



FASTAptameR 2.0 - User Interface Tutorial

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1 Introduction

FASTAptameR 2.0 is an R-based update of FASTAptamer (Alam KK 2015). Like its predecessor, FASTAptameR 2.0 is an open-source toolkit designed to analyze populations of sequences resulting from combinatorial selections. This updated version features a user interface (UI), interactive graphics, more modules, and a faster implementation of the original clustering algorithm.

This user guide walks you through installation and each of the analytical modules, and it highlights what options you have when analyzing your data through the UI.

1.1 Overview

- **FASTAptameR-Count**
 - *This module is the entry point into FASTAptameR 2.0*
 - Input: preprocessed FASTQ/A
 - Workflow:
 1. count all unique sequences (**Reads**)
 2. sort by counts (**Rank**)
 3. normalize counts as reads per million (**RPM**)
 - Plotting:
 1. line plot of reads-per-rank
 2. histograms of sequence lengths - one for the unique sequences and one for all reads
 3. sequence abundance bar plot
 - Output: FASTA or CSV
- **FASTAptameR-Translate**
 - Input: FASTA from FASTAptameR-Count, which will be referred to as a “counted FASTA”
 - Workflow: translate D/RNA sequences to amino acid sequences
 - Plotting:
 1. line plot of reads-per-rank
 2. histograms of sequence lengths - one for the unique sequences and one for all reads
 - Output: FASTA or CSV
- **FASTAptameR-Motif_Search**
 - Input: counted FASTA and comma-separated patterns
 - Workflow: search for user-defined patterns in sequences
 - Output: FASTA or CSV
- **FASTAptameR-Motif_Tracker**
 - Input: at least two counted FASTAs and query list
 - Workflow: track how user-defined motifs or sequences change across populations
 - Plotting: line plot of each query’s RPM across the populations
 - Output: CSV
- **FASTAptameR-Distance**
 - Input: counted FASTA and query sequence
 - Workflow: compute the Levenshtein edit distance (LED) between a single query sequence and all other provided sequences
 - Plotting: histograms of edit distances - one for the unique sequences and one for all reads
 - Output: FASTA or CSV
- **FASTAptameR-Enrich**

- Input: at least two counted or clustered FASTAs
 - Workflow: calculate how each sequence enriches across populations
 - Plotting:
 1. bar plot of sequence persistence
 2. histogram(s) of $\log_2(\text{Enrichment})$
 3. scatter plot(s) of RPM
 4. MA plot, displaying average log-RPM (A) and fold change (M)
 5. box plot(s) of sequence enrichment per cluster; only available if clustered FASTAs are provided
 - Output: CSV
- **FASTAptameR-Position_Enrich**
 - Input: enrichment CSV and reference sequence
 - Workflow: for each position of the reference sequence, compute the average enrichment of non-reference residues in the data
 - Plotting:
 1. bar plot of average enrichment per position of reference sequence
 2. heat map of average enrichment per position of reference sequence, further grouped by residues
 - Output: None
- **FASTAptameR-Cluster**
 - Input: counted FASTA
 - Workflow:
 1. filter out low-read sequences based on user-defined input
 2. treat most abundant, non-clustered sequence as cluster seed
 3. add all sequences within a user-defined LED of the seed to the cluster
 4. Repeat until all sequences are clustered or a maximum number of clusters are created
 - Output: FASTA or CSV
- **FASTAptameR-Cluster_Diversity**
 - Input: clustered FASTA
 - Workflow: provide metadata for each cluster
 - Plotting:
 1. metaplots for count of unique sequences, count of total reads, and average LED per cluster
 2. k-mer PCA plot, colored by cluster identity
 - Output: CSV
- **FASTAptameR-Cluster_Enrich**
 - Input: at least two cluster-analysis CSVs
 - Workflow: calculate how each cluster enriches across populations
 - Plotting: line plot of each the total RPM per cluster for each seed per population
 - Output: CSV

Fig. 1 gives an overview of how these modules connect to one another. Additionally, a summary of input and output file types is given in **Table 1**. Please note that each module requires the user to upload a file or, in the case of **FASTAptameR-Count**, optionally provide a GitHub link to the data. At present, none of this data will “live” on the server to be passed between modules.

Table 1: Module Inputs and Outputs

| Module | Input Files | Output Files |
|-----------------------|----------------------|--------------|
| FASTAptameR-Count | Preprocessed FASTQ/A | FASTA or CSV |
| FASTAptameR-Translate | Counted FASTA | FASTA or CSV |

| Module | Input Files | Output Files |
|-----------------------------------|--------------------------------|--------------|
| FASTAptameR-Motif_Search | Counted FASTA | FASTA or CSV |
| FASTAptameR-Motif_Tracker | 2+ counted FASTAs | CSV |
| FASTAptameR-Distance | Counted FASTA | FASTA or CSV |
| FASTAptameR-Enrich | 2+ counted FASTAs | CSV |
| FASTAptameR-Positional_Enrichment | Enrich CSV | Plots |
| FASTAptameR-Cluster | Counted FASTA | FASTA or CSV |
| FASTAptameR-Cluster_Diversity | Clustered FASTA | CSV |
| FASTAptameR-Cluster_Enrich | 2+ CSVs from Cluster_Diversity | CSV |

Many function inputs / outputs are simply FASTA files, so FASTAptameR 2.0 can be easily integrated into most analytical pipelines. Note that *counted* FASTA files are the minimum input for most modules (*e.g.*, FASTAptameR-Translate needs *at least* a counted FASTA but could also accept a searched or clustered FASTA file).

Importantly, FASTAptameR 2.0 does not provide any functions that are easily addressed by other software (*e.g.*, merging paired-end reads, trimming constant regions, predicting structures, *etc.*). Rather, the focus of this application is to provide flexible downstream analyses for the selections field.

1.2 A note on plotting

Though many plots are initially created with `ggplot2`, they are all shown as interactive `plotly` plots. As such, you will see a number of options appear along the top of the image in response to your mouse hover. These options will allow you to zoom in and out, select regions of interest, and, most importantly, download the plots. This last functionality is provided by the camera icon (1st icon on the left as you hover over the image). Finally, double-clicking the image should reset it (*e.g.* remove zoom or crop effects).

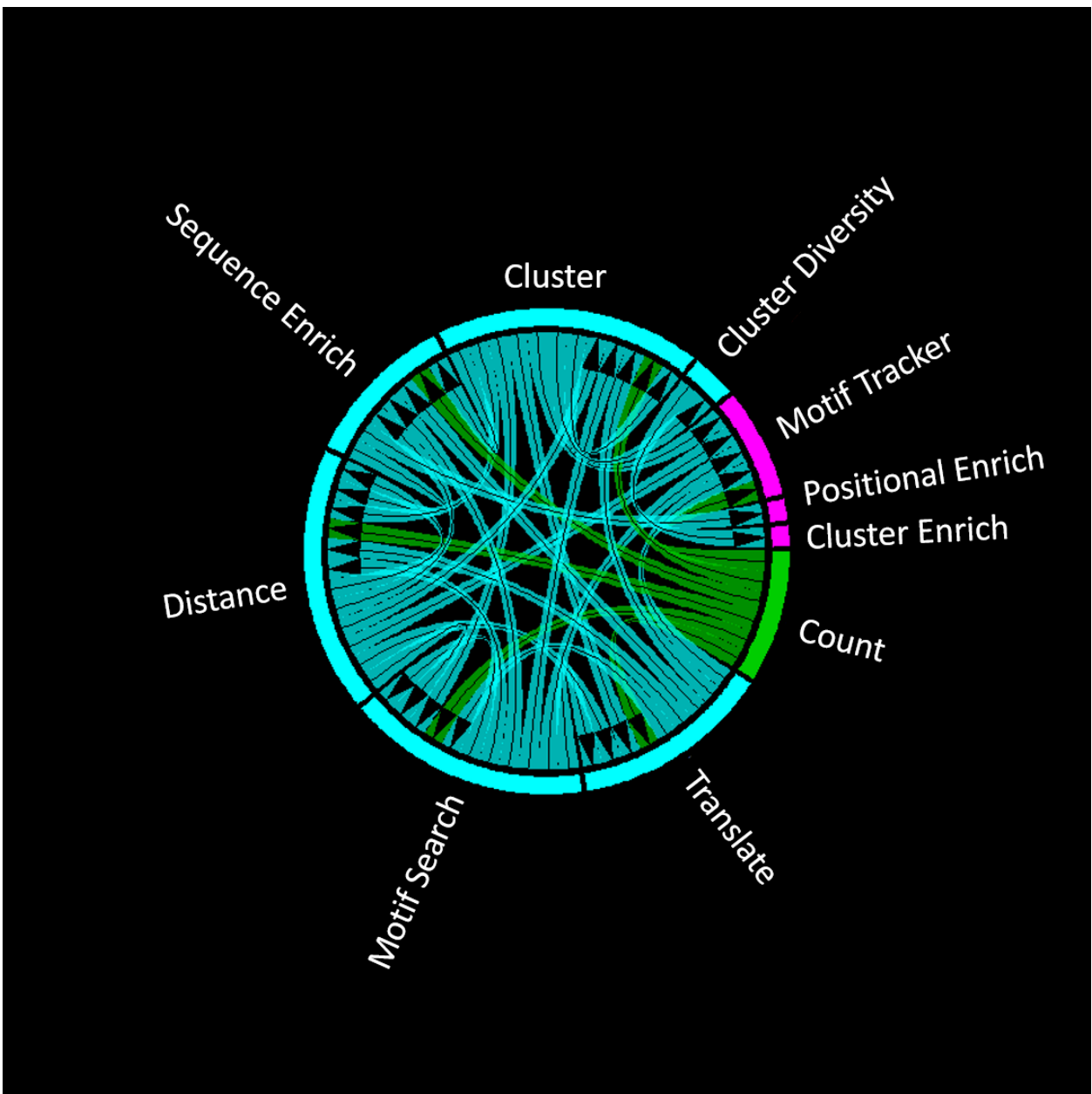


Figure 1: Screenshot of Module Connections.

2 How to get started

Exactly like its predecessor, FASTAptameR 2.0 is designed to be **easy** to use. There are three ways for users to interact with FASTAptameR 2.0. The *web server* is the easiest way to interact with this application because it only requires an internet connection and browser. The user interface can also run on your local system as a *Docker container*. Finally, all code can be pulled from GitHub at <https://github.com/SkylerKramer/FASTAptameR-2.0>.

2.1 Web server interface

This is the easiest way to use the FASTAptameR 2.0 UI. The web server can be accessed from <https://fastaptamer2.missouri.edu/>, which is hosted by the Digital Biology Laboratory under the direction of Dr. Dong Xu. However, this option only works if the files you need to upload are less than 2 GB.

2.2 Docker interface

The web server can also be run locally on your machine(s) via Docker. This interface functions identically to the web server variant and does not depend on the strength of your internet connection to run. In brief, Docker is a convenient tool that may be used to construct *images* of software. The *image* essentially functions as the blueprint for an application. The *image* of FASTAptameR 2.0, for example, contains all relevant software (*e.g.*, R), files (*e.g.*, this PDF), and packages (*e.g.*, Shiny).

Here are the **three steps** required for initial installation of FASTAptameR 2.0 via Docker. **Step 1** is required for the initial installation of Docker. **Step 2** is required to get access to FASTAptameR 2.0 and all subsequent versions. **Step 3** is required to run the application.

