set the –H option to a very high number to keep QIIME from throwing away valid sequences. This is a bigger issue in fungal samples than bacterial.

How do I use MOTHUR with RTLGenomics Data?

MOTHUR’s Trim.seqs command is used to demultiplex sequence data and format it to work with the downstream analysis commands. This command requires two files to work, a FASTA formatted sequence file and an oligos mapping file. If you provide this command with the FASTA sequence file and oligos file provided by RTLGenomics, then you should be able to use this data in all downstream analysis.

What exactly does No Hit mean in my analysis?

The RTLGenomics analysis pipeline generates a number of files as discussed in the section “Analysis Archive File Descriptions” on page 22. The taxonomic information given in the “FullTaxa” and “TrimmedTaxa” are often the assigned taxonomic information, but in some cases the taxonomic information may contain the “No Hit” keyword for each taxonomic level. The “No Hit” keyword simply means that there exists no matches in our database that meet the minimum criteria to be considered
likely. As such we cannot make any confident calls regarding the taxonomic classification of the OTU – even at the Kingdom/Domain level. There exists a number of reasons why this occur which we will describe below:

1. The organism’s sequence is missing from our database
   a. New sequences and organisms are added to NCBI/EMBL/DDBJ on a daily basis and, while RTLGenomics makes every effort to keep our databases current, it may take a few months before new sequences are in our database.
   b. Many sequences in NCBI/EMBL/DDBJ are too short or contain no taxonomic information and are excluded from being added to our database.

2. The organism’s sequence data is not yet in NCBI/EMBL/DDBJ.
   a. NCBI/EMBL/DDBJ contain a vast amount of data, however they require researchers to have already sequenced an organism before they have sequence data to provide. If no one has sequenced the organism and submitted it to those repositories, then we will not yet have the sequence.
   b. While this is not often the case, there exists the possibility that your sequence data contains an organism not yet known to science, i.e. a novel species

3. Low quality sequence
   a. While RTLGenomics does perform quality and chimera checking on your data, these algorithms are not fool-proof and low quality or chimeric sequences may have managed to make it to the taxonomic analysis stage. These sequences will often fail to identify as any organism due to their low quality or chimeric nature, causing them to be marked as “No Hit”.

What is the difference between Unknown and Unclassified in my analysis files?

The RTLGenomics analysis pipeline generates a number of files as discussed in the section “Analysis Archive File Descriptions” on page 22. The taxonomic information given in the “FullTaxa” and “TrimmedTaxa” are often the assigned taxonomic information, but in some cases the taxonomic information may contain the “Unknown” and “Unclassified” keywords which can cause some confusion. These keywords are described as follows:

- **Unknown**
  - Our algorithm was unable to make a confident determination regarding the taxonomic classification at a certain level.
  - See section “USEARCH Global Search (Default)” on page 10 for more information for how we determine confidence.

- **Unclassified**
The taxonomic information retrieved from NCBI contains missing information at this level.

For instance, if the best match in our database is classified in NCBI down to the Family level then our database will mark the Genus and Species as “Unclassified”.

How are confidence values determined?

The RTLGenomics analysis pipeline generates a number of files as discussed in the section “Analysis Archive File Descriptions” on page 22. Once each OTU has been aligned to our database, our algorithm will select the top/best six matches for the OTU and attempt to assign a confidence value to each taxonomic level. The top match is then compared against the other five to determine the number of matches at agree with the base match at each taxonomic level. The number of agreements is then converted into a confidence value using the following equation:

\[
\text{confidence} = \frac{\text{Number of Matching Taxa}}{\text{Total Number of Taxa}}
\]

For instance, if OTU ID #13 has 6 top matches with the following taxa:

1) Bacteria; Proteobacteria; Alphaproteobacteria; Rhizobiales; Bradyrhizobiaceae; Bradyrhizobium; Bradyrhizobium sp
2) Bacteria; Proteobacteria; Alphaproteobacteria; Rhizobiales; Bradyrhizobiaceae; Bradyrhizobium; Bradyrhizobium sp
3) Bacteria; Proteobacteria; Alphaproteobacteria; Rhizobiales; Bradyrhizobiaceae; Bradyrhizobium; Bradyrhizobium sp
4) Bacteria; Proteobacteria; Alphaproteobacteria; Rhizobiales; Bradyrhizobiaceae; Bradyrhizobium; Bradyrhizobium japonicum
5) Bacteria; Proteobacteria; Alphaproteobacteria; Rhizobiales; Bradyrhizobiaceae; Bradyrhizobium; Bradyrhizobium japonicum
6) Bacteria; Proteobacteria; Alphaproteobacteria; Rhizobiales; Bradyrhizobiaceae; Rhodopseudomonas; Rhodopseudomonas palustris

