AMOEBAE documentation

Lael D. Barlow

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5 References 30

1 Introduction

$_{\scriptscriptstyle 2}$ 1.1 What is AMOEBAE?

- ³ Analysis of MOlecular Evolution with BAtch Entry (AMOEBAE) is a bioinformatics software
- 4 toolkit composed primarily of scripts written in the Python3 language. AMOEBAE scripts
- 5 use existing Python packages including Biopython (Cock et al., 2009), the Environment for
- ⁶ Tree Exploration (ETE3) (Huerta-Cepas et al., 2016), pandas, and Matplotlib (Hunter, 2007)
- ⁷ for setting up, running, and summarizing analyses of molecular evolution using bioinformat-
- s ics software packages including MUSCLE (Edgar, 2004), BLAST+ (Camacho et al., 2009),
- 9 HMMer3 (Eddy, 1998), and IQ-Tree (Nguyen et al., 2015). Applications include identifying
- $_{10}$ and classifying predicted peptide sequences according to their evolutionary relationships with
- 11 homologues. All dependencies are freely available, and AMOEBAE code is open-source (see
- subsection 1.9) and available on GitHub (https://github.com/laelbarlow/amoebae).

13 1.2 Why use AMOEBAE?

Webservices such as those provided by NCBI (https://blast.ncbi.nlm.nih.gov/Blast. cgi) (Camacho et al., 2009) provide a means to investigate the evolution of one or a few genes 15 via similarity searching, and automated pipelines such as orthoMCL (Li, 2003) attempt to 16 rapidly perform orthology prediction for all genes in several genomes. AMOEBAE addresses 17 mid-scale analyses which are too cumbersome to be done via webservices and yet require a 18 level of detail and flexibility not offered by automated pipelines. AMOEBAE may be useful for analyzing the distribution of orthologues of up to perhaps 30 genes/proteins among a 20 sampling of no more than approximately 100 eukaryotic genomes. However, you may need to carefully define the scope of your analysis depending on what additional steps you may 22 find necessary beyond those that may be performed using AMOEBAE (30 queries and 100 23 genomes may in fact be unmanageable). AMOEBAE provides many options which can be 24 tailored to the specific genes/proteins being analyzed, and allow analyses using complex sets of customized criteria to be reproduced more practically. 26

$_{27}$ 1.3 Key features

- The core functionality is to run sequence similarity searches with multiple algorithms, multiple queries, and multiple databases simultaneously and to allow highly customizable imple-
- mentation of reciprocal-best-hit search strategies. The output includes detailed summaries
- of results in the form of a spreadsheet and plots.
- $_{32}$ A particular advantage of AMOEBAE over other tools is the functionality for parsing re-
- 33 sults of TBLASTN (searching in nucleotide sequences with peptide sequence queries) search
- results. This allows rapid identification of High-scoring Segment Pair (HSP) clusters at sepa-
- rate gene loci (identified according to user-defined criteria), automatic checking of those loci

- against information in genome annotation files, and systematic use of Exonerate (Slater and
- ² Birney, 2005) where possible for obtaining better exon predictions.

3 1.4 A word of caution

- 4 AMOEBAE is not optimized for ease of use, but is meant to be highly configurable. The
- 5 many options available to AMOEBAE users inevitably provide many opportunities for errors
- 6 in specifying search criteria, and errors in interpreting output files. Some prior experience
- 7 with similarity searching and with running software using the command line is essential
- 8 for using AMOEBAE, and experience writing scripts in Bash and Python would be highly
- 9 advantageous. Moreover, AMOEBAE is still under active development, so some features may
- not yet be thoroughly tested.

11 1.5 User support

- For specific issues with the code, please use the issue tracker on the GitHub webpage here:
- https://github.com/laelbarlow/amoebae/issues.
- 14 If you have general questions regarding AMOEBAE, please email the author at lael (at)
- ualberta.ca.

1.6 Documentation

- 17 This document provides an overview of AMOEBAE and describes the functionality of the var-
- ious commands/scripts. For a tutorial which includes a working example of a similarity search
- analysis run using AMOEBAE, see the Jupyter Notebook: amoebae/notebooks/similar-
- 20 ity search tutorial 2.ipynb. For code documentation, please see the html file(s), which can
- be opened with your web browser: amoebae/doc/code_documentation/html/index.html.

22 1.7 How to cite AMOEBAE

- 23 Please cite the GitHub webpage https://github.com/laelbarlow/amoebae (or alternative
- 24 permanent repositories if relevant). Also, the first publication to make use of a version of
- ²⁵ AMOEBAE was an analysis of Adaptor Protein subunits in embryophytes by Larson *et al.*
- 26 (2019).
- Also, you may wish to cite the software packages which are key dependencies of AMOEBAE,
- since AMOEBAE would not work without these (see subsection 2.2).

1 1.8 Acknowledgments

- ² AMOEBAE was initially developed at the Dacks Laboratory at the University of Alberta, and
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12 1.9 License

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- 20 permissions and limitations under the License.

$_{\scriptscriptstyle 21}$ 2 How to start using AMOEBAE

22 2.1 System requirements

- ²³ Please note that the commands shown likely only work on macOS or Linux operating systems
- 24 (you may have trouble running AMOEBAE directly on Windows).

$_{\scriptscriptstyle{25}}$ 2.2 Dependencies

- All dependencies are free and open-source, and can be automatically installed in a virtual
- environment (see subsection 2.3).

- ¹ These are the main dependencies of AMOEBAE:
- Python3 (the Anaconda distribution works well).
- Biopython, a Python package for bioinformatics (Cock et al., 2009).
- The Environment for Tree Exploration 3 (ETE3), a Python package for working with phylogenetic trees (Huerta-Cepas *et al.*, 2016).
- Matplotlib, a Python package for generating plots (Hunter, 2007).
- (gffutils).

12

- NCBI BLAST+, a software package for sequence similarity searching (Camacho *et al.*, 2009).
- HMMer3, a software package for profile sequence similarity searching (Eddy, 1998).
- MUSCLE, for multiple sequence alignment (Edgar, 2004).
 - IQ-TREE, for phylogenetic analysis (Nguyen et al., 2015).