1. Install Docker on a Linux machine or install Docker Desktop on a Windows or Mac machine to establish a “Docker-active terminal” on your local system.
2. Pull the FASTAptameR 2.0 image from the Docker Hub repository.
3. Execute the `docker run ...` command and navigate to `localhost:3838` in your preferred web browser.

Step 1. Install Docker to establish a “Docker-active terminal” on your local system.

For extensive details on Docker or its installation, please see <https://www.docker.com/> and <https://docs.docker.com/get-docker/>, respectively. Importantly, the FASTAptameR 2.0 *image* is built on Linux. Thus, it is necessary to run it from a Linux environment or virtual machine. This should naturally happen after installing Docker for Linux or Docker Desktop for Mac. Windows users must also install Docker Desktop and will typically need to enable virtualization through their Bios. Successful completion of this step will yield a “Docker-active terminal”. Linux, Windows, or Mac users can check if their terminal is Docker-active by running the following command, which should return which version of Docker has been installed if it was installed successfully:

```
docker version
```

Step 2. Pull the FASTAptameR 2.0 image from the Docker Hub repository.

The FASTAptameR 2.0 *image* must be pulled from a repository (*i.e.*, Docker Hub) by running the following command in a Docker-active terminal:

```
docker pull skylerkramer/fastaptamer2:publicupload04
```

Please note that this version (`publicupload04`) will change over time. For the most recent version of FASTAptameR 2.0, please refer to the dynamic User Guide at <https://github.com/SkylerKramer/FASTAptameR-2.0>.

FASTAptameR-2.0 or the main Docker Hub page at <https://hub.docker.com/repository/docker/skylerkramer/fastaptamer2>.

Step 3. Launch FASTAptameR 2.0.

Once you have this application's image, run the following from a Docker-active terminal:

```
docker run -d --rm -p 3838:3838 skylerkramer/fastaptamer2:publicupload04
```

This will launch a local instance - a *container* - of FASTAptameR 2.0. You will then interact with this container in the same fashion as the web server by typing `localhost:3838` into your web browser address bar.

Explanation of flags from Step 3:

- `-d`: enable detached mode, which allows you to use your command line / terminal even with the active *container* (*i.e.*, *container* is detached from your terminal and runs in the background)
- `--rm`: automatically stop the container upon exit
- `-p 3838:3838`: publish 3838 host port (1st number) to the 3838 container port (2nd number)
- `skylerkramer/fastaptamer2:publicupload04`: the local path to the **FASTAptameR 2.0** Docker *image*; please see **Step 2** for the most recent version
- `localhost:3838`: navigate here from your web browser to start interacting with **FASTAptameR 2.0**

2.3 R interface

All code is publicly available at <https://github.com/SkylerKramer/FASTAptameR-2.0>. This will allow R developers to adjust the code to their specific needs. Please note that all dependencies in this app (*e.g.*, Shiny, ggplot2, *etc.*) must be installed to use this app. A full list of dependencies and their installation instructions are available on the GitHub page.

2.4 Software usage

If you use, adapt, or modify FASTAptameR 2.0, please cite: **CITATION**.

For any questions or concerns, please email burkelab@missouri.edu or stk7c9@umsystem.edu.

3 Tutorial

3.1 Data requirements

FASTAptameR 2.0 utilizes many string-based functions in its backend. Thus, this program can be used to analyze many types of biological populations. However, all libraries must be initially saved in a FASTA or FASTQ format and passed through **FASTAptameR-Count** prior to any subsequent analyses. Further, any data preprocessing steps, such as trimming flanking sequences or filtering for read quality, must be made outside of this application.

3.2 Sample Data and Uploading User Data

All data shown in this tutorial come from the 14th and 15th rounds of an aptamer selection against HIV-1 reverse transcriptase (Burke DH 1996; Ditzler MA 2013; Whatley AS 2013). These data are preprocessed (trimmed and filtered) and available from <http://burkelab.missouri.edu/fastapamer.html>.

To start analyzing the sample data or your own data, please do one of two things. Either 1) upload a local copy of the file via the file browser in **FASTAptameR-Count** or 2) supply a link to the data via the text box labeled as **Online source** in **FASTAptameR-Count**. This module is the entry point to **FASTAptameR 2.0**, so each analysis should start here.

3.3 FASTAptameR-Count

3.3.1 Description

FASTAptameR-Count serves as the entry point into this suite of modules, and, thus, it should be run prior to any of the following modules. This function accepts either a FASTQ/A file chosen with the file browser or a link to such a file (*e.g.*, the default GitHub link in the text box labeled as **Online source**) and returns a *counted* data table as output that can be downloaded as a FASTA or CSV file.

Input FASTQ files should be properly formatted (4 lines per entry with the 2nd line of each entry being the sequence). Input FASTA files are not required to have sequence identifiers. No pre-existing sequence identifiers will be conserved by this module. Instead, output sequence identifiers are defined by the statistical representation of each sequence. Sample input files are shown in **Fig. 2**. A screenshot of the module interface is shown in **Fig. 3**.

A) FASTA sequence without identifiers:

```
@HWI-700167R:5:11:C3N2DAXX:5:1101:4016:2214 1:N:0:GACGGCT  
AGCGCGCACCCAAAATCGAAATCCGAAGCGAACGGGAGAATCGGACCAAAGATAACCTGTGAATGGC  
AGCGCGCACCCAAAATCGAAATCCGAAGCGAACGGGAGAATCGGACCAAAGATAACCTGTGAATGGC  
ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTGAGACTGCCACGCTTGGTGT  
ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTGAGACTGCCACGCTTGGTGT  
ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTGAGACTGCCACGCTTGGTGT  
ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTGAGACTGCCACGCTTGGTGT
```

B) FASTA sequence with identifiers:

```
@HWI-700167R:5:11:C3N2DAXX:5:1101:4016:2214 1:N:0:GACGGCT  
AGCGCGCACCCAAAATCGAAATCCGAAGCGAACGGGAGAATCGGACCAAAGATAACCTGTGAATGGC  
+  
IIJJJJJJJJJJJJJJJJJJJJHHFF<ADBDDDDDDDDDDCDDDDDDDDDCACCDCDDDECD  
@HWI-700167R:5:11:C3N2DAXX:5:1101:4997:2191 1:N:0:GACGGCT  
ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTGAGACTGCCACGCTTGGTGT  
+  
IIJJJJGIIJJJJJJJJJJJJJJJJJJHHFFEEEDDDDDDDDDDDDBB@CDD
```

C) FASTQ file:

```
@HWI-700167R:5:11:C3N2DAXX:5:1101:4016:2214 1:N:0:GACGGCT  
AGCGCGCACCCAAAATCGAAATCCGAAGCGAACGGGAGAATCGGACCAAAGATAACCTGTGAATGGC  
+  
IIJJJJJJJJJJJJJJJJJJJJHHFFEEEDDDDDDDDDDDDBB@CDD
```

Figure 2: Valid inputs to FASTAptameR-Count. A) FASTA without sequence identifier lines. B) FASTA with sequence identifier lines. C) FASTQ file.

Choose data to count*:

A) FASTQ or FASTA file

*Do not start until loading bar shows 'Upload complete'.

Online source:

B)

FASTA or CSV download?

FASTA CSV

Min. number of reads to plot:



Max. rank to plot:



Adjust default bins?

Yes No

Figure 3: Screenshot of FASTAptameR-Count.

3.3.2 Usage

The input FASTQ/A file must be chosen with the file browser (**Fig. 3A**) or linked in the text box (**Fig. 3B**). A sample link is already provided in the text box. Note, if a file is uploaded via the file browser **AND** a link is provided, only the uploaded file (**NOT** the linked one) will be analyzed.

The **Start** button will begin the counting process. The results will be displayed as a data table on the right side of the screen. For file uploads, please wait for the loading bar to show *Upload complete* before using the **Start** button.

A sample output data table is shown in **Fig. 4**.

| Show 10 entries | | | | | | Search: |
|---------------------|------|--------|-----------|--------|--|---------|
| id | Rank | Reads | RPM | Length | seqs | |
| All | All | All | All | All | All | |
| >1-417696-193358.44 | 1 | 417696 | 193358.44 | 70 | ACGTTGTCGAAAGCCTATGCAAATTAGGACTGTGCGACGAAACCTTGCCTAGACTGCCACGCTTGGTGT | |
| >2-313312-145037.35 | 2 | 313312 | 145037.35 | 70 | CATAGCGACTGTCCACGAACTCGAACGCCAACGGGACAAAAGGCAAGAGCGCGATAACCATGCTGGACTG | |
| >3-174096-80591.94 | 3 | 174096 | 80591.94 | 70 | AACCGCAAGCAACACCCAGCAAGAACATCCGACGCACGACGGGAGAAAGTCGATTACACGATGTCGAT | |
| >4-94978-43966.9 | 4 | 94978 | 43966.9 | 70 | CATAGCGACTGCCACGAATCCGAAAGCCTAACGGGACAAAAGGCAAGAGCGCGATAACCATGCTGGACTG | |
| >5-74389-34435.91 | 5 | 74389 | 34435.91 | 70 | ACGTTGTCGAAAGCCTATGCAAATTAGGACTGTGCGACGAAACCTTGCCTAGACTGCCCGCTTGGTGT | |
| >6-57625-26675.57 | 6 | 57625 | 26675.57 | 69 | CCCTCCTGTATGACGCTAACTGAGAATCGAACGGGAGAAAGGACACTTATGACGTGGCG | |
| >7-53608-24816.04 | 7 | 53608 | 24816.04 | 70 | ACGTTGTCGAAAGCCTATGCAAATTAGGACTGTGCGACGAAACCTTGCCTAGACTGCCATGCTTGGTGT | |
| >8-39793-18420.84 | 8 | 39793 | 18420.84 | 69 | AGCGCGCACCCAAATCGAAATCGAACGGGAAACGGGAGATGCGACCAAAGATAACCTGTGAATGGC | |
| >9-33800-15646.58 | 9 | 33800 | 15646.58 | 70 | TTGACAATAACTCGAGAACCGAGGTGCAAACGGGAGAACACAATGGATTACACCGAGCTGGCTGAC | |
| >10-29794-13792.14 | 10 | 29794 | 13792.14 | 70 | GCGAACCAAACCCAGATTACTAACCGTGGGCTGAAACACGGGACAAAAGGCGATCAATGGAGTGTAC | |

Showing 1 to 10 of 72,921 entries

Previous 1 2 3 4 5 ... 7293 Next

Figure 4: FASTAptameR-Count Output.

Note that the *id* column has the following format: >Rank-Reads-RPM, where **Rank** is the order of sequences after sorting by **Reads**, which is the raw abundance of each sequence. **RPM** (*i.e.*, Reads per Million) is the value of **Reads**, normalized by the total population size: **RPM** = **Reads** / (populationSize / 1e6).

The total number of sequences, number of unique sequences, and module runtime will be displayed below the **Start** and **Download** buttons after running is finished. The **Download** button opens a file browser prior to downloading the output as a FASTA or CSV file (DEFAULT = FASTA, which is required for subsequent modules). Keep the *count* file in an easily accessible folder, as this file will serve as input for many other FASTAptameR 2.0 modules.

3.3.3 Troubleshooting

If you start the module before the upload is finished **AND** a file link is provided, then the file link will be analyzed. If you start the module before the upload is finished **AND** no link is provided, then you will get an error that says **No file or link provided!**. If you start the module before the upload is finished **AND** no file link is provided **AND** you have previously uploaded a file to this module, then the previously

uploaded file will be reanalyzed. In any of these cases, reuploading the file, *waiting for it to finish uploading*, and then starting the module should correctly analyze your data. If any errors persist, please refresh the page.