Then each taxonomic level would receive the following confidence:

<table>
<thead>
<tr>
<th>Kingdom</th>
<th>Phylum</th>
<th>Class</th>
<th>Order</th>
<th>Family</th>
<th>Genus</th>
<th>Species</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacteria</td>
<td>Proteobacteria</td>
<td>Alphaproteobacteria</td>
<td>Rhizobiales</td>
<td>Bradyrhizobiaceae</td>
<td>Bradyrhizobium</td>
<td>Bradyrhizobium sp</td>
</tr>
</tbody>
</table>
Using this example, we would assign a confidence of 1 (100%) to the kingdom, phylum, class, order and family taxa. We would then assign a confidence of .83 (83%) to the genus taxon and a confidence of .5 (50%) to the species. These confidence values are then used when the “TrimmedTaxa” files are generated.

**What is the difference between FullTaxa and TrimmedTaxa Files?**

The RTLGenomics analysis pipeline generates a number of files as discussed in the section “Analysis Archive File Descriptions” on page 22. In our analysis data we provide most of our analysis in duplicate files, one containing the analysis using the “FullTaxa” and the other containing the analysis using the “TrimmedTaxa”. Please see “How are confidence values determined?” found on page 14 as this discussion will assume you have an idea of what confidence values are and how we assign them.

The FullTaxa files are generated under the assumption that the best match is the correct one. As such each taxonomic level is assigned using that match without taking the confidence values into account. We provide this file for two reasons: 1) this method for assigning taxa is similar to the method we used many years ago and we continue to provide these files for legacy purposes and 2) this helps you see what the original best match was before confidence trimming occurred. We believe these files are a powerful tool in allowing you to better get an idea of what our algorithm originally thought the data contained before confidence values were taken into account, which can help you get a better feel for the data. However, we do advise that you do not use only the FullTaxa data to perform your analysis as the lack of confidence makes the data considerably less accurate.

The TrimmedTaxa files are generated using the FullTaxa data after the confidence values have been taken into account. As such each taxonomic level is assigned only if the confidence value is greater than or equal to .51 (51%). If a taxon falls below .51, it is replaced with the “Unknown” keyword. Using the example data provided in the section “How are confidence values determined?” on page 14, the FullTaxa would read *Bacteria ; Proteobacteria ; Alphaproteobacteria ; Rhizobiales ; Bradyrhizobiaceae ; Bradyrhizobium ; Bradyrhizobium sp* and the TrimmedTaxa would read *Bacteria ; Proteobacteria ; Alphaproteobacteria ; Rhizobiales ; Bradyrhizobiaceae ; Bradyrhizobium ; Unknown*, where the species is now assigned “Unknown” due to the low confidence value.

**Brief Synopsis**

- FullTaxa
o Generated under the assumption that the top/best match is completely accurate.
 o Does not take confidence values into account.
 o Provided primarily for legacy purposes and not recommended to be used for detailed analysis.

- TrimmedTaxa
  o Takes the confidence value into account at each taxonomic level.
  o Replaces low confidence taxa with the “Unknown” keyword.

Do my sequences contain the primer, barcodes or adapters?

Upon the completion of an order at RTLGenomics, clients will receive two zip archives containing their sequence data, the files are described in the section “

File Descriptions and Formatting” starting on page 17. The raw data archive contains your sequence data directly from the sequencer with no post-processing done on our end. This data is packaged as one SFF or one pair of FASTQ files per sample. The FASTA data archive contains your sequence data after we have performed denoising and some basic quality checking on the data. This data is provided as a single FASTA formatted sequence and quality file that contains all of your sequences multiplexed together.

