QC and downstream analysis of the Rp-Bp results

Rp-Bp includes a number of additional scripts for quality control and downstream analysis.

- Creating read length-specific profiles
- Preprocessing analysis
 - Counting and visualizing reads filtered at each step
 - Creating and visualizing read length distributions
 - Visualizing read length metagene profiles
- Predictions analysis
 - Counting and visualizing the predicted ORF types

Creating read length-specific profiles

As described in the usage instructions, Rp-Bp writes the unsmoothed ORF profiles to a matrix market file. This profile merges reads of all lengths.

The create-read-length-orf-profiles script can be used to create profile files which also include counts of individual read lengths.

create-read-length-orf-profiles <config> <sample or condition name> <out> [--is-condition]

Command line options

- config. A yaml config file
- sample or condition name. The name of either one of the riboseq_samples or riboseq_biological_replicates from the config file
- out. The output (txt.gz) file, containing the read-length specific profiles. The format is a sparse coordinate format inspired by the matrix market format. See below for details about the output format.
- [--is-condition]. If the sample or condition name is a condition, that is, if it is a key from riboseq_biological_replicates, then this flag must be given.

Additionally, the command can be given logging and slurm options.

Output format

Each line in the output file is a tuple containing the following values.

- read_length. The (trimmed) read lengths for this position.
- orf_num. An identifier which maps to orf_num in the (static) list of ORFs
 for the reference, <genome_base_path>/transcript-index/<genome_name>.genomic-orfs.<orf_note>
- orf_position. The base-0 position with respect to the spliced transcript (so position % 3 == 0 implies the position is in-frame)
- read_count. The sum of counts across all replicates for the condition (if --is-condition is given) or the single sample (otherwise) after adjusting according to P-sites and removing multimappers.

Preprocessing report

The create-rpbp-preprocessing-report script can be used to create several plots which summarize the preprocessing and ORF profile construction. The script creates all of the following plots and generates a latex document including all of them.

- Counting and visualizing reads filtered at each step
- Creating and visualizing read length distributions
- Visualizing read length metagene profiles

Optionally, the script can also call FastQC. See more details below.

create-rpbp-preprocessing-report <config> <out> [--show-orf-periodicity] [--show-read-lengt]

- config. A yaml config file
- out. A *directory* where the latex report will be created. If the directory does not exist, it will be created.
- [--show-orf-periodicity]. If this flag is present, metagene periodicity plots will be created for ORFs of each type. (This is similar to Figure S2 in the supplement, although this will include all ORFs of the respective type, regardless of whether they are predicted as translated or not.) These plots can be quite time-consuming to create.
- [--show-read-length-bfs]. If this flag is present, plots showing the Bayes factor for each possible P-site offset for each read length will be included.
- [--overwrite]. By default, if an image file is already present, it will not be recreated. If this flag is given, any existing images will be overwritten.
- [--min-visualization-count]. The minimum number of reads of a given length necessary to include the relevant plots for that read length in the report. Default: 500

- [--image-type]. The extension for the image files. Matplotlib uses this to guess the type of the images. Default: eps. Other common types: png, pdf.
- [--note]. An optional note to include in image file names. This takes precedence over the note specified in the config file.
- [--num-cpus]. The number of samples to process at once.
- [--create-fastqc-reports]. If this flag is present, the FastQC reports described below will be created. This can be rather time-consuming.
- [--tmp]. A temp location for FastQC. It is not used by any of the other reporting scripts.

FastQC reports

If the -c/--create-fastqc-reports flag is given, then FastQC reports will be created for the following files for each sample.

- Raw data. Files from riboseq_samples in the config file.
- Trimmed and filtered reads. <riboseq_data>/without-adapters/<sample-name>[.<note>].fastq.gz
- Reads aligning to ribosomal sequences. <riboseq_data>/with-rrna/<sample-name>[.<note>].fastq.g
- Reads not aligning to ribosomal sequences. <riboseq_data>/without-rrna/<sample-name>[.<note>].f
- Reads aligned to the genome. <riboseq_data>/without-rrna-mapping/<sample-name>[.<note>].bam
- Reads uniquely aligned to the genome. <riboseq_data>/without-rrna-mapping/<sample-name>[.<not

Counting and visualizing reads filtered at each step

Counting

The get-all-read-filtering-counts script counts reads filtered at each step of the preprocessing pipeline.

This script requires samtools to be present in \$PATH.

get-all-read-filtering-counts <config> <out> [--num-cpus <num_cpus>]

- config. A yaml config file
- out. The output file, in csv.gz format. See below for details.
- [--num-cpus]. The script is parallelized at the sample level. If specified, this many samples will be processed at once.

Output format

The output is a "wide" data frame which contains one row for each sample. The fields are as follows.

- note. The name of the sample.
- raw_data_count. The number of reads in the original fastq files.
- without_adapters_count. The number of reads remaining after running flexbar to remove adapters and low-quality reads.
- without_rrna_count. The number of reads remaining after removing ribosomal and other reads with bowtie2.
- genome_count. The number of reads with at least one genome alignment.
- unique_count. The number of reads with exactly one genome alignment.
- length_count. The number of uniquely mapping reads which also have a "periodic" read length, as determined by BPPS.