3.3.4 Plotting

This module can also generate three types of interactive plots based on the counted data: a line plot of reads-per-rank (**Fig. 5A, B**), two histograms of sequence lengths (**Fig. 5C**), and a sequence abundance bar plot (**Fig. 5D**). Line plots are filterable by 1) minimum number of reads to plot and 2) maximum rank to plot, and both values are chosen with a slider bar. The histograms, however, are not filterable.

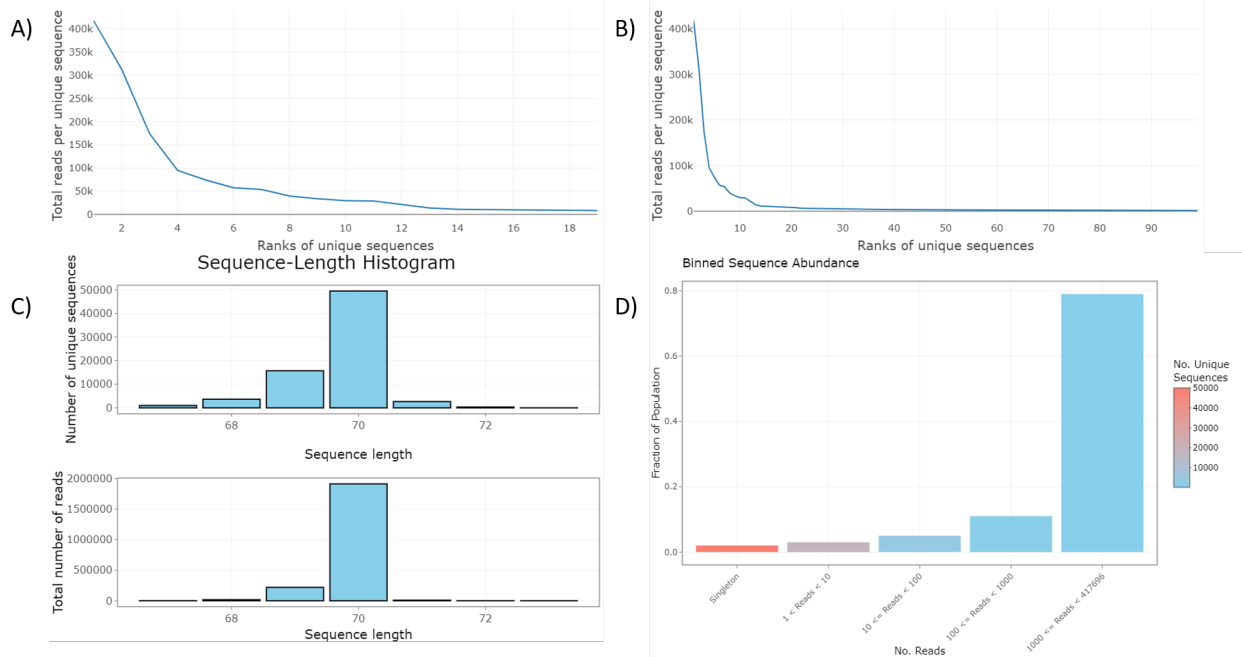


Figure 5: FASTAptameR-Count Plots. A) A line plot showing the total number of reads for the 20 most abundant sequences. B) The same plot with the 100 most abundant sequences. C) Histograms of sequence lengths for unique sequences (top) and all reads (bottom). D) Binned sequence abundance bar plot where x corresponds to discrete bins of read counts, y corresponds to the fraction of the total population, and color corresponds to the number of unique sequences per bin.

The sequence abundance bar plot first bins sequences based on their read counts and then plots these bins against their relative abundance (as fractions of the total population). Finally, the bars are colored according to the number of unique sequences in each bin. These breakpoints forming these bins are, by default, set to the following: `Reads = 1, 1 < Reads < 10, 10 <= Reads < 100, 100 <= Reads < 1000, 1000 <= max(Reads)`. However, the user can toggle singletons on/off or alter these breakpoints by selecting Yes to the respective prompt. New breakpoints should be entered as a comma-separated list.

3.4 FASTAptameR-Translate

3.4.1 Description

FASTAptameR-Translate translates input nucleotide sequences into amino acid sequences following the standard genetic code, treating the input nucleotide sequences as positive-sense mRNA. This module accepts a *counted* FASTA file and returns a *translated* data table that can be downloaded as a FASTA or CSV file. A screenshot of the module interface is shown in **Fig. 6**.

3.4.2 Usage

The input FASTA file must be chosen with the file browser. The open reading frame may be selected by the 1st set of radio buttons (`DEFAULT = 1`) (**Fig. 6A**). The 2nd set of radio buttons indicates whether nucleotide sequences that encode the same amino acid sequence should be merged (`DEFAULT = Yes`) (**Fig. 6B**). If `Yes`, then redundantly encoded amino acid sequences are converged, and a new column (`Unique.Nt.Count`) will specify how many non-unique nucleotide sequences from the *counted* input were merged into each amino acid sequence. If `No`, then each unique nucleotide sequence is treated separately, even if multiple sequences encode the same amino acid sequence.

The dropdown menu in this UI (**Fig. 6C**) allows the user to select one of sixteen genetic codes for translation (`DEFAULT = Standard`):

1. Standard
2. Vertebrate mitochondrial
3. Yeast mitochondrial
4. Mold, protozoan, and coelenterate mitochondrial + Mycoplasma / Spiroplasma
5. Invertebrate mitochondrial
6. Ciliate, dasycladacean and Hexamita nuclear
7. Echinoderm and flatworm mitochondrial
8. Euplotid nuclear
9. Alternative yeast nuclear
10. Ascidian mitochondrial
11. Alternative flatworm mitochondrial
12. Blepharisma nuclear
13. Chlorophycean mitochondrial
14. Trematode mitochondrial
15. Scenedesmus obliquus mitochondrial
16. Pterobranchia mitochondrial

The user may also alter these codes by selecting `Yes` in the 3rd set of radio buttons (**Fig. 6D**) prior to translating. If `Yes`, then comma-separated codon / translation pairs may be entered in the resulting text box (*e.g.*, GAT,Z). If the codon already exists in the standard genetic code, then the user-supplied mapping will take precedence. If the codon does not exist in the standard genetic code, then it will be added to it. Please note that only 3-letter codons and 1-letter translations are currently accepted.

The `Start` button begins the translation process. The *translated* data table will be shown on the right side of the screen. The `Download` button opens a file browser prior to downloading the output as a FASTA or CSV file (`DEFAULT = FASTA`, which is required for subsequent modules).

3.4.3 Plotting

This module generates the same two types of plots as FASTAptameR-Count: a line plot of reads-per-rank and two histograms of sequence lengths. See that section for more details.

Choose data to translate:

Open reading frame:

A) 1 2 3

Should non-unique sequences be merged?

B) Yes No

Genetic code selection:

C)

Do you want to use non-standard translations?

D) Yes No

FASTA or CSV download?

FASTA CSV

Min. number of reads to plot:

Max. rank to plot:

Figure 6: Screenshot of FASTAptameR-Translate.

3.5 FASTAptameR-Motif_Search

3.5.1 Description

FASTAptameR-Motif_Search identifies sequences that contain one or more user-specified sequence motifs, or ‘patterns.’ The module accepts a *counted* FASTA file and returns a *searched* data table that can be downloaded as a FASTA or CSV file. Sequences in the output must have at least one occurrence of each pattern or at least one occurrence of at least one pattern (see details below for the **partial match** radio button). A screenshot of the module interface is shown in **Fig. 7**.

The screenshot shows the user interface for the FASTAptameR-Motif_Search module. At the top, there are two tabs: "Search" (highlighted in orange) and "Tracker" (in blue). Below the tabs, there is a section labeled "Input data:" with a "Browse..." button (highlighted in black) and a "FASTA file" label. The main area contains several configuration options:

- A)** A text input field labeled "Comma-separated patterns:".
- B)** A question "Place patterns in parentheses in output?" with radio buttons: Yes (unselected) and No (selected).
- C)** A question "If multiple patterns, return partial matches?" with radio buttons: Yes (unselected) and No (selected).
- D)** A question "Type of pattern?" with radio buttons: Nucleotide (selected), AminoAcid (unselected), and String (unselected).
- A question "FASTA or CSV download?" with radio buttons: FASTA (selected) and CSV (unselected).

At the bottom are two buttons: "Start" (highlighted in black) and "Download" (with a download icon).

Figure 7: Screenshot of FASTAptameR-Motif_Search.

3.5.2 Usage

The input FASTA file must be chosen with the file browser. The following text box (**Fig. 7A**) must contain at least one pattern (*e.g.*, AAA). If the user wishes to search for multiple patterns, the patterns must be separated by commas (*e.g.*, AAA, GTG).

The 1st set of radio buttons (**Fig. 7B**) determines whether the output has parentheses set around identified patterns (DEFAULT = No). For example, when **pattern** = GGC and **sequence** = AAAGGCT, the output is AAA(GGC)T. Note, when two or more patterns overlap, output only displays parentheses around the first search term that is matched. For example, when **pattern** = AGGC,GGCT and **sequence** = AAAGGCT, the output is AA(AGGC)T. Note that parentheses will be treated as individual characters by subsequent modules and may alter downstream analyses.

The 2nd set of radio buttons (**Fig. 7C**) governs how the software deals with multiple search terms. When the query contains multiple patterns, the search can be carried out either as a Boolean AND function by requiring all parts of the query to be present within a given sequence (this is the **default**, with button set to No), or as a Boolean OR function to identify sequences that contain any part of the query (set button to Yes). If Yes, filtered sequences must have at least one occurrence of **at least one** of the listed patterns. If No (DEFAULT), filtered sequences must have at least one occurrence of **each** of the listed patterns.

The 3rd set of radio buttons (**Fig. 7D**) determines the type of pattern (DEFAULT = Nucleotide). If Nucleotide, then degenerate nucleotide codes are allowed, and T/U are interchangeable. Degenerate search patterns are **not** allowed for other sequence types. Importantly, all patterns are converted to uppercase and have white spaces removed regardless of the pattern type.

1. **A/T/G/C/U** - single bases
2. **R** - puRine (A/G)
3. **Y** - pYrimidine (C/T)
4. **W** - Weak (A/T)
5. **S** - Strong (G/C)
6. **M** - aMino (A/C)
7. **K** - Keto (G/T)
8. **B** - not A
9. **D** - not C
10. **H** - not G
11. **V** - not T/U
12. **N** - aNy base (not gap)

The **Start** button begins the search process. The *searched* data table will be shown on the right side of the screen. The **Download** button opens a file browser prior to downloading the output as a FASTA or CSV file (DEFAULT = FASTA, which is required for subsequent modules).

A sample output data table is shown in **Fig. 8** with the following parameters: **comma-separated patterns** = UCCG,CGGGAnAA, **parentheses** = No, **partial filtering** = No, and **pattern type** = Nucleotide.