³ 2.3 Setting up an environment for AMOEBAE using Singularity

- Follow the steps below to set up AMOEBAE on your personal computer. This setup process should take approximately 20 minutes to complete. Additional instructions for setting up AMOEBAE on a remote server will soon be added as well.
- 1. Ensure that Git is installed on your computer If you do not already have git installed, 17 then your computer will prompt you with instructions for how to install it when you 18 type git into the command line. If you have a newer version of macOS it may prompt 19 you to install developer tools, which may take up a considerable amount of storage 20 space. Documentation for Git is available here: https://git-scm.com/doc. You can 21 check which version you have (or whether it is installed at all) by running the command 22 below. Please note: Here ">>>" is used to indicate that the following text in the line 23 is to be entered in you terminal command prompt. 24
- 26 2. Clone the AMOEBAE repository using Git. If you simply download the code from
 Code GitHub, instead of cloning the repository, then AMOEBAE cannot record specifically
 what version of the code you use, and will not run properly. Make sure to use the
 appropriate directory path (the path shown is just an example). Also, replace the path
 shown below with the path to the directory on your system where you wish to put the
 main AMOEBAE directory.
- 32 >>> cd /path/to/directory/where/you/keep/files

- 3. Set up AMOEBAE. This performs several steps including checking for whether singularity is installed and attempting to use VirtualBox and Vagrant to run Singularity in a pre-built Ubuntu virtual machine with Singularity installed. This is because Singularity does not run on MacOS (or Windows), and installation of Singularity on Linux is complex, as several dependencies are required. This script downloads a pre-built singularity container, which was built using the singularity recipe file, and provided on the Singularity Library (https://cloud.sylabs.io/library/_container/5e8ca8fff0f8eb90a8a7b60d).
- 9 >>> cd amoebae
 10 >>> bash setup.sh

22

23

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4. If you are setting up AMOEBAE on a high performance computing cluster, then you will not be able to install Singularity yourself, and may need to use specific procedures to load Singularity prior to use.

14 2.4 Running AMOEBAE using Jupyter notebooks

- 1. After setting up AMOEBAE according to the instructions above, the easiest way to start running analyses using AMOEBAE is via the tutorials, which are in the form of Jupyter notebooks (https://jupyter.org/). These Jupyter notebooks can be run using the installation of Jupyter in the Singularity container, and can be accessed using your browser (on a personal computer). To start a Jupyter server, run the bash script as indicated below (assuming your current working directory is the main amoebae directory that you cloned with Git).
 - >>> bash singularity_jupyter.sh
 - 2. Copy and past the resulting URL (the one at the bottom of the output) into the address bar of your web browser (either Firefox, Chrome, or Safari will work). This will open Jupyter to the notebooks subdirectory, which contains several tutorial and example notebooks (.ipynb files). These files are the files on your regular (host) filesystem, as the amoebae directory is synced with the Singularity container. Thus changes to files will persist after you shut down the Jupyter server and the Singularity container. Documentation on Jupyter is available here: https://jupyternotebook.readthedocs.io/en/stable/.
- 3. Click on one of the tutorial files (.ipynb). These Jupyter notebooks include information on how to use them once opened. The first tutorial (amoebae_tutorial_1.ipynb) provides a simple example of similarity searching with BLASTP using a Jupyter notebook.

 The second tutorial (amoebae_tutorial_2.ipynb) provides an example using most of the similarity searching functionality that AMOEBAE provides.
- 4. To shut down the Jupyter server, click the logout button in the jupyter browser tab(s), and then go to the terminal window that you used to startup the Jupyter server, and press CTRL-C to kill the Jupyter kernel. This will close the Jupyter notebooks, but the

- analysis output files will remain, because they are saved to your amoebae/notebooks folder which is on your host machine and accessed from within the container.
- 5. Working with the Jupyter notebooks interactively in this manner on high-performance computing clusters is likely possible but inconvenient, and procedures will vary. Also, running the tutorial notebooks would require access to the internet from compute nodes (as opposed to login nodes) which may not be supported. Therefore, it is recommended that you run the tutorials on a personal laptop/desktop computer if possible. To run your own notebooks on a cluster, you will need to write a job submission script that 8 will be specific to the cluster, the job scheduler it uses, and your account details. 9 Please refer to documentation provided by your system administrators for this. For an 10 example script that writes a script for running a notebook as a job to a SLURM job 11 scheduler see https://github.com/laelbarlow/amoebae/blob/master/notebooks/ 12 write_notebook_slurm_script.sh. 13

4 2.5 Running AMOEBAE via the command line

- 1. The easiest way to access AMOEBAE dependencies via the command line is to use the bash script provided. If you are running singularity in a virtual machine (e.g., on MacOS), then only one shell session may be opened at once (and these cannot be opened at the same time as the singularity_jupyter.sh script is running Singularity in a virtual machine). Running the script as indicated below will open a shell session in the Singularity container, with the amoebae directory being the only one accessible. Also, the amoebae executable script is added to the \$PATH in the container, so you can run amoebae commands from any directory.

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- 2. You may find it useful to explore and test the environment using the following commands.
 - Print the paths included in the \$PATH variable in the container.

```
>>> tr ':' '\n' <<< "$PATH"
```

• Check the location of the amoebae executable being run from within the container.

```
>>> command -v amoebae
```

• Check that the amoebae executable script can be run (print the help message).

```
>>> amoebae -h
```

• Check that all modules can be imported in all python files in the AMOEBAE code.

```
>>> amoebae check_imports
```

• Check that key dependencies such as BLASTP can be accessed (they are installed in the Singularity container).

3. Again, running AMOEBAE commands on high-performance computing clusters will require you to write custom job submission scripts. Please refer to documentation provided by your system administrator(s) regarding details specific to your cluster, including the job scheduler used. Also, refer to the Singularity documentation for formulating Singularity commands (https://sylabs.io/docs/).

₇ 3 Command reference

- Documentation for each AMOEBAE command and the various options may be accessed from
- 9 the command line via the "-h" options. The following command reference information is the
- output of running amoebae (and each command) with the "-h" option.