Visualizing (script)

The visualize-read-filtering-counts script visualizes the read counts from get-all-read-filtering-counts.

visualize-read-filtering-counts <read_counts> <out> [--without-rrna] [--title <title>] [--fo

- read_counts. The output from get-all-read-filtering-counts
- out. The output image file. The extension should be something recognized by matplotlib, such as png or pdf.
- [--without-rrna]. If this flag is given, then the bar chart will not include reads filtered due to low quality or mapping to ribosomal sequences.
- [--title]. A title placed at the top of the plot
- [--fontsize]. The fontsize used for most of the text on the plot, including the tick labels (sample names and read counts), axis labels and title.
- [--legend-fontsize]. The fontsize to use for the entries in the legend (the filtering steps).
- [--ymax]. The maximum number of reads displayed on the y-axis. Typically, this value should be around 10% higher than the largest read count. However, some other value may be more appropriate if one of the samples has many more reads than the others.
- [--ystep]. The frequency of tick marks on the y-axis.

Visualizing (ipython notebook)

The notebooks/preprocessing/create-read-filtering-bar-chart notebook can be used to visualize the read counts. It functionality is essentially the same as the visualize-read-filtering-counts script; however, the properties of the plot, such as the exact location of the legend, are much easier to manipulate in the notebook.

Additionally, the notebook will attempt to use the riboseq_sample_name_map from the config file to find "pretty" names for the samples. In particular, this should be a map from the sample name given in the riboseq_samples to a string that will be used for the x-tick labels in the plot. If a sample name is not present in the name map, it will be left unchanged.

Control variables

In the third cell, the config_files, alignment_counts_files, out_files and without_rrna_files dictionaries must be updated to include the relevant files. The key in the dictionary should be the same for all of the new files.

In the fourth cell, the data variable should be changed to the key used in the dictionaries. The other variables (without-rrna, etc.) have the same interpretation as for the script.

In the sixth cell, visualization aspects such as the colors, legend location, figure size, etc., can be set using the respective matplot lib options.

Example visualization

Creating and visualizing read length distributions

Creating distributions

The get-read-length-distribution script (part of the misc package) counts the number of reads of each length in a given bam file. It can be used to count the read length distribution for both all aligned reads and only uniquely-aligning reads.

 ${\bf N.B.}$ The script handles multi-mappers to ensure they only contribute to the counts once.

counts onec.

get-read-length-distribution <bam_1> [<bam_2> ...] -o/--out <length-counts.csv.gz> [-p/--nur

Command line options

• bam_i. The bam files which contain the aligned reads.

- out. The output file, in csv.gz format, which contains the counts. See below for the column specifications.
- [--num-cpus]. The number of CPUs to use; this many files will be processed at once.

Output format

The output is a "long" ("tidy") data frame with the following fields.

- basename. The name of the bam file, excluding the ".bam" extension.
- length. The read length.
- count. The number of reads of that length in the indicated file.

Visualizing the distributions (script)

The plot-read-length-distribution script creates a bar chart of the counts from get-read-length-distribution.

Command line options

- distribution. The csv file created by get-read-length-distribution.
- basename. The basename to visualize.
- out. The output (image) file
- [--title]. The title of the plot.
- [--{min, max}_read_length]. The minimum and maximum read lengths to include in the plot, inclusive. Defaults: [22, 35]
- [--ymax]. The maximum value for the y-axis. Default: 1.5e6+1
- [--fontsize]. The size of the fonts for the title, axis labels and ticks

Visualizing the distributions (ipython notebook)

The notebooks/preprocessing/create-read-length-distribution-bar-chart notebook can be used to visualize the read counts. Its functionality is essentially the same as plot-read-length-distribution; however, the properties of the plot, such as the colors, are much easier to manipulate in the notebook.

Additionally, the notebook will attempt to use the riboseq_sample_name_map from the config file to find "pretty" names for the samples. In particular, this should be a map from the sample name given in the riboseq_samples to a

string that will be used for the x-tick labels in the plot. If a sample name is not present in the name map, it will be left unchanged.

Control variables

All of the relevant control variables in the third cell should point to the appropriate files.