Please refer to the following graph in order to determine whether the primer, barcodes or adapters are on your sequences:

<table>
<thead>
<tr>
<th>Data Type</th>
<th>Primer/Barcode</th>
<th>illumina(^1) Single PCR</th>
<th>illumina(^1) Double PCR</th>
<th>Illumina Library</th>
</tr>
</thead>
<tbody>
<tr>
<td>Raw</td>
<td>Forward Primer</td>
<td>Not Present</td>
<td>Present(^2)</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>Reverse Primer</td>
<td>Not Present</td>
<td>Present(^2)</td>
<td>N/A</td>
</tr>
<tr>
<td>Raw</td>
<td>Barcodes</td>
<td>Not Present</td>
<td>Not Present</td>
<td>Not Present</td>
</tr>
<tr>
<td>FASTA</td>
<td>Forward Primer</td>
<td>Not Present</td>
<td>Not Present</td>
<td>N/A</td>
</tr>
<tr>
<td>FASTA</td>
<td>Reverse Primer</td>
<td>Not Present</td>
<td>Not Present</td>
<td>N/A</td>
</tr>
<tr>
<td>FASTA</td>
<td>Barcodes</td>
<td>Present(^3)</td>
<td>Present(^3)</td>
<td>Present(^3)</td>
</tr>
</tbody>
</table>

\(^1\) Most samples are run using the MiSeq Double PCR method, if you need to confirm which method your samples were run on then please contact us at lab@rtlgenomics.com.

\(^2\) The forward primer will be found in the R1 files, the reverse primer will be found in the R2 files.

\(^3\) The barcode is present however it is not the barcode used during sequencing.
Term Definitions

Terms used within this guide are defined as follows:

- **Tag**
  - The term tag refers to the 8-10 bp sequence at the 5’ end of the sequence read.
  - The tag is also known as the barcode in some programs.

- **ASCII value**
  - ASCII (American Standard Code for Information Interchange) is a character encoding scheme based on the English alphabet to encode the following: the numbers 0-9, the letters a-z, the letters A-Z, basic punctuation, control codes (such as new line), and the blank space.
  - Each letter, number and punctuation mark on a keyboard is assigned a numeric value (mostly between 0 and 127) using the ASCII table in order to create a way of encoding/decoding character symbols into computer readable digital bit patterns.

File Descriptions and Formatting

**Zip Archives**

The following archives will be passed along to you upon completion of your order:

- `<Name>_<OrderNumber>Raw<SequencingDate>.zip`
  - This archive contains the raw FASTQ and/or raw SFF as described in the “Raw Sequence Data File Formats” section found on page 18.

- `<Name>_<OrderNumber>Fasta<SequencingDate>.zip`
  - This archive contains the denoised sequence data for your entire order in FASTA/Qual format as described in the “FASTA Archive File Descriptions” section found on page 21.

- `<Name>_<OrderNumber>Analysis<SequencingDate>.zip`
  - This archive contains the analysis data described in the “Analysis Archive File Descriptions” section found on page 22.
  - This archive will only be sent if you used a standard primer set that we have a working database for. Custom or functional assays will likely not be analyzed.
Split Zip Archives

If any zip archive is larger than 10GB in size, we will be unable to upload the file to our file server without breaking the file into smaller chunks. In order for you to open these files you will need to download each file in the archives set (denoted with <ArchiveName>.zip.XXX where XXX is a number starting at 001 and counting upwards) and then stitch them back together prior to unzipping the archive. The following commands can be used to rebuild the zip file prior to unzipping.

**Windows**

Stitching the files together in Windows requires you to do the following:

1. Open a command/DOS prompt
   - In most versions of windows go to “Start menu” then type in cmd and run cmd.exe.
2. Navigate to the folder you downloaded the files into.
3. Type in the following: `copy /B ArchiveName.zip.* ArchiveName.zip`
4. Unzip the ArchiveName.zip file as you normally would.

**Linux / Mac**

Stitching the files together in Linux or Mac requires you to do the following:

1. Open a command terminal.
2. Navigate to the folder you downloaded the files into.
3. Type in the following: `cat ArchiveName.zip.* > ArchiveName.zip`
4. Unzip the ArchiveName.zip file as you normally would.