$_{11}$ 3.1 amoebae

1

```
usage: amoebae <command> [<args>]
12
13
   Commands for setting up data structure:
       mkdatadir
                         Make a directory with subdirectories and CSV files for
15
                         storing sequence data, etc.
16
17
   Commands for similarity searching:
18
      setup_hmmdb
                         Construct an HMM database (with hmmpress).
19
      add_to_dbs
                         Format and add a file to a formatted directory.
20
      list_dbs
                         Print a list of all usable database files in the database
21
                         directory as defined in the settings file.
                         Add a query file to a formatted directory.
      add_to_queries
23
      list_queries
                         Print a list of all usable query files in the query
24
                         directory as defined in the settings file.
25
      get_redun_hits
                         Run searches with queries to find redundant hits in
26
                         databases (for interpreting results).
27
      setup_fwd_srch
                         Make directory in which to perform forward searches.
28
      run_fwd_srch
                         Perform searches with given queries into given dbs.
      sum_fwd_srch
                         Append information about forward searches to csv summary
30
                         file (this is used to organize reverse searches).
31
                         Make a directory in which to perform reverse searches.
      setup_rev_srch
32
      run_rev_srch
                         Perform searches with given forward search hits into given db.
33
      sum_rev_srch
                         Append information about reverse searches to csv summary
34
35
                         Interpret search results based on summary.
      interp_srchs
36
                         Identify sequences likely encoded on redundant loci
      find_redun_seqs
                         predicted for the same species.
38
      plot
                         Plot search results.
39
```

```
Commands for phylogenetic analysis using a reference tree:
2
      add_to_models
                         Add an alignment, tree, substitution model, names of
3
                         clade-defining sequences to a directory with other models.
                         Print a list of all usable model/reference tree names in
      list_models
5
                         the models directory as defined in the settings file.
      get_alt_topos
                         Take a tree and make copies with every alternative
7
                         topology for the branches connecting the clades of
8
                         interest.
9
10
   Commands for phylogenetic analysis without a reference tree:
11
                         Identify sequences in a tree, and remove them from a
12
      prune
                         given alignment for further phylogenetic analysis.
13
      auto_prune
                         Automatically identify sequences in a tree, and remove
14
                         them from a given alignment for further phylogenetic
15
                         analysis.
16
      reduce_tree
                         Remove terminal nodes from a given tree if there are
17
                         not any sequences with the same name in a given multiple
18
                         sequence alignment file.
19
                         Add constraint commands to MrBayes input file based on a
      constrain_mb
20
                         given tree topology.
      visualize_tree
                         Parse phylogenetic analysis output files for a single
22
                         alignment in a given directory, and write human-readable
23
                         tree figures to PDF files.
24
      replace_seqs
                         Replace sequences in an alignment with their top hits in a
25
                         given fasta file (useful if genomes or taxon selection has
26
                         been updated).
27
   Miscellaneous commands:
29
      csv_to_fasta
                         Generate a fasta file from sequences detailed in a
30
                         spreadsheet of similarity search results.
31
      check_depend
                         Check that all the dependencies are properly installed and
32
                         useable.
33
      check_imports
                         Check that all the import statements used in the AMOEBAE
34
                         repository run without error.
35
      regen_genome_info Write a new genome info spreadsheet file using filenames
36
                         from the Genomes directory.
37
38
   positional arguments:
39
     command
                 Specify one of the functionalities of amoebae.
40
   optional arguments:
42
     -h, --help show this help message and exit
   Copyright 2018 Lael D. Barlow Licensed under the Apache License, Version 2.0
45
   (the "License"); you may not use this file except in compliance with the
46
   License. You may obtain a copy of the License at
   http://www.apache.org/licenses/LICENSE-2.0 Unless required by applicable law
   or agreed to in writing, software distributed under the License is distributed
```

1

```
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```

3 permissions and limitations under the License.

4 3.2 amoebae mkdatadir

```
s usage: amoebae [-h] new_dir_path

Make a directory with subdirectories and CSV files for storing sequence data,
etc.

positional arguments:
    new_dir_path Specify the full file path that you want the new directory to
    have.

potional arguments:
    -h, --help show this help message and exit
```

₆ 3.3 amoebae setup hmmdb

```
usage: amoebae [-h] indirpath
18
   Construct an HMM database (with hmmpress). This is for later sorting of given
19
   sequences into categories based on which HMM the score highest against.
20
21
   positional arguments:
22
     indirpath
                 Path to directory containing amino acid sequence alignment
23
                 file(s) to be constructed into an HMM database using hmmpress
                 from the HMMer3 software package.
25
26
   optional arguments:
27
     -h, --help show this help message and exit
28
```

29 3.4 amoebae add to dbs

```
usage: amoebae [-h] [--split_char SPLIT_CHAR] [--split_pos SPLIT_POS]
                   [--skip_header_reformat] [--auto_extract_accs]
31
                  new_file
   Format and add a file to a formatted directory.
34
35
   positional arguments:
36
     new_file
                            Can be a fasta file (prot or nucl) or HMM databases,
37
                            generated using the hmmpress program in the HMMer
                            software package. Or a GFF3 annotation file.
39
40
   optional arguments:
```

```
-h, --help
                            show this help message and exit
1
     --split_char SPLIT_CHAR
2
                            Character to split the header string on for extracting
3
                            the accession. (default: )
4
     --split_pos SPLIT_POS
5
                            Position that the accession will be in after
                            splitting. (default: 0)
7
     --skip_header_reformat
8
                            Skip reformatting of header lines in input fasta file.
9
                            (default: False)
10
                            Automatically identify accessions/IDs in sequence
     --auto_extract_accs
11
                            headers (overrides split_char and split_pos options
12
                            above). (default: False)
```

$_{14}$ 3.5 amoebae list dbs

```
usage: amoebae [-h]

running

running

running

runn
```

$_{2}$ 3.6 amoebae add_to_queries

```
usage: amoebae [-h] query_file
23
   Add a query file to a formatted directory. This command adds a given sequence
25
   file to the directory with the path that you have specified in the settings.py
   file, and appends a corresponding line to the CSV file that you specified
   (e.g., '0_query_info.csv') to indicate the query title, etc.
  positional arguments:
30
     query_file Path to a sequence file in FASTA format that can be used as a
31
                 similarity search query file. Or path to a directory containing
32
                 only files for addition to the queries. Note: By default, the
33
                 portion of the input filename preceding the first underscore
                 character will be recorded as the "query title", the remaining
35
                 substring preceding the second underscore character will be
36
                 recorded as the taxon (e.g., "Hsapiens"), and the rest of the
37
                 filename preceding the filename extension will be recorded as
38
                 the sequence ID. So the filename might look like this:
39
                 "QUERYTITLE_HSAPIENS_SEQUENCEID.fa". However, the relevant
40
                 information can be revised in the "Queries/O_query_info.csv"
41
                 file afterward if necessary.
42
```

2 3.7 amoebae list queries

```
usage: amoebae [-h]

Print a list of all usable query files in the query directory as defined in the settings file.

optional arguments:

-h, --help show this help message and exit
```

$_{\circ}$ 3.8 amoebae get redun hits

```
usage: amoebae [-h] [--csv_file CSV_FILE] [--query_name QUERY_NAME]
                   [--query_list_file QUERY_LIST_FILE] [--db_name DB_NAME]
12
                   [--db_list_file DB_LIST_FILE] [--query_title QUERY_TITLE]
                   [--outdir OUTDIR]
                   [--blast_report_evalue_cutoff BLAST_REPORT_EVALUE_CUTOFF]
15
                   [--blast_max_target_seqs BLAST_MAX_TARGET_SEQS]
16
                   [--hmmer_report_evalue_cutoff HMMER_REPORT_EVALUE_CUTOFF]
17
                   [--hmmer_report_score_cutoff HMMER_REPORT_SCORE_CUTOFF]
18
                   [--max_number_of_hits_to_summarize MAX_NUMBER_OF_HITS_TO_SUMMARIZE]
19
                   [--num_threads_similarity_searching NUM_THREADS_SIMILARITY_SEARCHING
      ]
21
                   [--predict_redun_hit_selection]
22
                   srch_dir
23
24
   Run searches with queries to find redundant hits in databases (for
25
   interpreting results).
26
   positional arguments:
     srch_dir
                            Path to directory that will contain output directory
29
                            as a subdirectory.
30
31
   optional arguments:
32
     -h, --help
                            show this help message and exit
33
     --csv_file CSV_FILE
                            Path to spreadsheet to append summary of result to for
34
                            manual annotation. (default: None)
35
     --query_name QUERY_NAME
36
                            Query filename to use (not full path). (default: None)
37
     --query_list_file QUERY_LIST_FILE
38
                            Path to file containing a list of query files to use,
39
                            if no query_name is specified (or all queries by
40
                            default). (default: None)
41
                            Name of database file in the database directory in
     --db_name DB_NAME
42
                            which to do searches (not full path). (default: None)
     --db_list_file DB_LIST_FILE
```