Show 10 entries Search: TCCG|CGGGA[ACGT]AA

| id | Rank | Reads | RPM | seqs |
|---------------------|------|--------|-----------|---|
| All | All | All | All | All |
| >2-313312-145037.35 | 2 | 313312 | 145037.35 | CATAGCGACTGTCCACGA TCCG AAGCCTAACGGGACA AA AGGCAAGAGCGCGATA CCAATGCTGGACTG |
| >3-174096-80591.94 | 3 | 174096 | 80591.94 | AACCGCAAGCAACCCCAGCAAGAACAT TCCG ACGCACGA CGGGAGAA AGTG CATTACCACGATGTCGAT |
| >4-94978-43966.9 | 4 | 94978 | 43966.9 | CATAGCGACTGCCACGA TCCG AAGCCTAACGGGACA AA AGGCAAGAGCGCGATA CCAATGCTGGACTG |
| >6-57625-26675.57 | 6 | 57625 | 26675.57 | CCCTCTTGTATGACGCTAACTGAGAAT TCCG AAGCTAACGGGACA CGGGAGAA AGG ACACTTATGACGTGGCGG |
| >8-39793-18420.84 | 8 | 39793 | 18420.84 | AGCGCGCACCCAAATCGAAAT TCCG AAGGCAGGGACA CGGGAGAA ATGC GACCAAAGATA CCCTGTGAATGGC |
| >12-22089-10225.37 | 12 | 22089 | 10225.37 | CATAGCGACTGTCCACGA TCCG AAGCCTAACGGGACA AA AGGCAAGAGCGCGATA CCAATGCTGGACTG |
| >13-14115-6534.07 | 13 | 14115 | 6534.07 | CATAGCGACTATCCACGA TCCG AAGCCTAACGGGACA AA AGGCAAGAGCGCGATA CCAATGCTGGACTG |
| >14-11313-5236.98 | 14 | 11313 | 5236.98 | AGCGCGCACCCAAATCGAAAT TCCG AAGGCAGGGACA CGGGAGAA ATGC GTCAAAGATA CCCTGTGAATGGC |
| >15-10818-5007.83 | 15 | 10818 | 5007.83 | CATAGCGACTGTCCACGA TCCG AAGCCTAACGGGACA AA AGGCAAGAGTGCG GATA CCAATGCTGGACTG |
| >16-10514-4867.11 | 16 | 10514 | 4867.11 | CATAGCGACCGTCCACGA TCCG AAGCCTAACGGGACA AA AGGCAAGAGCGCGATA CCAATGCTGGACTG |

Showing 1 to 10 of 36,025 entries

Previous 1 2 3 4 5 ... 3603 Next

Figure 8: FASTAptameR-Motif_Search Output.

3.6 FASTAptameR-Motif_Tracker

3.6.1 Description

FASTAptameR-Motif_Tracker reports on the occurrence of one or more query patterns / sequences across multiple populations. The module accepts at least two *counted* FASTA files as input and returns a data table of metadata related to the enrichment of the query pattern(s) across multiple populations. Multiple FASTA files should be selected from the file browser at the same time. Columns of the output data table include the following:

1. Population
2. File name
3. Query
4. Rank
5. Reads
6. RPM

Optionally, an alias list can be provided and will be included as a separate column. These aliases will be used in the legend of the line plot. If provided, there must be one alias per query per line.

This output can be downloaded as a CSV file and will include appended enrichment scores. A screenshot of the module interface is shown in **Fig. 9**.

3.6.2 Usage

The input FASTA files must be chosen with the file browser. The following text box must contain at least one pattern or sequence. If the user wishes to search for multiple patterns or sequences, they must be included on separate lines. The set of radio buttons determines the type of query (**DEFAULT = Nucleotide**). If **Nucleotide**, then degenerate nucleotide codes are allowed. Note, the query is converted to uppercase and white spaces are removed regardless of the type.

The **Start** button begins the motif enrichment process. The resulting data table will be shown on the right side of the screen. The **Download** button opens a file browser prior to downloading the output as a CSV file.

Search Tracker

Input data:

FASTA files

Holding ctrl (Windows) or command (Mac) will allow you to click multiple files.

Select file order.

Motif or sequence list:

Alias list:

Search for motifs or whole sequences?

Motif Sequence

Type of pattern?

Nucleotide AminoAcid String

Figure 9: Screenshot of FASTAptameR-Motif_Tracker.

A screenshot of a sample output data table from tracked sequences is shown in **Fig. 10** with the three most abundant sequences from the 70HRT14 population.

| Show 10 entries | | | | | | Search: |
|-----------------|---------------------|---|-------|------|--------|-----------|
| Population | FileName | seqs | Alias | Rank | Reads | RPM |
| 1 | 70HRT14-count.fasta | AACCGCAAGCAACCCAGCAAGAAACATCGACGCACGACGGGAGAAGTCATTACACGATGTCGAT | 3rd | 3 | 174096 | 80591.94 |
| 2 | 70HRT15-count.fasta | AACCGCAAGCAACACCCAGCAAGAAACATCCGACGCACGACGGGAGAAGTCATTACACGATGTCGAT | 3rd | 5 | 104932 | 52786.23 |
| 1 | 70HRT14-count.fasta | ACGTTGTCGAAAGCCTATGCAAATTAGGACTGTCGACGAAACCTTGCCTAGACTGCCACGCTTGGTGT | 1st | 1 | 417696 | 193358.44 |
| 2 | 70HRT15-count.fasta | ACGTTGTCGAAAGCCTATGCAAATTAGGACTGTCGACGAAACCTTGCCTAGACTGCCACGCTTGGTGT | 1st | 3 | 161830 | 81408.87 |
| 1 | 70HRT14-count.fasta | CATAGCGACTGTCCACGAATCCGAAAGCTAACGGGACAAAGGCAAGAGCGCGATAACCATGCTGGACTG | 2nd | 2 | 313312 | 145037.35 |
| 2 | 70HRT15-count.fasta | CATAGCGACTGTCCACGAATCCGAAAGCTAACGGGACAAAGGCAAGAGCGCGATAACCATGCTGGACTG | 2nd | 1 | 382391 | 192362.47 |

Showing 1 to 6 of 6 entries

Previous 1 Next

Figure 10: FASTAptameR-Motif_Tracker Output.

3.6.3 Plotting

This module can generate an interactive line plot showing the query's RPM across each population (**Fig. 11**).

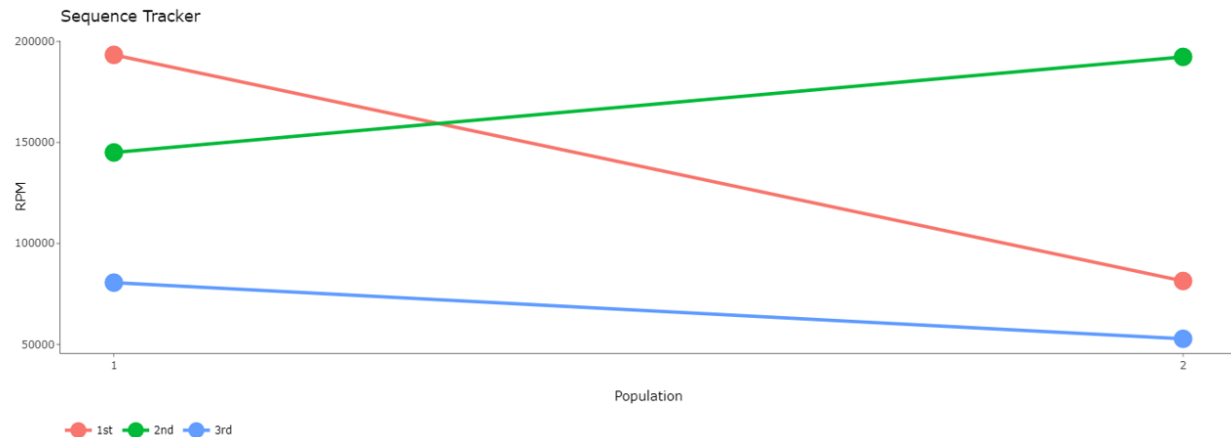


Figure 11: Sequence Tracking Line Plot. Shows the RPM of three sequences across the 70HRT14 and 70HRT15 populations. The aliases - 1st, 2nd, 3rd - refer to the 1st, 2nd, and 3rd most abundant sequences from the 70HRT14 population.

3.7 FASTAptameR-Distance

3.7.1 Description

FASTAptameR-Distance tabulates the distribution of distances from a user-defined reference sequence for all sequences in a population. The module accepts a *counted* FASTA file as input and returns a data table that contains a column for the Levenshtein edit distance (LED) between each input sequence and a query sequence. The output can be downloaded as a FASTA or CSV. A screenshot of the module interface is shown in Fig. 12.

The screenshot shows the user interface for the FASTAptameR-Distance module. At the top, there is a section labeled "Input data:" with two buttons: "Browse..." (black background with white text) and "FASTA or CSV file" (light gray background with black text). Below this is a section labeled "Query sequence:" containing a large, empty text input field. Underneath is a "Sequence range:" section featuring a horizontal slider with numerical markers at 1, 31, 61, 91, 121, 151, 181, 211, 241, 271, and 300. The slider has two circular endpoints, one at position 1 and another at position 300. Below the slider is a "FASTA or CSV download?" section with two radio buttons: "FASTA" (selected, indicated by a blue outline) and "CSV". At the bottom of the interface are two buttons: "Start" (black background with white text) and "Download" (white background with black text and a download icon). A horizontal line separates this from a final section labeled "Distance Histogram" in a dark gray box.

Figure 12: Screenshot of FASTAptameR-Distance.

3.7.2 Usage

The input FASTA file must be chosen with the file browser, and the following text box must contain a single query sequence. Note, this query sequence may not have any degenerate nucleotide codes. The **Start** button

begins the distance calculations. The resulting data table will be shown on the right side of the screen. The **Download** button opens a file browser prior to downloading the output as a FASTA or CSV file.

A sample output data table is shown in **Fig. 13** with the following query sequence (the most abundant sequence from the 70HRT14 dataset):

ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT.

| Show 10 entries | Search: | | | |
|---|---------|--------|-----------|----------|
| seqs | Rank | Reads | RPM | Distance |
| All | All | All | All | All |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT | 1 | 417696 | 193358.44 | 0 |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT | 5 | 74389 | 34435.91 | 1 |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT | 7 | 53608 | 24816.04 | 1 |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT | 20 | 8003 | 3704.72 | 1 |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT | 21 | 7815 | 3617.69 | 1 |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT | 23 | 6177 | 2859.44 | 1 |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT | 26 | 5487 | 2540.02 | 1 |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT | 28 | 5302 | 2454.38 | 1 |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAAACCTTGCCTAGACTCGCCACGCTTGGTGT | 29 | 5079 | 2351.15 | 1 |
| ACGTTGTCGAAAGCCTATGCAAATTAAAGGACTGTCGACGAGACCTTGCCTAGACTCGCCACGCTTGGTGT | 31 | 4504 | 2084.98 | 1 |

Showing 1 to 10 of 72,921 entries

Previous 1 2 3 4 5 ... 7293 Next

Figure 13: FASTAptameR-Distance Output.

The slider bar allows the user to select a range of positions to query. For example, setting the two ends of this slider bar to 10 and 60 will truncate all of the sequences (**including the query sequence**) to be in that specific range. Thus, the resulting distance value will be the LED between positions 10-60 of the query sequence and positions 10-60 of every other sequence in the data. Note, this option is best used if the sequences have already been length-filtered in FASTAptameR-Count since this module will not align any sequences.

3.7.3 Plotting

This module can also generate interactive histograms of distances (**Fig. 14**). The top plot corresponds to the distances between the query and unique sequences, whereas the bottom plot corresponds to the distances between the query and all sequences. In both cases, the query sequence is displayed at the top of the plot.