Raw Sequence Data File Formats

**FASTQ File**

FASTQ files are text based formatted data files that store the nucleotide sequences generated by the sequencer and their corresponding quality scores encoded as ASCII characters. A FASTQ file contains 4 lines per read that contain the following information:

- Line 1 contains the sequence ID (read definition) and is prepended with an “at” symbol, ‘@’.
- Line 2 contains the sequence data.
- Line 3 acts as a separator line between the sequence data and the quality score, it contains a single plus sign, ‘+’.
• Line 4 encodes the quality values for the sequence in line 2 with each quality score being represented by a single character. As such Line 2 and Line 4 must be the same length.

Decoding of the quality scores requires you to know the phred score offset that was used when the file was generated. Once you know the offset, you can take the ASCII value for the given character and subtract the offset value to obtain the quality score. For example, if the phred offset is +33 and the character ‘B’ is encountered, then the quality score for that position would be 33 as ‘B’ is represented by the ASCII value 66 and the offset is 33 (66 – 33 = 33). Using the same logic, ‘A’ (represented by the value 65) would be 65-33=32 meaning the ‘A’ character represents a quality score of 32. A free to view ASCII table can be found here: http://www.ascii-code.com/.

Paired End FASTQ Files

Paired end reads are stored in two FASTQ files with the first file storing the forward “half” of the read and the second file which stores the reverse “half” of the read. It should be noted that both reads are provided in forward order, meaning if you wish to link the two reads together you will first need to take the reverse complement of the reads in the second file. Depending on the insertion size and sequencing read length, the forward and reverse reads may or may not overlap at some point. Unlike other FASTQ files, the order of reads within these files must be kept in a specific order to avoid issues with most post processing programs. The reads within these two files must stay the same between the two files, meaning if you remove or move a sequence in one file, you must remove it or move it to same place in the other file to preserve the same order in both files.

Because the insert size for a paired end sequence matters, we provide two examples of how your sequences may, or may not, line up. In both examples it is assumed that you have already taken the reverse complement of the reverse reads.

**Example 1** – Insert Size approx. 500bp using a 2x300 kit.

Forward----------------------------------------------------------

~250 BP Forward Only | ~50 BP Alignment | ~250 BP Reverse Only

-------------------------------------------------------------------Reverse

**Example 2** – Insert Size approx. 800bp using a 2x300 kit.

Forward-----------------------------------------------


**BAM/SAM files**

BAM files are binary files used to store nucleotide reads in SAM format. The SAM format was developed to store alignment information for each sequence, but it is possible to have unaligned SAM files as well. Each read in the file has a header section with a header (@HD), a sequence dictionary (@SQ), read group (@RG), program (@PG) and may have comments (@CO), and an alignment section describing the read’s alignment and quality. The full file specification can be found at [http://samtools.github.io/hts-specs/SAMv1.pdf](http://samtools.github.io/hts-specs/SAMv1.pdf). BAM files cannot be read with any text editor and have to first be converted to SAM format before they are human readable. Programs like samtools can convert BAM to SAM or other sequence formats, losing some of the metadata associated if the target format does not store that information.

**Raw Sequencer Output**

**Illumina MiSeq, HiSeq, NextSeq**

The Illumina MiSeq produces FASTQ files with a phred offset of +33. While the FASTQ file(s) generated do contain all of the raw sequence data generated by the sequencer, they may contain information regarding the primer (forward or reverse) for amplicon sequencing. If primer-specific adapters are used for sequencing, the generated files will not contain the forward or reverse primer. To allow more flexibility of runs, RTL Genomics has used a two-step PCR process for Illumina sequencing that uses universal adapters and does sequence the forward and reverse primers. This allows additional QC to be done to remove reads with incorrect primers from tag hopping.

FASTQ files generated by the Illumina sequencers come in two forms depending on the sequencing method used – either paired end or single end. Single end reads are stored in a single FASTQ file with each read in the file representing a read from the sequencer. Paired end reads, however, are slightly more complex and are covered in paired end FASTQ section. Reads from the Illumina sequencer are stored per sequence and are demultiplexed by the Illumina Software, thus your raw data will be missing all barcode information.