```
Path to file containing a list of database files to
1
                            use (if no db_name specified). (default: None)
2
     --query_title QUERY_TITLE
3
                            Name to be assigned to hits in databases that may be
                            considered redundant with a search query to which the
                            same title is assigned, otherwise it is taken from the
                            query info spreadsheet specified in the settings.py
7
                            file ('query_info_csv'). (default: None)
8
     --outdir OUTDIR
                            Path to directory to write search results to.
9
                            (default: None)
10
     --blast_report_evalue_cutoff BLAST_REPORT_EVALUE_CUTOFF
11
                            Maximum E-value for reporting BLAST hits. (default:
12
                            0.05)
13
     --blast_max_target_seqs BLAST_MAX_TARGET_SEQS
14
                            Maximum BLAST target sequences to consider. (default:
15
                            500)
16
     --hmmer_report_evalue_cutoff HMMER_REPORT_EVALUE_CUTOFF
17
                            Maximum E-value for reporting HMMer hits. (default:
18
                            0.05)
19
     --hmmer_report_score_cutoff HMMER_REPORT_SCORE_CUTOFF
20
                            Minimum sequence score for reporting HMMer hits.
21
                            (default: 5)
22
     --max_number_of_hits_to_summarize MAX_NUMBER_OF_HITS_TO_SUMMARIZE
23
                            Absolute maximum number of hits (BLAST, HMMer, etc) to
24
                            summarize in the output spreadsheet. This is important
25
                            when working with sequences with WD40 domains, for
26
                            example. (default: 50)
27
     --num_threads_similarity_searching NUM_THREADS_SIMILARITY_SEARCHING
28
                            Number of threads to use for running searches.
29
                            (default: 4)
30
     --predict_redun_hit_selection
31
                            Write a copy of the output spreadsheet with '+' in
32
                            rows for hits that may be specific to each query
33
                            title, due to not being retrieved as top hits by
34
                            queries associated with different query titles.
35
                            (default: False)
36
37
   Recommendation: For most analyses, use the --query_name option and the
38
   --db_name option, and run the get_redun_hits command for each query
39
   separately. Otherwise, there will be redundant information in the output
40
   spreadsheet(s).
         amoebae setup fwd srch
   3.9
   usage: amoebae [-h] [--outdir OUTDIR] srch_dir query_list_file db_list_file
43
  Make a directory in which to write output files from similarity searches.
```

46

positional arguments:

```
srch_dir
                      Path to directory that will contain output directory as a
1
                      subdirectory.
2
     query_list_file Path to file with list of queries to search with.
3
     db_list_file
                      Path to file with list of databases to search with.
   optional arguments:
     -h, --help
                      show this help message and exit
7
     --outdir OUTDIR
                      Path to directory to put search results into (so that this
8
                      step can be piped together with other commands). (default:
                      None)
10
   Note: Use the bash script to run forward searches on a remote server.
   3.10
           amoebae run fwd srch
   usage: amoebae [-h] [--blast_report_evalue_cutoff BLAST_REPORT_EVALUE_CUTOFF]
14
                   [--blast_max_target_seqs BLAST_MAX_TARGET_SEQS]
15
                   [--hmmer_report_evalue_cutoff HMMER_REPORT_EVALUE_CUTOFF]
16
                   [--hmmer_report_score_cutoff HMMER_REPORT_SCORE_CUTOFF]
17
                   [--num_threads_similarity_searching NUM_THREADS_SIMILARITY_SEARCHING
18
      ]
19
                  fwd_srch_dir
   Perform searches with original queries into subject databases.
22
23
   positional arguments:
24
                            Path to directory that will contain forward search
     fwd_srch_dir
25
                            output files.
26
   optional arguments:
28
     -h, --help
                            show this help message and exit
29
     --blast_report_evalue_cutoff BLAST_REPORT_EVALUE_CUTOFF
30
                            Maximum E-value for reporting BLAST hits. (default:
31
                            0.05)
32
     --blast_max_target_seqs BLAST_MAX_TARGET_SEQS
33
                            Maximum BLAST target sequences to consider. (default:
                            500)
35
     --hmmer_report_evalue_cutoff HMMER_REPORT_EVALUE_CUTOFF
36
                            Maximum E-value for reporting HMMer hits. (default:
37
                            0.05)
38
     --hmmer_report_score_cutoff HMMER_REPORT_SCORE_CUTOFF
39
                            Minimum sequence score for reporting HMMer hits.
40
                            (default: 5)
     --num_threads_similarity_searching NUM_THREADS_SIMILARITY_SEARCHING
42
                            Number of threads to use for running searches.
                            (default: 4)
```

3.11amoebae sum fwd srch

48

```
usage: amoebae [-h] [--max_evalue MAX_EVALUE]
                   [--max_gap_between_tblastn_hsps MAX_GAP_BETWEEN_TBLASTN_HSPS]
3
                   [--do_not_use_exonerate]
4
                   [--exonerate_score_threshold EXONERATE_SCORE_THRESHOLD]
5
                   [--max_hits_to_sum MAX_HITS_TO_SUM]
6
                   [--max_length_diff MAX_LENGTH_DIFF]
                  fwd_srch_out csv_file
   Append information about forward searches to csv summary file (this is used to
10
   organize reverse searches). For TBLASTN searches (protein queries, nucleotide
11
   target sequences), HSPs are clustered into groups that are close enough within
12
   the target sequence to potentially represent exons from the same coding
   sequence. The nucleotide subsequences in which these clusters of HSPs are
   found are then analyzed using exonerate to identify and translate potential
   exons, in "protein2genome" mode, because exonerate, unlike TBLASTN, attempts
   to identify exon boundaries, yielding translations that are less likely to
17
   include translations of non-coding regions outside exons (which might include
   apparent stop codons).
19
20
   positional arguments:
21
                            Path to directory where forward search results were
     fwd_srch_out
22
                            written.
23
     csv_file
                            Path to summary spreadsheet (CSV) file, which may
                            already contain search summaries, or may not exist
25
                            yet.
26
27
   optional arguments:
28
     -h, --help
                            show this help message and exit
29
     --max_evalue MAX_EVALUE
30
                            Maximum E-value threshold for reporting forward search
31
                            hits. (default: 0.0005)
32
     --max_gap_between_tblastn_hsps MAX_GAP_BETWEEN_TBLASTN_HSPS
33
                            Maximum number of nucleotide bases between TBLASTN
34
                            HSPs to be considered part of the same gene locus.
35
                            This is important, because it will be assumed that HSP
36
                            separated by more than this number of nucleotide bases
37
                            are not part of the same gene or TBLASTN "hit".
38
                            (default: 10000)
39
     --do_not_use_exonerate
40
                            Override the default use of exonerate to identify
41
                            coding sequences and translations, and just use
42
                            TBLASTN instead. This option is provided because
43
                            concatenated TBLASTN HSPs may be more inclusive of
44
                            sequences within the target sequence, and the results
45
                            of TBLASTN and exonerate may need to be compared.
46
                            Also, note that HSPs identified by TBLASTN but for
47
                            which exonerate yields no alignments will be ignored
```