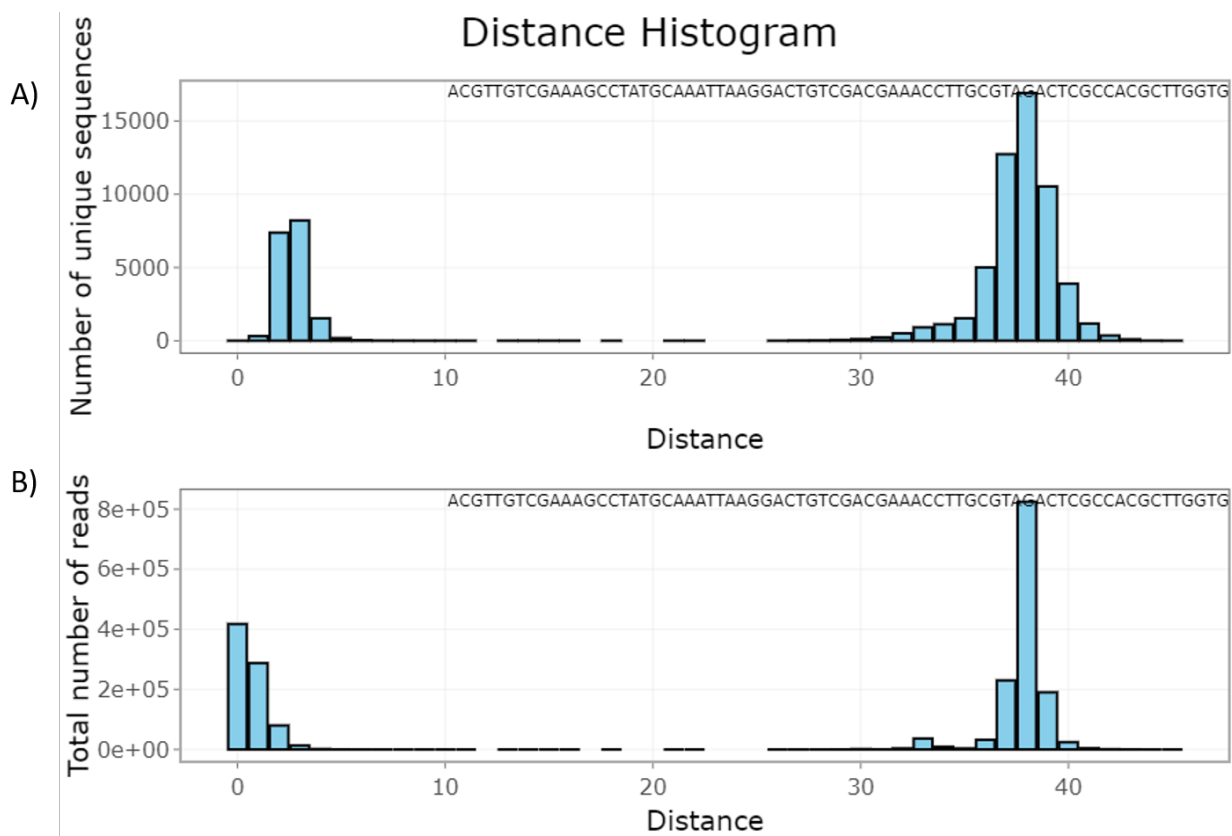


Figure 14: Distance Histograms with the 70HRT14 population as the target and the most abundant sequence as the query. A) Distance histogram with only unique sequences. B) Distance histogram with all reads.

3.8 FASTAptameR-Enrich

3.8.1 Description

FASTAptameR-Enrich calculates the enrichment (or depletion) of each sequence in one population relative to other populations. The module at least two *counted* FASTA files as input and returns a single data table after merging by sequences. Column headers for output data are appended with *.a*, *.b*, *.c*, etc., depending on the order in which they are uploaded. Additional columns include enrichment scores (“Enrichment” = $\text{RPM}_2 / \text{RPM}_1$) and the base-two logarithm of the Enrichment (“ $\log_2(E)$ ” = $\log_2(\text{Enrichment})$). For simplicity, comparisons are only made between consecutive populations (*e.g.*, 2:1, 3:2, etc.). This output can be downloaded as a CSV.

A screenshot of the module interface is shown in **Fig. 15**.

3.8.2 Usage

The input FASTA files must be chosen with the file browser. The following set of radio buttons determine whether missing values are allowed in the output. Missing values result from sequences that are only present in a subset of the input files.

The **Start** button begins the enrichment calculations, and the resulting data table will be shown on the right side of the screen. All numeric columns in this data table are filterable by typing into the corresponding text box (*e.g.*, 1 . . . 10 to keep values in the range [1:10]) or by using the slider bar that is displayed after clicking in the corresponding text box. Note, these filters apply the mask only to the displayed data, so calculations will **not** be repeated when the filters are altered. To display all data again, delete the filters from the text boxes. Note, many other outputs are similarly filterable.

The **Download** button opens a file browser prior to downloading the output as a CSV file. A sample filtered output data table is shown in **Fig. 16**.

3.8.3 Plotting

This module can generate five types of interactive plots (**Fig. 17, 18**): sequence persistence bar plots (**Fig. 17A**), $\log_2(\text{Enrichment})$ histograms (one per comparison - **Fig. 17B**), RPM scatter plots (one per comparison - **Fig. 17C**), RA plots (one per comparison - **Fig. 17D**), and a cluster box plot in the case where **clustered** FASTAs are provided (**Fig. 18**).

The sequence persistence bar plot (**Fig. 17A**) bins sequences by how many rounds they are found in all uploaded FASTA files. The slider bar just above this plot button filters these sequences by their respective read count. These results suggest that approximately 25000 sequences were found in both populations, whereas approximately 80000 sequences were found in only one population.

The spread of the $\log_2(\text{Enrichment})$ histogram (**Fig. 17B**) relative to a vertical line at $x = 0$ can indicate the magnitudes of enrichment (or depletion), while displacement of the centroid of the distribution from this line can indicate possible directionality of the population’s evolution. Similarly, the spread of the RPM scatter plot (**Fig. 17C**) relative to the diagonal line at $y = x$ can also indicate the magnitudes of enrichment and possible directionality. Finally, the RA plot (**Fig. 17D**) is used to show the relationship between the average log-RPM and $\log_2(\text{Enrichment})$ for each sequence. Note that missing sequences (those with **RPM** = 0 in one round) are treated as having **RPM** = 0.1 for the sake of getting their log2.

The cluster box plot (**Fig. 18**) shows the distribution of enrichment values for sequences after grouping by cluster. The red marker indicates where the seed sequence of the cluster falls.

Sequence Enrichment [Positional Enrichment](#)

Input data:

[Browse...](#) FASTA files

Holding ctrl (Windows) or command (Mac) will allow you to click multiple files.

Keep missing sequences?

Yes No

[Start](#) [!\[\]\(952835fe0c32eed2cddf0a857395a954_img.jpg\) Download](#)

Minimum number of reads to consider for persistence plot?

0 1,000

[Seq. Persistence](#)

[log2\(Enrichment\) Histogram](#) [RPM Scatter Plot](#) [MA Plot](#)

[Cluster Boxplot](#)

Figure 15: Screenshot of FASTAptameR-Enrich.

| Show 10 entries | | Search: <input type="text"/> | | | | | | | |
|---|-----|------------------------------|-----------|-------|--------|-----------|-------|---------------|----------|
| seqs | | Rank.a | Reads.a | RPM.a | Rank.b | Reads.b | RPM.b | enrichment.ba | log2E_ba |
| All | All | All | All | All | All | All | All | All | All |
| ACGTTGCGAAAGCTATGCAAATTAGGACTGCGACGAAACCTTGCGTAGACTGCCACGCTGGTGT | 1 | 417696 | 193358.44 | 3 | 161830 | 81408.87 | 0.421 | -1.248 | |
| CATAGCGACTGTCCACGATCGAAGCTAACGGACAAAAGGCAAGGCGCGATAACATGCTGGACTG | 2 | 313312 | 145037.35 | 1 | 382391 | 192362.47 | 1.326 | 0.407 | |
| AACCGCAGAACACCCAGCAAGAAACATCCGACGCGACGGGAGAAAGTCGATTACACGATGTCGAT | 3 | 174096 | 80591.94 | 5 | 104932 | 52786.23 | 0.655 | -0.61 | |
| CATAGCGACTGCCACGATCCGAGCTAACGGACAAAAGGCAAGGCGCGATAACATGCTGGACTG | 4 | 94978 | 43966.9 | 6 | 42954 | 21608.09 | 0.491 | -1.025 | |
| ACGTTGCGAAAGCTATGCAAATTAGGACTGCGACGAAACCTTGCGTAGACTGCCACGCTGGTGT | 5 | 74389 | 34435.91 | 9 | 32821 | 16510.66 | 0.479 | -1.061 | |
| CCCTCTTGATGCGCTAACTGAGAACATCGAACGGAGAAAGGACACTTACGCTGGCG | 6 | 57625 | 26675.57 | 7 | 37701 | 18965.55 | 0.711 | -0.492 | |
| ACGTTGCGAAAGCTATGCAAATTAGGACTGCGACGAAACCTTGCGTAGACTGCCACGCTGGTGT | 7 | 53608 | 24816.04 | 10 | 30749 | 15468.34 | 0.623 | -0.682 | |
| AGCGCGCACCCAAATCGAACATCGAACGGGAGAACACAATGGATTACCGAGCTGGCTGAC | 8 | 39793 | 18420.84 | 15 | 15414 | 7754.04 | 0.421 | -1.248 | |
| TTGACAATACTCGAGAGAACCGAGGTGAAACACGGGACAAACAGGCATCAATGGAGTGGTAC | 9 | 33800 | 15646.58 | 4 | 136505 | 68669.08 | 4.389 | 2.134 | |
| GCGAACCAACCCAGATTACTAACCGTGGCCTGAAACACGGGACAAACAGGCATCAATGGAGTGGTAC | 10 | 29794 | 13792.14 | 176 | 732 | 368.23 | 0.027 | -5.227 | |

Showing 1 to 10 of 21,924 entries

Previous 1 2 3 4 5 ... 2193 Next

Figure 16: Filtered FASTAptameR-Enrich Output.

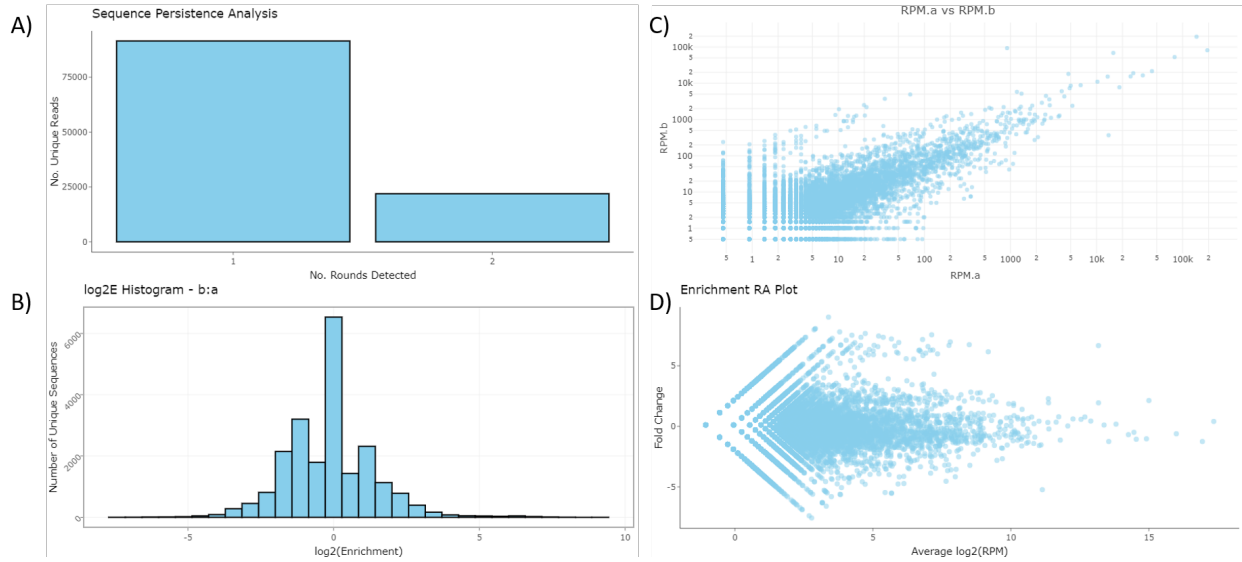


Figure 17: FASTAptameR-Enrich Plots. A) The sequence persistence bar plot shows across how many rounds unique sequences persist. B) The histogram shows the distribution of fold-change between rounds. C) The RPM scatter plot shows the RPM of sequences across two rounds. D) The RA plot has fold change on the y-axis (R for Ratio) and average log2(RPM) on the x-axis (A for average).