**PacBio Sequel**

The Sequel generates its primary output in unaligned BAM format. PacBio specific information regarding the BAM header values may be found at [https://pacbiofileformats.readthedocs.io/en/5.1/BAM.html](https://pacbiofileformats.readthedocs.io/en/5.1/BAM.html). This document is periodically updated by PacBio as new versions of SMRT Link are released. Each SMRT cell has a folder containing all the related data for the cell. Many of the files are metadata and are in xml
These contain information about the run, and many run metrics for use by PacBio support. The primary output file is the `<movie_name>.subreads.bam` file, which contains all the high quality subreads from the cell. The subreads are grouped by ZMW in the order that they were sequenced. The `<movie_name>.scraps.bam` file contains the low-quality regions, hairpin adapters, and barcode regions if applicable. This file is only used for barcoding and can be ignored otherwise. It is typically the largest file generated for a cell, so it can be excluded from unmultiplexed cells to reduce raw data size without losing relevant information. The `.pbi` files are bam index files, used by Pacbio tools in later data analysis, and can be ignored if using other tools to analyze your data.

**FASTA Archive File Descriptions**

The FASTA archive you receive with your data will contain the following files:

- **<Name>_<OrderNumber>.fna**
  - This file contains your sequence data after it has undergone quality trimming, denoising and quality checking. Each sequence in this file has had an 8 nucleotide barcode prepended to the front of it. This barcode is not the barcode that was originally used during sequencing, instead we use a faux barcode for each sample to ensure that they each have a unique barcode.
  - **File Type:** FASTA formatted sequence data. Sequence definition lines begin with a ‘>' (greater than) character followed by 1 or more lines of sequence data.
  - **Recommended Program:** Any text editor.

- **<Name>_<OrderNumber>.qual**
  - This file contains the quality scores for your sequence data after it has undergone quality trimming, denoising and quality checking. Each quality score set in this file has had 8 fake 40 scores prepended to the front of it to account for the faux barcode added to the sequence file.
  - **File Type:** FASTA formatted sequence data. Sequence definition lines begin with a ‘>' (greater than) character followed by 1 or more lines of sequence quality data.
  - **Recommended Program:** Any text editor.

- **<Name>_<OrderNumber>.txt**
  - This file stores a table that maps each faux barcode generated in the sequence file to the sample it corresponds to. The table contains four columns and has been formatted to be compatible with QIIME, however some changes may be required if your sample names contain characters or symbols that are disallowed in QIIME.
    - The table contains the following four columns:
      - Column 1 contains the sample name (SampleID) for your sample.
      - Column 2 contains the faux barcode sequence added to you sequence file.


- Column 3 contains the forward primer sequence that was used for that sample.
- Column 4 contains the reverse primer sequence that was used for that sample.
- Column 5 contains the description of the sample which is required for QIIME. Because we do not know exactly what your samples are, we instead just fill this column with the assay name.

  o **File Type:** Table written in tab-separated value (TSV) format.
  o **Recommended Program:** Any spreadsheet program or text editor.

- **<Name>_<OrderNumber>.<AssayName>.oligos**
  o This file stores mapping information that maps each faux barcode generated in the sequence file to the sample it corresponds to. The file contains the forward and reverse primers followed by the barcode and samples name for each sample that used that forward and reverse primer combination. This file has been formatted to be compatible with MOTHUR, however some changes may be required if your sample names contain characters or symbols that are disallowed in MOTHUR.

  o **File Type:** A standard text file.
  o **Recommended Program:** Any text editor.

### Analysis Archive File Descriptions

The analysis archive you receive with your data will contain the following files:

- For each taxonomic level (<level>) where level is Kingdom, Phylum, Class, Order, Genus or Species.
  - FullTaxa.<level>.counts.txt
    - This file contains a table with the columns representing each sample in your order and the rows representing each unique taxonomic information for the top hit listed down to <level>, e.g. if <level> is Phylum then it will give each unique Kingdom/Phylum combination.
    - Each row/column intersection defines the number of sequences in the sample that matched that particular unique taxonomic information.
    - Keep in mind that the Full Taxa data shows only the taxonomic information for the top hit, regardless of what the confidence values were.
    - **File Type:** Table written in tab-separated value (TSV) format.
    - **Recommended Program:** Any spreadsheet program or text editor.
  - FullTaxa.<level>.percent.txt
    - This file contains a table with the columns representing each sample in your order and the rows representing each unique taxonomic information for the top hit listed down to <level>, e.g. if <level> is Phylum then it will give each unique Kingdom/Phylum combination.
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- Each row/column intersection defines the percent of sequences in the sample that matched that particular unique taxonomic information.
- Keep in mind that the Full Taxa data shows only the taxonomic information for the top hit, regardless of what the confidence values were.
- **File Type:** Table written in tab-separated value (TSV) format.
- **Recommended Program:** Any spreadsheet program or text editor.
  - TrimmingTaxa.<level>.counts.txt
    - This file contains a table with the columns representing each sample in your order and the rows representing each unique taxonomic information for the top hit listed down to <level>, e.g. if <level> is Phylum then it will give each unique Kingdom/Phylum combination.
    - Each row/column intersection defines the number of sequences in the sample that matched that particular unique taxonomic information.
    - Keep in mind that the Trimmed Taxa data shows the taxonomic information after the confidence values are taken into account. The USEARCH method rejects the taxonomic information at a level if the confidence is below 51% while the RDPClassifier uses a minimum confidence of 80%.
  - **File Type:** Table written in tab-separated value (TSV) format.
  - **Recommended Program:** Any spreadsheet program or text editor.
  - TrimmingTaxa.<level>.percent.txt
    - This file contains a table with the columns representing each sample in your order and the rows representing each unique taxonomic information for the top hit listed down to <level>, e.g. if <level> is Phylum then it will give each unique Kingdom/Phylum combination.
    - Each row/column intersection defines the percent of sequences in the sample that matched that particular unique taxonomic information.
    - Keep in mind that the Trimmed Taxa data shows the taxonomic information after the confidence values are taken into account. The USEARCH method rejects the taxonomic information at a level if the confidence is below 51% while the RDPClassifier uses a minimum confidence of 80%.
  - **File Type:** Table written in tab-separated value (TSV) format.
  - **Recommended Program:** Any spreadsheet program or text editor.
- **OTU/Derep tables**
  - FullTaxa.otu_table.txt
    - This file contains a table with the columns representing each sample in your order and the rows representing each unique OTU or Dereplication Cluster. The final column contains the taxonomic information for that particular OTU/Cluster listed down to the Species level.
Keep in mind that the Full Taxa data shows only the taxonomic information for the top hit, regardless of what the confidence values were.

- **File Type:** Table written in tab-separated value (TSV) format.
- **Recommended Program:** Any spreadsheet program or text editor.
  - FullTaxa.otu_table.biom
    - This file contains a BIOM formatted copy of the OTU Table stored in FullTaxa.otu_table.txt.
    - Keep in mind that the Full Taxa data shows only the taxonomic information for the top hit, regardless of what the confidence values were.
    - **File Type:** OTU Table in BIOM Format.
    - **Recommended Program:** Any text editor or program that accepts BIOM files (e.g. QIIME).
  - TrimmedTaxa.otu_table.txt
    - This file contains a table with the columns representing each sample in your order and the rows representing each unique OTU or Dereplication Cluster. The final column contains the taxonomic information for that particular OTU/Cluster listed down to the Species level.
    - Keep in mind that the Trimmed Taxa data shows the taxonomic information after the confidence values are taken into account. The USEARCH method rejects the taxonomic information at a level if the confidence is below 51% while the RDPClassifier uses a minimum confidence of 80%.
    - **File Type:** Tables written in tab-separated value (TSV) format.
    - **Recommended Program:** Any spreadsheet program or text editor.
  - TrimmedTaxa.otu_table.biom
    - This file contains a BIOM formatted copy of the OTU Table stored in TrimmedTaxa.otu_table.txt.
    - Keep in mind that the Trimmed Taxa data shows the taxonomic information after the confidence values are taken into account. The USEARCH method rejects the taxonomic information at a level if the confidence is below 51% while the RDPClassifier uses a minimum confidence of 80%.
    - **File Type:** OTU table in BIOM format.
    - **Recommended Program:** Any Text Editor or program that accepts BIOM files (e.g. QIIME).

- **Krona Folder**
  - **Raw Data Folder**
    - This folder contains the raw data files that were passed to Krona in order to generate the FullTaxa and TrimmedTaxa Krona HTML files. These files were derived directly from the FullTaxa.species.counts.txt and
TrimmedTaxa.species.counts.txt files. These files are provided for transparency purposes regarding how your visualization data was created.