```
if exonerate is used. (default: False)
1
     --exonerate_score_threshold EXONERATE_SCORE_THRESHOLD
2
                            Set score threshold to be applied when running
3
                            exonerate on nucleotide sequences identified by
                            TBLASTN. The default for setting of exonerate is 100,
                            but a lower score is set as default here, because
                            otherwise exonerate cannot identify some of the
7
                            sequences identified by TBLASTN. This option is only
8
                            relevant if using exonerate. (default: 10)
9
     --max_hits_to_sum MAX_HITS_TO_SUM
10
                            Maximum number of forward search hits to list in the
11
                            summary spreadsheet. If zero, then reverse searches
12
                            will be performed for all hits. (default: 0)
13
     --max_length_diff MAX_LENGTH_DIFF
14
                            Maximum number of amino acid residues length
15
                            difference allowed between the original query and the
16
                            forward hit sequence. If -1, then a maximum length
17
                            cutoff will not be applied. (default: -1)
18
           amoebae setup rev srch
   3.12
   usage: amoebae [-h] [--outdir OUTDIR] [--aasubseq] [--nafullseq]
20
                  srch_dir csv_file databases
21
22
   Make directory in which to write results of reverse searches.
23
24
   positional arguments:
25
     srch_dir
                      Path to directory that will contain output directory as a
26
                      subdirectory.
                      Path to summary spreadsheet (CSV) file, which contains a
     csv_file
28
                      summary of forward search(es).
29
     databases
                      Database filename (in database directory) or path to file
30
                      with list of database filenames. Note that filenames are
31
                      needed, not file paths.
32
33
   optional arguments:
     -h, --help
                      show this help message and exit
35
     --outdir OUTDIR
                      Path to directory to put search results into (so that this
36
                      step can be piped together with other commands). (default:
37
                      None)
38
     --aasubseq
                      Use only the portion of each (amino acid) forward hit
39
                      sequence that aligns to the original query used (top HSP
40
                      subject sequence). This is default for nucleotide hits.
41
                       (default: False)
42
     --nafullseq
                      Use the full (nucleic acid) forward hit sequence. This is
43
```

default for amino acid hits. (default: False)

~ 3.13 amoebae run rev srch

rev_srch_out

46

```
usage: amoebae [-h] [--blast_report_evalue_cutoff BLAST_REPORT_EVALUE_CUTOFF]
                   [--blast_max_target_seqs BLAST_MAX_TARGET_SEQS]
3
                   [--hmmer_report_evalue_cutoff HMMER_REPORT_EVALUE_CUTOFF]
                   [--hmmer_report_score_cutoff HMMER_REPORT_SCORE_CUTOFF]
                   [--num_threads_similarity_searching NUM_THREADS_SIMILARITY_SEARCHING
      ]
                  rev_srch_dir
8
   Perform searches with forward search hit sequences as queries into the
10
   original query databases.
   positional arguments:
13
     rev_srch_dir
                            Path to directory that will contain output of
14
                            searches.
15
16
   optional arguments:
17
     -h, --help
                            show this help message and exit
18
     --blast_report_evalue_cutoff BLAST_REPORT_EVALUE_CUTOFF
19
                            Maximum E-value for reporting BLAST hits. (default:
                            0.05)
     --blast_max_target_seqs BLAST_MAX_TARGET_SEQS
22
                            Maximum BLAST target sequences to consider. (default:
23
24
     --hmmer_report_evalue_cutoff HMMER_REPORT_EVALUE_CUTOFF
25
                            Maximum E-value for reporting HMMer hits. (default:
26
                            0.05)
27
     --hmmer_report_score_cutoff HMMER_REPORT_SCORE_CUTOFF
28
                            Minimum sequence score for reporting HMMer hits.
29
                            (default: 5)
30
     --num_threads_similarity_searching NUM_THREADS_SIMILARITY_SEARCHING
31
                            Number of threads to use for running searches.
32
                            (default: 4)
33
   3.14
           amoebae sum rev srch
   usage: amoebae [-h] [--redun_hit_csv REDUN_HIT_CSV]
35
                   [--min_evaldiff MIN_EVALDIFF] [--aasubseq] [--nafullseq]
36
                   [--max_rev_srchs MAX_REV_SRCHS]
                  csv_file rev_srch_out
38
39
   Append information about reverse searches to csv summary file. Use information
40
   from redundant hit csv file to interpret results.
41
42
  positional arguments:
43
     csv_file
                            Path to summary spreadsheet (CSV) file, which may
44
                            already contain reverse search summaries.
45
```

Path to directory where reverse search results were

written. 1 2 optional arguments: 3 -h, --help show this help message and exit --redun_hit_csv REDUN_HIT_CSV Path to spreadsheet (CSV) file, which specifies which hits are redundant positive hits for a given query 7 (query title) in a given database. If this is not 8 provided, then it is assumed that any and all reverse 9 search hits are equivalent to/redundant with the 10 original query. (default: None) 11 --min_evaldiff MIN_EVALDIFF 12 Minimum difference in E-value order of magnitude 13 between top reverse search hit and first reverse 14 search hit that is not redundant with the original 15 query. (default: 5) 16 --aasubseq Use only the portion of each (amino acid) forward hit 17 sequence that aligns to the original query used (top 18 HSP subject sequence). This is default for nucleotide 19 hits. Must be selected if selected when the 20 setup_rev_srch command was run. (default: False) Use the full (nucleic acid) forward hit sequence. This --nafullseq 22 is default for amino acid hits. Must be selected if 23 selected when the setup_rev_srch command was run. 24 (default: False) 25 --max_rev_srchs MAX_REV_SRCHS 26 Maximum number of forward search hits to perform 27 reverse searches for per query database. If zero, then reverse searches will be performed for all hits. 29 (default: 0) 30 3.15amoebae interp srchs usage: amoebae [-h] [--fwd_only] [--fwd_evalue_cutoff FWD_EVALUE_CUTOFF] 32 [--rev_evalue_cutoff REV_EVALUE_CUTOFF] 33 [--hmmer_cutoff HMMER_CUTOFF] [--no_overlapping_hits] 34 [--out_csv_path OUT_CSV_PATH] 35 csv_file 36 Interpret search results based on final summary, which provides a basis for further analyses of positive hits. 39 40 positional arguments: 41 csv_file Path to spreadsheet with forward and reverse search 42 results. 43 optional arguments: 45 -h, --help show this help message and exit

Interpret forward searches based on score (HMMer)

46

47

--fwd_only