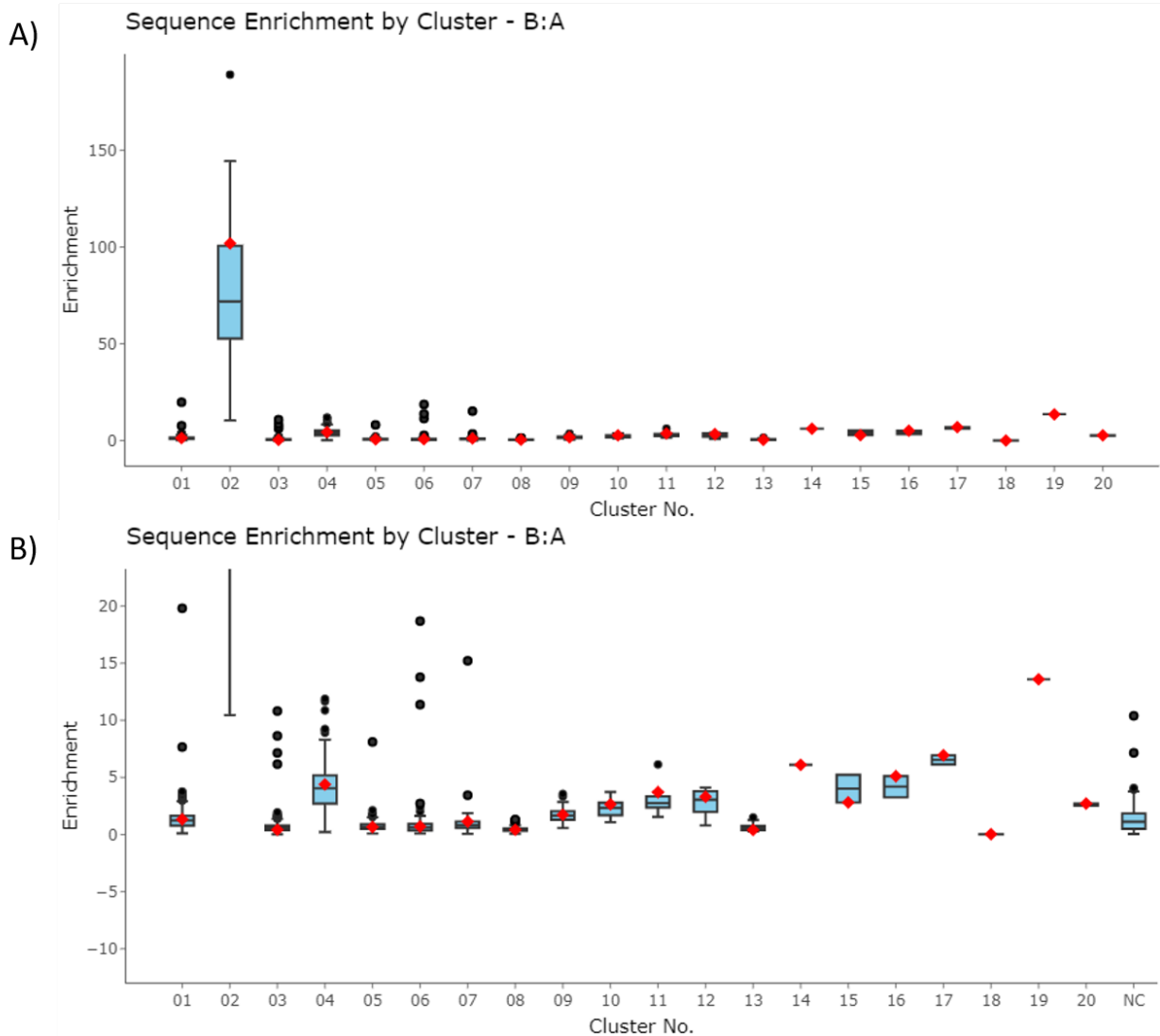


Figure 18: Cluster Box Plots of Sequence Enrichment. A) Enrichment distribution for the top 20 clusters. B) The same plot after zooming in on the y-axis.

3.9 FASTAptameR-Position_Enrich

3.9.1 Description

FASTAptameR-Position_Enrich calculates the average enrichment (or depletion) for each position that does not match the corresponding residue in the user-defined reference sequence. For example, if the first residue of the reference sequence is E, then this module will calculate the average enrichment of all sequences that do not have an E in the first position. Thus, it is recommended that all sequences are of the same length (can be done by applying a filter to the output table from FASTAptameR-Count).

This module accepts a CSV from the previous module, though it exclusively operates on the `enrichment_ba` column. Thus, an enrichment CSV with >2 populations can be uploaded here, but only the first enrichment column will be used.

The outputs of this module are two plots. The first is a bar plot showing the average enrichment value for each position. The second is a heat map that shows the average enrichment per position per residue.

A screenshot of the module interface is shown in [Fig. 19](#).

3.9.2 Usage

The input CSV file must be chosen with the file browser, and the reference sequence must be added in the subsequent text box. The first set of radio buttons allows the standard genetic code to be altered. Each change should occupy a single line. To add a residue, enter its single-letter code. To replace a residue, enter a comma-separated pair (*e.g.*, A,B will replace A with B in the algorithm and resulting plots).

The slider bar allows the user to set the minimum and maximum enrichment values (*e.g.*, 0-10 means that any value greater than 10 is made equal to 10). The final set of radio buttons determines the set of residues for which the algorithm searches. The next three text boxes allow the user to set the “low”, “middle”, and “high” colors for the plots.

Finally, the `Start` button generates the two plots.

3.9.3 Plotting

FASTAptameR-Position_Enrich generates two types of plots ([Fig. 20](#)): 1) average enrichment bar plot and 2) average enrichment heat map. The bar plot ([Fig. 20A](#)) shows the reference sequence on the x-axis and the average enrichment on the y-axis. The heat map ([Fig. 20B](#)) shows the reference sequence on the x-axis, possible residues on the y-axis, and average enrichment in the color axis. The workflow to generate these plots is given below:

1. Count 70HRT14.fastq and 70HRT15.fastq with FASTAptameR-Count. Filter for sequences with exactly 70 nucleotides.
2. Generate top five clusters for each counted population with FASTAptameR-Cluster. Include all sequences (min. reads > 0) and use LED = 7.
3. Use cluster 1 from 70HRT14 and cluster 2 from 70HRT15 in FASTAptameR-Enrich. The former population corresponds to 6/5 asymmetric loop aptamers, whereas the latter population corresponds to family 1 pseudoknots.
4. Use the output CSV in FASTAptameR-Position_Enrich.

Positions ~1-7 and ~40-44 have poor enrichment scores, which suggests the importance of these subsequences. Positions 13 and 55, however, have relatively high average enrichment scores, suggesting that these mutations (G->T for position 13 and C->G for position 55) may be selected for in the experiment.

Sequence Enrichment Positional Enrichment

Choose data to analyze:

Sequence Enrichment CSV file

Reference sequence:

Do you want to adjust the standard alphabet?

Yes No

Range of enrichment values:

0 10 200

Type of sequences?

Nucleotide AminoAcid

Select low colour

red2

Select middle colour

gold

Select high colour

yellow

Start

Figure 19: Screenshot of FASTAptameR-Position_Enrich.

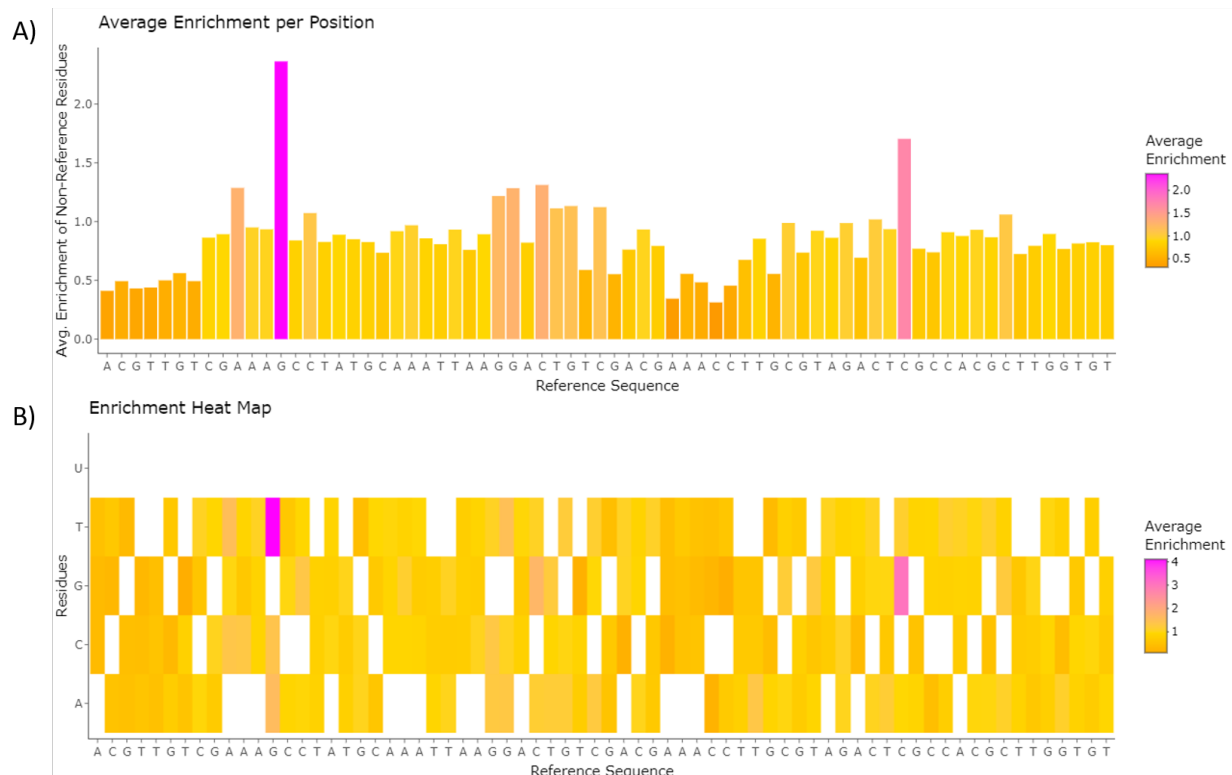


Figure 20: FASTAptameR-Position_Enrich Plots. A) The bar plot shows the user-defined reference sequence on the x-axis and average enrichment of non-reference residues on the y-axis. B) The heat map shows the user-defined reference sequence on the x-axis and all possible residues on the y-axis (nucleotides for this use case). Colors depict average enrichment of non-reference residues.