- **FullTaxa.krona.html**
  - This file contains the Krona visualization of the FullTaxa.species.count.txt file. The visualization file is a standard HTML file and should be accessible using any web browser. This visualization was generated using the FullTaxa.species.counts.txt file described before and contains data on all samples found in that file. You are able to switch between samples using the menu on the left hand side of the screen. Keep in mind that the “Collapse” checkbox is checked by default which can cause your taxonomic levels to look incorrect.
  - **This file requires an internet connection in order to work.**
  - **File Type:** HTML file with embedded code from Krona.
  - **Recommended Program:** Any Web Browser, however the developers suggest using FireFox.

- **TrimmedTaxa.krona.html**
  - This file contains the Krona visualization of the FullTaxa.species.count.txt. The visualization file is a standard HTML file and should be accessible using any web browser. This visualization was generated using the TrimmedTaxa.species.counts.txt file described before and contains data on all samples found in that file. You are able to switch between samples using the menu on the left hand side of the screen. Keep in mind that the “Collapse” checkbox is checked by default which can cause your taxonomic levels to look incorrect.
  - **This file requires an internet connection in order to work.**
  - **File Type:** HTML file with embedded code from Krona.
  - **Recommended Program:** Any Web Browser, however the developers suggest using FireFox.

- **OTUs Folder**
  - **OtuMap.txt**
    - This file contains the mapping of each OTU identification number used in the OTU Table files. Each line contains the following information separated by tabs: the OTU identification number, the number of sequences that make up the OTU, the sequence definition for the seed sequence and then the sequence definition for each member sequence.
    - **File Type:** Table written in tab-separated value (TSV) format. The first column is the OTU #, the 2nd can be ignored, the 3rd is the OTU centroid sequence definition and columns 4+ contain each member sequence definition.
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- **Recommended Program:** Any spreadsheet program or text editor.
  - **OTUs.fas**
    - **This file contains the OTU sequences selected during sequence clustering in fasta format.** For information regarding how this file was generated please see Error! Reference source not found. on page Error! Bookmark not defined.
    - **File Type:** FASTA formatted sequence data. Sequence definition lines begin with a ‘>’ (greater than) character followed by 1 or more lines of sequence data.
  - **Recommended Program:** Any text editor.

- **TreeData Folder**
  - **otu_map.condensed.txt**
    - **This file contains a condensed version of the OtuMap.txt file.** Each line contains two columns separated by tabs. The first column gives the OTU identification number and the second column contains the sequence definition for the seed sequence. Thus this file contains the equivalent of columns #1 and #3 from the OTU Map.
    - **File Type:** Table written in tab-separated value (TSV) format
    - **Recommended Program:** Any Spreadsheet program or Text Editor
  - **otus.msa**
    - **This file contains the multiple sequence alignment for each OTU described in the OTU table and OTU map.** This file was generated using MUSCLE as described above in the section titled Tree Building.
    - **File Type:** FASTA formatted sequence data. Sequence definition lines begin with a ‘>’ (greater than) character followed 1 or more lines of sequence data.
    - **Recommended Program:** This file is best viewed using a MSA viewer but can also be viewed using any text editor.
  - **otus.tre**
    - **This file contains the phylogenetic tree in Newick tree format created using the otus.msa file.** This file was generated using FastTree as described in the section titled Tree Building on page 9.
    - **File Type:** Newick tree formatted phylogenetic tree.
    - **Recommended Program:** This file is best viewed using a phylogenetic tree viewer but can also be viewed using any text editor.
    - **Disclaimer:** Numerous tree viewers apply restrictions on the Newick format that are not standard. As such this file may not be readable by all tree viewers. We do try to constrain the data to work with as many viewers as possible, but we can in no way make it work for all.
Database Maintenance Policy

RTLGenomics makes every effort to keep our databases as current as possible. This requires periodic updates to include new taxonomies and sequences as they become available. Database updates occur on an irregular schedule, so we cannot guarantee orders submitted at different times will be run against the same database version. If you have concerns about which version your samples were run against, or if you need multiple orders run against the most recent version, please contact us. Orders may be combined or rerun against a specified database for a nominal fee, however this fee may be waived if the need to merge orders is discussed at the beginning of a project.

Client Data Retention Policy

Data will made available for download (typically via a 90 day temporary link) upon completion of your project. RTL will make every reasonable effort to store all electronic data for your project for a period of 6 months from the date of notification that the project has been completed. If you have any questions regarding your data or if you need to discuss longer term storage, please contact us.