```
cutoff. (default: False)
1
     --fwd_evalue_cutoff FWD_EVALUE_CUTOFF
2
                            Specify an (more stringent) E-value cutoff for forward
3
                            search results. (default: None)
4
     --rev_evalue_cutoff REV_EVALUE_CUTOFF
5
                            Specify an (more stringent) E-value cutoff for reverse
                            search results. (default: None)
7
     --hmmer_cutoff HMMER_CUTOFF
8
                            Specify a score that hits must exceed to be included.
9
                            (default: 20)
10
     --no_overlapping_hits
11
                            If more than one query (query title) retrieves a given
12
                            sequence as a positive hit based on the search
13
                            criteria, make the sequence a negative hit for all
14
                            queries (query titles), except for the one that
15
                            retrieved the sequence with the lowest (strongest)
16
                            E-value. Warning: Do not use this option if you are
17
                            searching sequences that include genomic sequences
18
                            that may include more than one genomic locus per
19
                            sequence. False-negative results could occur in this
20
                            case, because different queries for non-orthologous
                            genes could retrieve subsequences in the same subject
22
                            sequence. (default: False)
23
     --out_csv_path OUT_CSV_PATH
24
                            Optionally specify an output file path, so that this
25
                            command can be piped together with others. (default:
26
                            None)
27
```

$_{88}$ 3.16 amoebae find_redun_seqs

47

```
usage: amoebae [-h] [--out_csv_path OUT_CSV_PATH]
29
                   [--remove_tblastn_hits_at_annotated_loci]
30
                   [--just_look_for_genes_in_gff3] [--ignore_gff3]
31
                   [--allow_internal_stops ALLOW_INTERNAL_STOPS]
32
                   [--min_length MIN_LENGTH]
33
                   [--min_percent_length MIN_PERCENT_LENGTH]
34
                   [--min_percent_query_cover MIN_PERCENT_QUERY_COVER]
35
                   [--overlap_required] [--max_percent_ident MAX_PERCENT_IDENT]
36
                   [--min_alig_res_in_overlap MIN_ALIG_RES_IN_OVERLAP]
37
                   [--min_ident_res_in_overlap MIN_IDENT_RES_IN_OVERLAP]
38
                   [--min_sim_res_in_overlap MIN_SIM_RES_IN_OVERLAP]
39
                   [--min_ident_span_len MIN_IDENT_SPAN_LEN]
40
                   [--min_sim_span_len MIN_SIM_SPAN_LEN]
41
                   [--min_percent_ident_in_overlap MIN_PERCENT_IDENT_IN_OVERLAP]
42
                   [--min_percent_sim_in_overlap MIN_PERCENT_SIM_IN_OVERLAP]
43
                   [--min_percent_overlap MIN_PERCENT_OVERLAP]
44
                   [--plot_hit_exclusion] [--add_ali_col]
45
                   csv_file
46
```

```
Identify hit sequences likely encoded by the same gene loci in the genome of a
  given species, or otherwise not representing paralogous genes. Criteria are
   applied in this order: 1. Peptide hits with the same ID as higher-ranking hits
  for the same query (query title) are excluded. 2. Nucleotide hits for the same
  loci as peptide sequence hits are excluded. 3. Sequences with internal stop
  codons are excluded, as these are potentially pseudogenes. 4. Sequences are
   excluded if they do not meet several minimum length criteria: Absolute minimum
  length (in amino acids) and percent query cover. 5. Sequences are excluded if
   they do not overlap to a specified degree with all included higher-ranking
  hits for the same query (query title) in sequence data for the same
  species/genome. This is determined by algorithmically comparing pairs of
   sequences aligned to a reference alignment of homologues, and several minimum
  measures of alignment overlap may be specified. 6. Secondary hit sequences are
   excluded if they do not meet a specified maximum percent identity threshold.
  Highly identical sequences may result from false segmental duplications in the
15
   genome assembly, may represent alleles, etc. Note: Applying these criteria
16
   requires a column to be manually added to the input csv file prior to running
17
  with the header "Alignment for sequence comparison" and filled with the
   appropriate alignment name to use (one for each query title, as listed in the
   "Query title" column). Alternatively, you can run this command with the
   --add_ali_col option to automatically identify appropriate alignments among
   your aligned FASTA queries used for running HMMer searches. If no alignment
22
   (.afaa) file can be found, then the first single sequence query file (.faa)
23
   that appears in the summary CSV file will be used instead.
24
25
  positional arguments:
26
     csv_file
                           Path to spreadsheet with interpreted search results
27
                           outputted by the interp_srchs command.
28
29
   optional arguments:
30
                           show this help message and exit
     -h, --help
31
     --out_csv_path OUT_CSV_PATH
32
                           Optionally specify an output file path, so that this
33
                           command can be piped together with others. (default:
34
                           None)
35
     --remove_tblastn_hits_at_annotated_loci
36
                           Ignore tblastn hits that overlap with any previously
37
                           annotated loci. The rationale for this would be that
38
                           the corresponding protein sequences should have been
39
                           retrieved if the tblastn hit were a true positive
40
                           anyway. If this option is not specified, then
                           sequences will still be excluded if they specifically
42
                           correspond to the same loci as do higher-ranking hits.
                           (default: False)
     --just_look_for_genes_in_gff3
45
                           When looking for records in GFF3 annotation files that
46
                           overlap with subsequences identified by similarity
47
                           searching (TBLASTN), ignore records that are not
48
                           explicitly "gene" (for example, "CDS", "mRNA", and
49
```

1		"exon"). This option should probably not be selected,
2		because in some GFF3 annotation files do not include
3		"gene" records, but do include predicted coding
4		sequences for genes. (default: False)
5	ignore_gff3	Disregard any information regarding redundancy of
6		identified nucleotide sequences with identified
7		protein sequences that may be found in GFF3 annotation
8		files. (default: False)
9	allow_internal_stops	s ALLOW_INTERNAL_STOPS
10	_	Include sequences that have internal stop codons
11		(anywhere other than the N-terminal position).
12		(default: True)
13	min_length MIN_LENGT	TH
14		Absolute minimum length (in AA) of a hit sequence to
15		be considered a potential distinct paralogue.
16		(default: 55)
17	min_percent_length M	MIN_PERCENT_LENGTH
18	-	Minimum length (in AA) of a hit sequence as a
19		percentage of query length for the hit to be
20		considered a potential distinct paralogue. (default:
21		15)
22	min_percent_query_co	over MIN_PERCENT_QUERY_COVER
23		Minimum number of residues aligning with the original
24		query as a percentage of the original query sequence
25		length. (default: 0)
26	overlap_required	True if hits must overlap with a higher-ranking hit to
27		be considered potential unique paralogues. (default:
28		False)
29	max_percent_ident MA	AX_PERCENT_IDENT
30		Maximum percent identity (among aligning residues) for
31		evaluating whether two sequences are redundant or not
32		(secondary hits showing a percent identity with a
33		higher-ranking hit exceeding this value will be
34		excluded). (default: 98.0)
35	min_alig_res_in_over	rlap MIN_ALIG_RES_IN_OVERLAP
36		Minimum number of residues which must align for two
37		sequences to be considered as potentially distinct
38		hits. This is only relevant if the overlap_required
39		option is specified. (default: 20)
40	min_ident_res_in_ove	erlap MIN_IDENT_RES_IN_OVERLAP
41		Minimum number of aligning residues which must be
42		identical for two sequences to be considered as
43		potentially distinct hits. This is only relevant if
44		the overlap_required option is specified. (default:
45		10)
46	min_sim_res_in_over	ap MIN_SIM_RES_IN_OVERLAP
47		Minimum number of aligning residues which must be
48		similar for two sequences to be considered as
49		potentially distinct hits. This is only relevant if