Example visualization

Visualizing read length metagene profiles

As described in the usage instructions, metagene profiles for each read lengths are created as a part of the pipeline. These can be visualized with the create-read-length-metagene-profile-plot script. In particular, it shows the reads aligned around the annotated translation initiation and termination sites.

create-read-length-metagene-profile-plot <metagene_profile> <length> <out> [--title <title>]

- metagene_profile. The metagene profile file (<riboseq_data>/metagene-profiles/<sample-name>[.<
- length. The length to visualize
- out. The output (image) file
- [--title]. A title for the plot
- [--xlabel-{start,end}]. Messages to include beneath the x-axis around the initiation and termination sites, respectively. Defaults: "Position of P-site relative to start (nt)\nRed: TIS. Green: TIS -12", "Position of P-site relative to stop (nt)\nBlue: Translation termination"
- [--ylabel]. The label for the y-axis. Default: "Read count (starting at bp x)"
- [--step]. The step size for the x-axis. Default: 10
- [--font-size]. The font size for everything in the plot. Default: 15
- [--{start,end}-{up,down}stream]. The position (in bp) to start and end the visualization around the translation initiation (start) and terminiation (end) sites. N.B. The upstream options must be negative, and the downstream options must be position. Defaults: [-50, 21] (for both)
- [--use-entire-profile]. If this option is given, then the entire profile in the file will be used, rather than the positions given by the other options.

There is not currently an ipython notebook to create these plots.

Example visualization

Predictions report

The create-rpbp-predictions-report script can be used to create several plots which summarize the predictions made by Rp-Bp. The scripts creates the following plots and generates a latex document including all of them.

- Predicted ORF types bar chart
- Predicted ORF types length distributions (Not documented yet)
- [Predicted ORF types metagene profiles(#predicted-orf-types-metageneprofiles) (Not documented vet)

create-rpbp-predictions-report <config> <out> [--show-unfiltered-orfs] [--show-orf-periodic;

- config. A yaml config file
- out. A *directory* where the latex report will be created. If the directory does not exist, it will be created.
- [--show-unfiltered-orfs]. By default, only the "filtered" ORF predictions (longest ORF at each stop codon and highest Bayes factor among overlapping ORFs; see "Final prediction set" in the paper). If this flag is given, then additional plots will be included showing the relevant statistics for all ORFs predicted as translated. Typically, the canonical_truncated type dominates these plots, so they are often not informative.
- [--show-orf-periodicity]. If this flag is present, metagene periodicity plots will be created for ORFs predicted as translated of each type. (This is similar to Figure S2 in the supplement.) These plots can be somewhat time-consuming to create, especially if the --show-unfiltered-orfs flag is given.
- [--show-chisq]. As described in the usage instructions, the pipeline also makes predictions using a simple chi square test. This is very similar to the ORFscore [Bazzini et al., The EMBO Journal, 2014]. If this flag is given, then all plots will be created using both the Rp-Bp and the chi square predictions (filtered and unfiltered for both, if the --show-unfiltered-orfs flag is given).
- [--uniprot]. Optionally, the ORF type length distributions can include the distribution of Uniprot (or other "reference") transcript sequences.

If given, then the KL-divergence will be calculated between the length distributions. This is similar to Figure S3 in the paper, though the ORFs will be split by type.

This should be a tab-delimited file which includes at least the fields "Status" and "Length". For the paper, we created this file on the UniProtKB by filtering on the relevant organism and using an identity of "90%" for the protein clusters (under the "UniRef" heading on the left panel on the UniProtKB results page).

- [--uniprot-label]. The label to use for the --uniprot sequence lengths, if they are given.
- [--image-type]. The extension to use for the image files. This must be something matplotlib can interpret. The figures do not include large scatter plots, etc., so the default is probably fine. Default: pdf. Other common types: eps, png
- [--note]. An optional note to include in image file names. This takes precedence over the note specified in the config file.
- [--overwrite]. By default, if an image file is already present, it will not be recreated. If this flag is given, any existing images will be overwritten.
- [--num-cpus]. The number of samples to process at once.

Predicted ORF types bar chart

The create-orf-types-bar-chart and create-orf-types-pie-chart scripts can be used to show the count of each type of ORF in a given bed file (which includes the orf_type field). For example, this can be used for both the filtered and unfiltered prediction files.

Both scripts show the number of ORFs of each type on both strands. Typically, there should not be a strong bias between the strands.

```
create-orf-types-bar-chart <orfs> <out> [--title <title>] [--use-groups] [--legend-fontsize
create-orf-types-bar-chart <orfs> <out> [--title <title>] [--use-groups]
```

Command line options

The shared command line options are the same for both scripts.

- $\bullet\,$ orfs. The bed file containing the ORFs
- out. The image file
- [--title]. A title for the plot

- [--use-groups]. If this flag is present, then ORF types will be combined as described in the supplement of the paper. In particular, the following groups are used:
 - Canonical: canonical
 - Canonical variant: canonical_extended, canonical_truncated
 - uORF: five_primedORF: three_primencRNA: noncoding
 - Other: five_prime_overlap, suspect_overlap, three_prime_overlap,
 - de novo only: novel
 - de novo overlap: all other "novel" types
- [--{legend-}fontsize]. The fontsize to use in the respective places in the bar chart. Default: 15, 20
- [--ymax]. The maximum value for the y-axis in the bar chart. Default:1e4

ipython notebooks

The notebooks/rpbp-predictions/create-orf-type-{bar,pie}-chart.ipynb notebooks can be used to create the same plots. The relevant variables in the third cell should be updated. The notebooks allow easier control over the colors, etc.

Example visualizations