3.10 FASTAptameR-Cluster

3.10.1 Description

FASTAptameR-Cluster groups sequences according to sequence relatedness for all sequences in the population within a user-defined threshold of similarity. The module accepts a *counted* FASTA file as input. If no output directory is specified (the default setting), the module returns a *clustered* data table to the screen. This data table contains all sequences and clusters and can be downloaded as a single FASTA or CSV file. If an output directory is specified, then no data table will be created, and one FASTA file per cluster will be written to the output directory.

Briefly, the module identifies clusters in an iterative manner. During each iteration, the most abundant sequence that has not yet been clustered becomes a cluster “seed” for that iteration. Any other sequences that have not yet been clustered and that are within a user-defined edit distance of this seed sequence are added to this cluster. This process repeats until all sequences are clustered or a predefined number of clusters is created.

A screenshot of the module interface is shown in **Fig. 21**.

3.10.2 Usage

The input FASTA file must be chosen with the file browser. The 1st slider bar (**Fig. 21A**) sets the minimum number of reads a sequence must have to be clustered (**DEFAULT = 10**). Sequences with the chosen number or fewer reads are removed prior to clustering. The 2nd slider bar (**Fig. 21B**) sets the maximum Levenshtein edit distance to consider between a seed sequence and all other sequences (**DEFAULT = 7**). The 3rd slider bar (**Fig. 21C**) sets the total number of desired clusters (**DEFAULT = 20**). Note, any remaining sequences will be grouped as NC (“not clustered”).

The 1st set of radio buttons (**Fig. 21D**) indicates whether non-clustered sequences should be kept (**DEFAULT = No**). If **Yes** then the sequence IDs of non-clustered sequences will be appended with NC.

The 2nd set of radio buttons (**Fig. 21E**) indicate whether each cluster should be written to a different FASTA file (**DEFAULT = No**). If **No**, then all clusters are grouped together and downloaded in a single file. If **Yes**, then each cluster will be written to its own FASTA file, and no data table will be displayed. Note, a directory path **must be copied or typed** into the corresponding text box (**Fig. 21F**) if this option is **Yes**. A sample directory path could be **C:/Users/Kramer/Desktop/Data/directory/**, though this will depend on your system. **Please note** that this requires backslashes (/), and forward slashes (\) will cause errors. Also note that the path should end with a backslash.

The **Start** button will begin the clustering process. The results will be displayed as a data table on the right side of the screen. The **Download** button opens a file browser prior to downloading the output as a FASTA or CSV file (**DEFAULT = FASTA**, which is required for subsequent modules).

Algorithm progress will be shown below these buttons and will update after each cluster finishes. These notifications occur regardless of whether the module is writing to one or many files.

A sample output data table is shown in **Fig. 22**.

Note that the new *id* column is the old *id* with three new values appended to the end: **Cluster Number**, **Rank in Cluster**, and **Distance to Cluster Seed**.

Cluster Diversity Enrichment

A) Input data:
 FASTA file

Min. number of reads to cluster:
 1,000

Max. LED:
 20

Total clusters:
 1,000

Keep non-clustered sequences?
 Yes No

One file per cluster?
 Yes No

If Yes, please provide an absolute path to a directory below. No output will be displayed if Yes.

B) Directory path:

FASTA or CSV download?
 FASTA CSV

C)

Figure 21: Screenshot of FASTAptameR-Cluster.

| Show 10 entries | | | | | | | | Search: |
|---------------------------|-------|--------|-----------|---------|---------------|-------|--|---------|
| id | Rank | Reads | RPM | cluster | rankInCluster | LED | seqs | |
| [All] | [All] | [All] | [All] | [All] | [All] | [All] | [All] | |
| >1-417696-193358.44-1-1-0 | 1 | 417696 | 193358.44 | 1 | 1 | 0 | ACGTTGTCGAAGCTATGCAAATTAAAGGACTGTGACGAAACCTTGCCTAGACTGCCACGCTGGTGT | |
| >2-313312-145037.35-2-1-0 | 2 | 313312 | 145037.35 | 2 | 1 | 0 | CATAGCGACTGTCACGAATCCGAAGCCTAACGGACAAAAGGCAAGAGCGCGATACCAATGCTGGACTG | |
| >3-174096-80591.94-3-1-0 | 3 | 174096 | 80591.94 | 3 | 1 | 0 | AACCGCAAGCAACACCCAGCAAGAACATCCGACGCCACGACGGAGAAAGTCGATTACACGATGTCGAT | |
| >4-94978-43966.9-2-2-1 | 4 | 94978 | 43966.9 | 2 | 2 | 1 | CATAGCGACTGCCACGAATCCGAAGCCTAACGGACAAAAGGCAAGAGCGCGATACCAATGCTGGACTG | |
| >5-74389-34435.91-1-2-1 | 5 | 74389 | 34435.91 | 1 | 2 | 1 | ACGTTGTCGAAGCTATGCAAATTAAAGGACTGTGACGAAACCTTGCCTAGACTGCCACGCTGGTGT | |

Figure 22: FASTAptameR-Cluster Output.

3.11 FASTAptameR-Cluster_Diversity

3.11.1 Description

FASTAptameR-Cluster_Diversity evaluates diversity across the *clustered* population and sequence relationships within and between clusters. The module accepts a *clustered* FASTA file as input and returns a data table with metadata for each cluster. This data table can be downloaded as a CSV file.

A screenshot of the module interface is shown in Fig. 23.

3.11.2 Usage

The input FASTA file must be chosen with the file browser. The Start button begins the analysis. The results will be displayed as a data table on the right side of the screen and include the following columns: Cluster Number, Seed Sequence, Total Sequences, Total Reads, and Total RPM. The Download button opens a file browser prior to downloading the output as a CSV file, which can be used by FASTAptameR-Cluster_Enrich. A sample output data table is shown in Fig. 24.

3.11.3 Plotting

This module can generate metaplots of the analyzed data. These line plots correspond to the number of unique sequences per cluster, total reads per cluster, and average LED to seed sequence per cluster (Fig. 25A).

This module is also able to analyze clusters by converting all sequences into k-mer vectors and rendering an interactive 2D PCA plot, colored by cluster (Fig. 25B). The value of k can be chosen with the 1st set of radio buttons (DEFAULT = 3). The slider bar indicates how many of the top clusters should be plotted (max = 21 clusters due to graphics limitations). The 2nd set of radio buttons indicates whether non-clustered (NC) sequences should be plotted (DEFAULT = Yes). Note that non-clustered (NC) sequences in the output are marked as NA in this plot.

Importantly, only nucleotide sequences without ambiguities should be plotted in this module. The large k-mer matrix needed for peptide sequences may return errors related to memory usage. Further, this module will reject any set of sequences with characters outside of [A, C, G, T/U]. The resulting k-mer PCA plot will not display anything.

Cluster Diversity Enrichment

Input data:

Clustered FASTA file

kmer size for PCA plot:

3 4 5

Number of top clusters to plot:

1 20
1 3 5 7 9 11 13 15 17 19 20

Keep non-clustered sequences?

Yes No

*Characters outside of [A,C,G,T,U] converted to 'X'.

Figure 23: Screenshot of FASTAptameR-Cluster_Diversity.

| Cluster | | Seeds | TotalSequences | TotalReads | TotalRPM | AverageLED |
|---------|--|-------|----------------|------------|----------|------------|
| All | All | All | All | All | All | All |
| 1 | ACGTTGTGAAAGCCTATGCAAATTAAAGGACTGTGACGAAACCTTCCGTAGACTGCCACGGTTGGT | 1259 | 770383 | 356623.54 | 1.86 | |
| 2 | CATAGCGACTGTCCACGAATCGAACGCCAACGGGACAAAGGCAAGGCACGCGATACCATGCTGGACTG | 1295 | 652468 | 302038.75 | 1.97 | |
| 3 | AACCGCAAGAACACCCGACGAAAGAACATCCGACGACGACGGGAGAAAGTCATTACACGATGTCGAT | 528 | 257675 | 119282.23 | 1.63 | |
| 4 | CCCTCTTGATGACGCTAACTGAGAAATCGAACGGGAGAAAGGACACTTATGACGTGGCG | 324 | 101448 | 46962.14 | 1.49 | |
| 5 | AGCGCGCACCCAAATCGAAATCCGAAAGGCGAACGGGAGATGCGACCAAAGATAACCTGTGAATGCC | 262 | 73809 | 34167.47 | 1.47 | |
| 6 | TTGACAATAACTCGAGAAGAACCGAGGTGCAAACCGGGAGAACACAATGGATTACCCGAGCTCGGCTGAC | 275 | 67908 | 31435.87 | 1.51 | |
| 7 | GCGAACCAAAACCCAGATTACTAACCGTGGCTGAAACACGGGACAAAACAGGCATCAATGGAGTGTAC | 148 | 41566 | 19241.67 | 1.16 | |
| 8 | ACGTTGTGACGGATGCCACGGTGCACGAAACCTTGTGGATAGCGGAAATCGACGAGTGTGCC | 163 | 44693 | 20689.16 | 1.26 | |
| 9 | ACCAAATCCCACACTACAATCGAACGCTAACGGGACATTGCGAAATGGAACATACGGGCTGTTGA | 67 | 9114 | 4219.07 | 1.1 | |
| 10 | GTGCCCTACCACATGATCGAGGGAAAAGGGAAAGATGATCGATTACCGAACCGGCCACGGACA | 54 | 7292 | 3375.58 | 0.98 | |

Showing 1 to 10 of 21 entries

Previous 1 2 3 Next

Figure 24: FASTAptameR-Cluster_Diversity Output.

3.12 FASTAptameR-Cluster_Enrich

3.12.1 Description

FASTAptameR-Cluster_Enrich calculates the enrichment (or depletion) of each cluster in one population relative to other populations. The module accepts two or three *cluster-analysis* CSV files as input and returns a data table after merging by **Seed Sequence**. Thus, this module assumes that cluster seeds are consistent across populations, though this may not always be a valid assumption.

The output data table includes a column for **Enrichment** that can be downloaded as a CSV file. A screenshot of the module interface is shown in **Fig. 26**.

3.12.2 Usage

The input FASTA files must be chosen with the file browser. The **Start** button begins the enrichment calculation. The results will be displayed as a data table on the right side of the screen and include a column for **Enrichment**. The **Download** button opens a file browser prior to downloading the output as a CSV file.

A sample output data table is shown in **Fig. 27**.

Note that columns 3-5 and 7-9 refer to *total* values in the given cluster.

3.12.3 Plotting

After merging by seed sequence, this module will generate a line plot in which the x-axis corresponds to population, and the y-axis corresponds to the total RPM of the seed's cluster (**Fig. 28**).