```
the overlap_required option is specified. (default:
1
                            15)
2
     --min_ident_span_len MIN_IDENT_SPAN_LEN
3
                            Minimum number of aligning residues which are
                            identical that must exist in at least one continuous
                            span for two sequences to be considered as potentially
                            distinct hits (not counting positions where both
7
                            sequences have gaps). This is only relevant if the
8
                            overlap_required option is specified. (default: 0)
9
     --min_sim_span_len MIN_SIM_SPAN_LEN
10
                            Minimum number of aligning residues which are similar
11
                            (or identical) that must exist in at least one
12
                            continuous span for two sequences to be considered as
                            potentially distinct hits (not counting positions
14
                            where both sequences have gaps). This is only relevant
15
                            if the overlap_required option is specified. (default:
16
17
     --min_percent_ident_in_overlap MIN_PERCENT_IDENT_IN_OVERLAP
18
                            Minimum percent identity between the two sequences of
19
                            interest in the alignment. This is only relevant if the
20
                            overlap_required option is specified. (default: 0)
     --min_percent_sim_in_overlap MIN_PERCENT_SIM_IN_OVERLAP
22
                            Minimum percent similarity (including identity)
23
                            between the two sequences of interest in the
24
                            alignment. This is only relevant if the
25
                            overlap_required option is specified. (default: 0)
26
     --min_percent_overlap MIN_PERCENT_OVERLAP
27
                            Minimum number of aligning residues between the two
28
                            sequences of interest as a percentage of the length of
29
                            the second sequence (the last sequence in the
30
                            alignment), not including gaps, for the two sequences
31
                            to be considered as potentially distinct hits. This is
32
                            only relevant if the overlap_required option is
33
                            specified. (default: 0)
34
                           Plot number of hits excluded by the various criteria
     --plot_hit_exclusion
35
                            applied. (default: False)
36
     --add_ali_col
                            Add a column to the csv file listing which alignment
37
                            file in the queries directory to use for comparing
38
                            sequences. Aligned FASTA queries are selected that
39
                            match the query titles of the original queries used to
40
                            retrieve each of the relevant hits listed in the csv
41
                            file. No other options need to be specified in this
42
                            case. (default: False)
```

$_{*}$ 3.17 amoebae plot

```
usage: amoebae [-h] [--csv_file2 CSV_FILE2] [--complex_info COMPLEX_INFO]
[--row_order ROW_ORDER] [--out_pdf OUT_PDF]
csv_file
```

1 Plot results of similarity search and sequence classification analyses. The outputs are PDF files. positional arguments: csv_file Path to a spreadsheet with the relevant results to be plotted. This can be either a CSV file output of the 7 sum_rev_srch command or from the find_redun_seqs 8 command. If the output of the sum_rev_srch command is 9 used, however, redundant hits will be counted (e.g., 10 BLASTP and TBLASTN hits corresponding to the same or 11 highly identical genomic loci). 12 optional arguments: 14 -h, --help show this help message and exit 15 --csv_file2 CSV_FILE2 16 Path to a second spreadsheet with relevant results to 17 be compared to the first and plotted. (default: None) 18 --complex_info COMPLEX_INFO 19 Path to file that specifies which query titles represent components of which protein complexes (or otherwise grouped proteins). (default: None) 22 --row_order ROW_ORDER 23 Path to file that specifies the order in which data 24 for each species will be displayed. (default: None) 25 --out_pdf OUT_PDF Path to output pdf file. (default: None) 26 3.18amoebae add to models usage: amoebae [-h] 28 model_name alignment tree_topology subs_model type_seqs taxon 29 Add a phylogenetic model for relationships between members of a gene family (sequence_data matrix, data type, tree topology, type sequence defining each 32 clade of interest, and substitution model) to a directory for use in 33 classifying sequence (via the 'phylo_class' command. 34 35 positional arguments: 36 An arbitrary name for the model (which will refer to the model_name 37 alignment, tree, substitution model, etc. collectively). 38 A multiple amino acid sequence alignment in nexus format. alignment 39 tree_topology Text file containing a tree (identified previously using 40 MrBayes, etc) containing the names of all the sequences in 41 the alignment, in newick format. 42 The name of the substitution model used to recover the subs_model 43 provided topology (chosen with ModelFinder or similar 44 software). 45

type_seqs

46

47

Names of sequences (sequence headers) that are to be used to

define clades of interest. A csv file with seq names in one

```
column and clade names in the next column.
1
                    Taxonomic group represented in the model (e.g., "Eukaryotes",
     taxon
2
                    or "Amorphea").
3
   optional arguments:
     -h, --help
                    show this help message and exit
  3.19
           amoebae list models
   usage: amoebae [-h]
  Print a list of all usable model/reference tree names in the models directory
10
   as defined in the settings file.
   optional arguments:
     -h, --help show this help message and exit
           amoebae get alt topos
   3.20
   usage: amoebae [-h] [--polytomy] [--not_polytomy_clades]
16
                   [--keep_original_backbone] [--iqtree_au_test]
17
                  model_name out_dir_path
18
19
   Take a tree and make copies with every alternative topology for the branches
   connecting the clades of interest. Output as additional models in the Models
   directory.
22
23
   positional arguments:
24
     model_name
                           Name of model/backbone tree to modify (other info
25
                            provided in the model info csv file).
26
     out_dir_path
                           Path to directory in which output directory will be
27
                            written.
28
   optional arguments:
30
     -h, --help
                            show this help message and exit
31
                            Just make one big polytomy connecting the clades of
     --polytomy
32
                            interest intead of making alternative bifurcating
33
                            trees. (default: False)
34
     --not_polytomy_clades
35
                            Do not make subtrees/clades of interest polytomies in
36
                            output topologies. (default: False)
37
     --keep_original_backbone
38
                            Keep the original backbone topology instead of
39
                            generating a polytomy or alternative resolved
40
                            topologies. (default: False)
41
                            Test all the relevant alternative topologies against
     --iqtree_au_test
42
                            each other using Approximately Unbiased (AU) test with
                            IQ-tree. (default: False)
```

3.21 amoebae prune