Recommended Programs

This section will give a brief description of some programs that we suggest you use in order to view or edit the data we have passed along.

- Text Editors
  - Let us first note that a text editor and a word processor are vastly different programs. When we say text editor we are discussing programs that often do not allow for changes in font, size or stylization. As such we strongly advise you to avoid using Microsoft Word, OpenOffice Writer, Wordpad or Google Docs to view or edit any file we provide.
  - Recommended Text Editors (Paid Usage)
    - UltraEdit: http://www.ultraedit.com/
  - Recommended Text Editors (Free)
    - Notepad++: http://notepad-plus-plus.org/
    - Unix & Linux Text Editors: GEdit / vi / emacs

- Spreadsheet Applications
  - Recommended Spreadsheet Applications (Paid Usage)
  - Recommended Spreadsheet Applications (Free)
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- Browser
  - Krona supports most browsers however the developers suggest using FireFox.

- Tree Viewer
  - Recommended Tree Viewer (Local Installation)
    - MEGA: [http://www.megasoftware.net/](http://www.megasoftware.net/)
  - Recommended Tree Viewer (Web Based)
    - ETE Toolkit Tree Viewer: [http://etetoolkit.org/treeview/](http://etetoolkit.org/treeview/)

- Multiple Sequence Alignment Viewer
  - Recommended MSA Viewer (Local Installation)
    - MEGA: [http://www.megasoftware.net/](http://www.megasoftware.net/)
  - Recommended MSA Viewer (Web Based)
    - MView: [http://www.ebi.ac.uk/Tools/msa/mview/](http://www.ebi.ac.uk/Tools/msa/mview/)
  - For a more comprehensive list of options, see the Wikipedia article on the topic: [http://en.wikipedia.org/wiki/List_of_alignment_visualization_software](http://en.wikipedia.org/wiki/List_of_alignment_visualization_software)

Version Changelog

Current Version

Version 2.3.2 (04/15/2019)
- Added data descriptions for PacBio and illumina sequencer
- Removed references to 454 and IonTorrent data formats

Previous Versions

Version 2.3.1 (08/01/2016)
- Corrected the Paired-End illustrations.
- Updated the FAQ.

Version 2.3 (04/29/2016)
- Added reverse primer information to the FASTA archive.
- Updated the analysis pipeline graphic.
- Added the database maintenance policy.
• Added the FAQ (Frequently Asked Questions)
• Updated the file descriptions to add the new oligos files we provide.

**Version 2.2.4  (11/07/2014)**

• Updated the file descriptions to include the file type and recommended program for viewing the data.
• Updated the file descriptions to include all files provided by RTL.
• Added new section to cover recommended programs for viewing provided data.

**Version 2.2.3  (09/16/2014)**

• Updated the data archive to split files too large to fit on our webserver.
• Included instructions for how to handle split zip archives.

**Version 2.2.2  (09/03/2014)**

• Added the OTUs folder to the Analysis archive.
• Moved OTUmap.txt from Analysis/OTUMap.txt to Analysis/OTUs/OTUMap.txt
• Added OTUs.fas to the Analysis/OTUs archive.
• Corrected the otus.tre file. All ‘;’ within sequence definitions have been changed to ‘_’.

**Version 2.2.1  (08/29/2014)**

• Added the customer data retention policy.

**Version 2.2.0  (07/09/2014)**

• Added phylogenetic tree construction using MUSCLE and FastTree.
• Added Krona visualization to the Taxonomic Analysis pipeline.
• Added phylogenetic tree, multiple sequence alignment, and Krona visualizations to the analysis zip archive.
• Updated 454 and Ion Torrent PGM processing to run using the same workflow as MiSeq.
• Added description for the OTUMap.txt file in the Analysis Folder.

**Version 2.1.1  (05/20/2014)**

• Updated OTU Selection. Trimming to shortest sequence now performed before UPARSE OTU Selection.
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Version 2.1.0  (02/28/2014)

- Updated denoiser to use PEAR for paired-end read merging in place of USEARCH.

Version 2.0.0  (01/28/2014)

- Updated denoiser to use USEARCH 7, replacing USEARCH 5.
- Methodology now accounts for processing of 454, Ion Torrent PGM and Illumina MiSeq data.

References