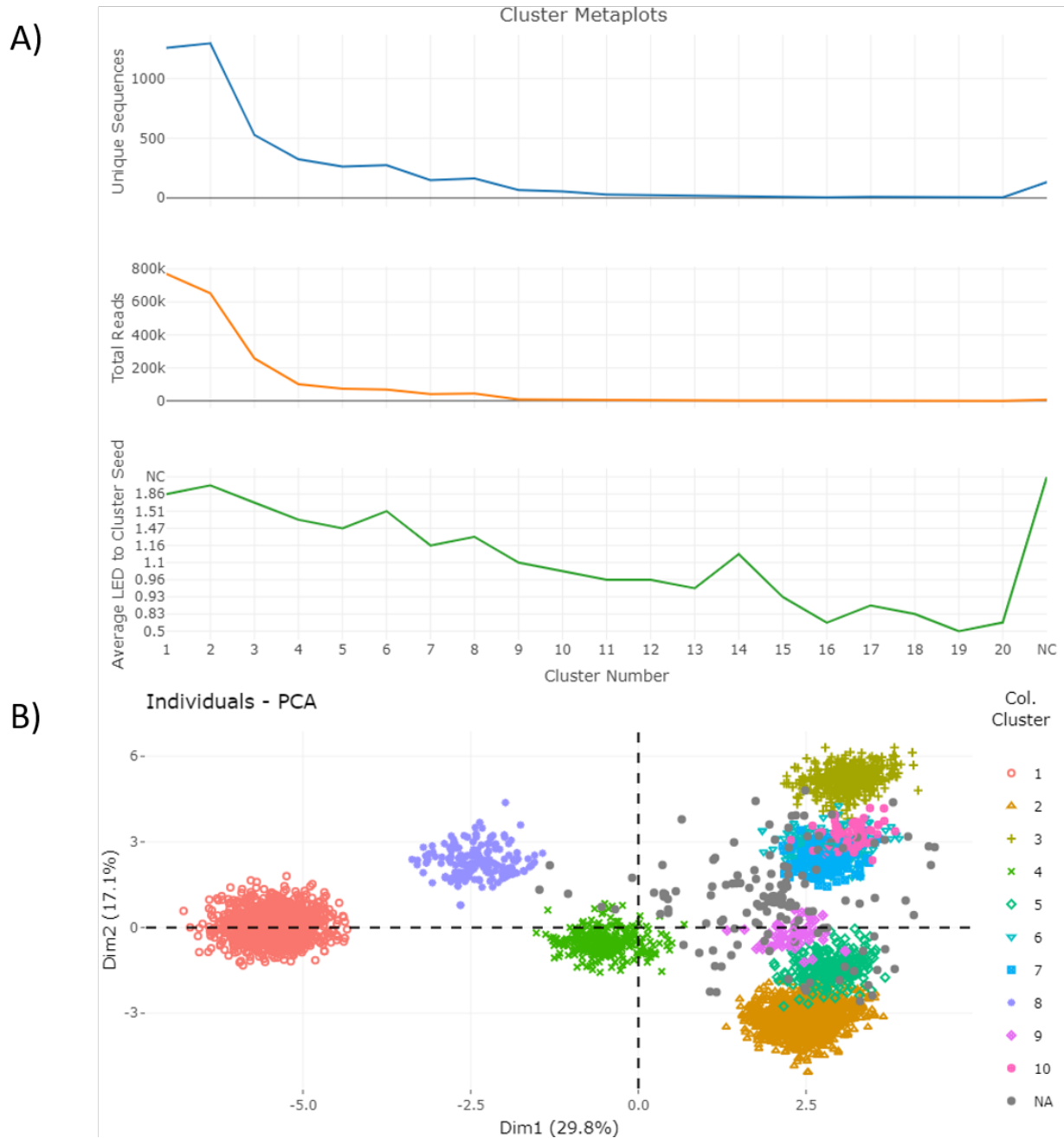


Figure 25: Cluster metaplots. A) Cluster metaplots depict number of unique sequences, total number of reads, and average LED to cluster seed per cluster. B) The k-mer PCA plot can qualitatively suggest how well the cluster algorithm performed.

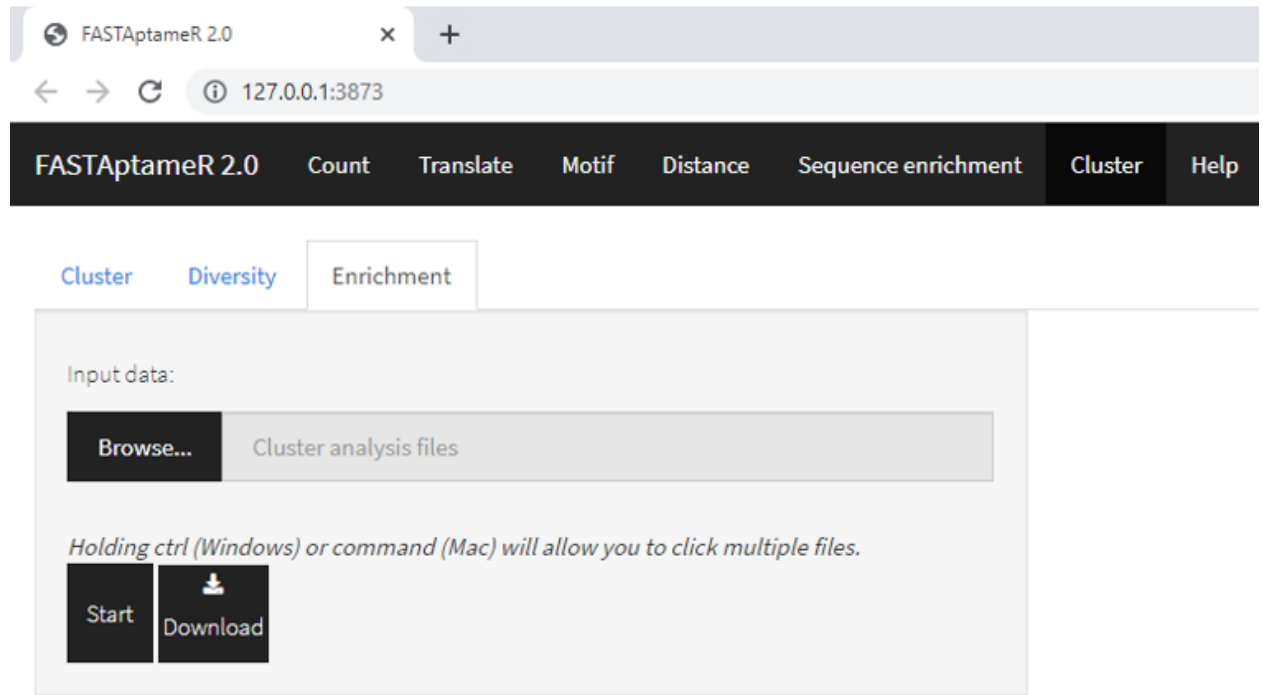


Figure 26: Screenshot of FASTAptameR-Cluster_Enrich.

| Cluster | | | | | | | | Seeds | | TotalSequences | | | | | | TotalReads | | TotalRPM | | AverageLED | | Population | | FileName | | | |
|---------|--|-----|-----|-----|-----|------|-----|--------|-----|----------------|------|-----|-----|-----|-----|------------|-----|----------|-----|------------|-----|------------|-----|----------|-----|--|-----|
| | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All | All |
| 1 | ACGTTGTCAAAGCCTATGCAAATTAAAGGACTGTGACGAAACCTTGCCTAGACTGCCACGCTTGGTGT | | | | | 1259 | | 770383 | | 356623.54 | 1.86 | | | | | | | | | | | | | | 1 | 70HRT14-count-cluster-clusterDiversity.csv | |
| 2 | CATAGCGACTGTCCACGAATCCGAAGCTAACGGACAAAAGGCAAGAGCGCGATAACATGCTGGACTG | | | | | 1295 | | 652468 | | 302038.75 | 1.97 | | | | | | | | | | | | | | 1 | 70HRT14-count-cluster-clusterDiversity.csv | |
| 3 | AACCGCAAGCACCCAGCAAGAACATCCGACGACGACGGGAGAAAGTCATTACACGATGTCGAT | | | | | 528 | | 257675 | | 119282.23 | 1.63 | | | | | | | | | | | | | | 1 | 70HRT14-count-cluster-clusterDiversity.csv | |
| 4 | CCCTCCTTGATGACGCTAACTGAGAACATCCGAAGTCCAACGGGAGAAAGGACACTTATGACGTGGCGCG | | | | | 324 | | 101448 | | 46962.14 | 1.49 | | | | | | | | | | | | | | 1 | 70HRT14-count-cluster-clusterDiversity.csv | |
| 5 | AGCGCGGCCACCCAAATCGAACGGCGAACGGGAGAATGCGACCAAAGATAACCTGTGAATGGC | | | | | 262 | | 73809 | | 34167.47 | 1.47 | | | | | | | | | | | | | | 1 | 70HRT14-count-cluster-clusterDiversity.csv | |

Figure 27: FASTAptameR-Cluster_Enrich Output.

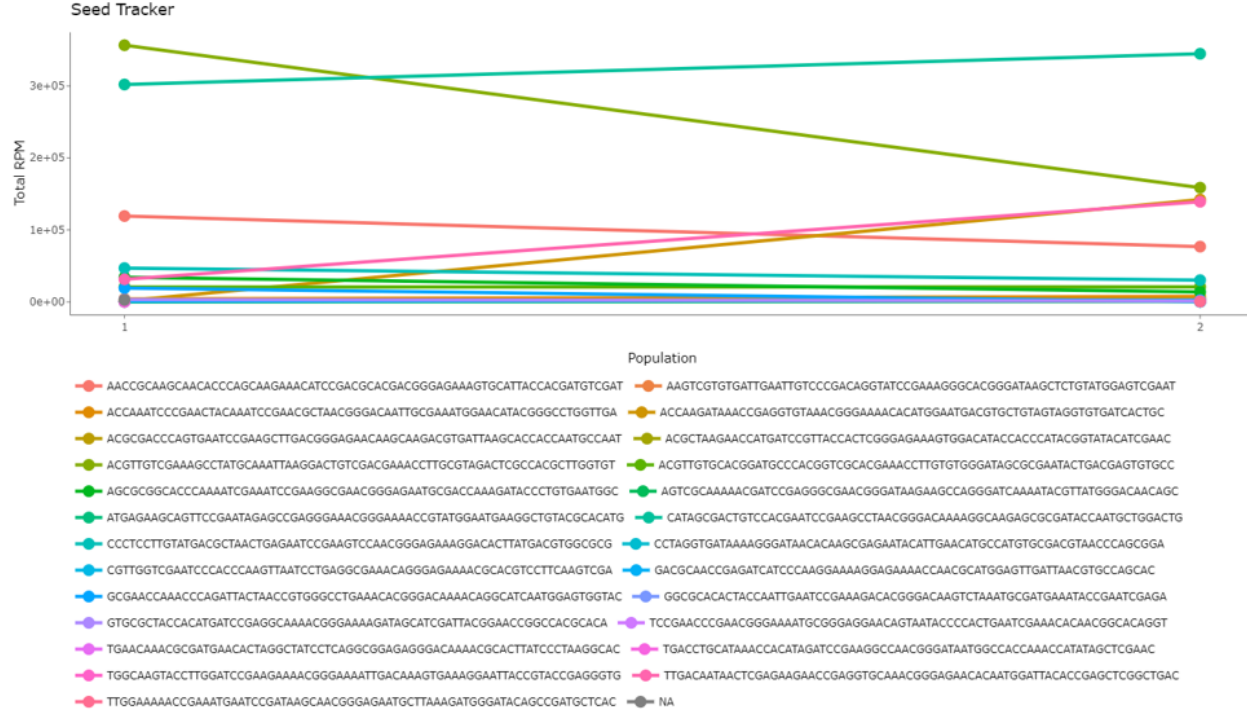


Figure 28: Cluster Tracker Line Plot.

4 Version history

References

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