```
usage: amoebae [-h] [--include_seqs] [--output_file OUTPUT_FILE]
                   tree_file alignment name_replace_table
3
   Identify sequences in a tree, and remove them from a given alignment for
5
   further phylogenetic analysis.
   positional arguments:
     tree_file
                            Tree in newick format (coded names, because ETE3
                            cannot parse taxon names with space characters without
10
                            quotation marks around them).
11
     alignment
                            Dataset used to make the tree (nexus alignment)
12
                            (original alignment with original taxon names either
13
                            trimmed or untrimmed).
     name_replace_table
                            File for decoding names in input tree file.
15
16
   optional arguments:
17
     -h, --help
                            show this help message and exit
18
     --include_seqs
                            Include only listed sequences/nodes instead of
19
                            removing them. (default: False)
20
     --output_file OUTPUT_FILE
21
                            Path to output file. (default: None)
22
```

3.22 amoebae auto_prune

```
usage: amoebae [-h]
24
                   [--max_bl_iqr_above_third_quartile MAX_BL_IQR_ABOVE_THIRD_QUARTILE]
25
                   [--remove_redun_seqs REMOVE_REDUN_SEQS]
26
                   [--remove_redun_seqs_threshold REMOVE_REDUN_SEQS_THRESHOLD]
27
                   [--output_file OUTPUT_FILE]
28
                   in_dir
29
30
   Automatically identify sequences in a tree, and remove them from a given
31
   alignment for further phylogenetic analysis.
32
33
   positional arguments:
34
     in_dir
                            Path to directory that contains the phylogenetic
35
                            analysis output files (sequence name conversion table
36
                            file and original nexus alignment file can be in the
                            parent directory to this directory as long as their
38
                            names are mostly identical.
39
40
   optional arguments:
41
     -h, --help
                            show this help message and exit
42
     --max_bl_iqr_above_third_quartile MAX_BL_IQR_ABOVE_THIRD_QUARTILE
43
                            Inclusion threshold for number of interquartile ranges
44
                            above the third quartile of terminal branch lengths
45
                            the length of a terminal branch can be before it is
46
```

```
considered an outlier (length is total distance from
1
                            root node after rooting on midpoint, or the longest
2
                            terminal branch on either side of the midpoint).
3
                            (default: 1.5)
     --remove_redun_segs REMOVE_REDUN_SEQS
5
                            Remove taxonomically redundant sequences (longest
                            branch of two sister branches when both are sequences
7
                            from the same species. (default: True)
8
     --remove_redun_seqs_threshold REMOVE_REDUN_SEQS_THRESHOLD
9
                            Minimum support required to consider one of two sister
10
                            branches/sequences taxonomically redundant. Note: only
11
                            used if the remove_redun_seqs option is specified.
12
                            (default: 0.95)
     --output_file OUTPUT_FILE
14
                            Path to output file. (default: None)
15
   3.23
           amoebae reduce tree
   usage: amoebae [-h] [--output_file OUTPUT_FILE] alignment tree_file
17
18
   Remove terminal nodes from a given tree if there are not any sequences with
   the same name in a given alignment.
   positional arguments:
22
     alignment
                            Alignment in nexus format with sequences representing
23
                            a subset of those represented in the input tree.
24
                            Tree in newick format.
     tree_file
25
26
   optional arguments:
     -h, --help
                            show this help message and exit
     --output_file OUTPUT_FILE
29
                            Path to output file. (default: None)
30
   3.24
           amoebae constrain mb
   usage: amoebae [-h] [--out_alignment OUT_ALIGNMENT] alignment tree
32
33
   Add constraint commands to MrBayes input file.
35
   positional arguments:
36
                            Nexus alignment for input to Mrbayes (without any
37
     alignment
                            constraint commands).
38
     tree
                            Tree in newick format with same taxon names as in
39
                            alignment. To be used as a topology constraint (all
40
                            nodes).
41
42
   optional arguments:
     -h, --help
                            show this help message and exit
```

```
--out_alignment OUT_ALIGNMENT
                           Path to nexus alignment for input to Mrbayes with
2
                            constraints added. (default: None)
3
   3.25
           amoebae visualize tree
   usage: amoebae [-h] [--root_taxon ROOT_TAXON] [--highlight_paralogues]
                   [--add_clade_names_from_file]
                  input_directory method
   Parse phylogenetic analysis output files in a given directory, and write
   human-readable tree figures to PDF files.
10
   positional arguments:
     input_directory
                           Path to directory containing input files (must contain
                            a .table file for decoding taxon names.
14
     method
                           Name of tree searching program used. Either iqtree,
15
                           raxml, or mrbayes accepted.
16
17
   optional arguments:
18
     -h, --help
                            show this help message and exit
19
     --root_taxon ROOT_TAXON
20
                            Name of species to root on (e.g.,
21
                            "Klebsormidium_nitens").
22
     --highlight_paralogues
23
                            Highlight clades that contain paralogues found in at
24
                            least one other clade in the tree.
25
     --add_clade_names_from_file
26
                            Use a file in the parent directory with clade names
27
                            corresponding to representative sequences to add clade
                            names to all the taxon names in the output trees.
   3.26
           amoebae replace seqs
   usage: amoebae [-h] [--fasta_file FASTA_FILE] alignment
31
32
  Replace sequences in an alignment the full-length sequences from the relevant
33
   file(s) in the Genomes directory, or with their top hits in a given fasta
   file. And, align, mask, and trim the identified sequences to the input
   alignment
36
37
   positional arguments:
38
     alignment
                           Path to multiple sequence alignment file in nexus
39
                            format (trimmed alignment).
40
   optional arguments:
42
```

show this help message and exit

-h, --help

--fasta_file FASTA_FILE

Path to file containing sequences with which to replace sequences in the alignment. If this option is not specified, then full-length sequences will be retrieved from files in the Genomes directory.

$_{5}$ 3.27 amoebae csv_to_fasta

```
usage: amoebae [-h] [--output_dir OUTPUT_DIR] [--abbrev] [--paralogue_names]
                   [--only_descr] [--subseq] [--all_hits] [--split_by_query_title]
                   [--split_by_top_rev_srch_hit SPLIT_BY_TOP_REV_SRCH_HIT]
                   [--split_to_query_fastas]
                   csv_file
10
11
   Extract sequences described in a spreadsheet output by AMOEBAE, and write to a
12
   file in FASTA format.
13
   positional arguments:
15
     csv_file
                            Path to csv file listing sequences.
16
17
   optional arguments:
18
                            show this help message and exit
     -h, --help
19
     --output_dir OUTPUT_DIR
20
                            Path for output directory to contain FASTA files.
21
                            (default: None)
22
     --abbrev
                            Add species name instead of sequence description from
23
                            fasta header. Applicable when the output file is to be
24
                            used for alignment and phylogenetic analysis.
25
                            (default: False)
26
                            Use species name, query title, and paralogue number
     --paralogue_names
27
                            instead of sequence description from fasta header.
28
                            Applicable when the output file is to be used for
29
                            alignment and phylogenetic analysis. Does not work if
30
                            the abbrev option is specified. (default: False)
31
     --only_descr
                            Use the description but not the ID as the new fasta
32
                            sequence header. Does not work if the abbrev option is
33
                            specified. (default: False)
34
     --subseq
                            Write subsequences that aligned to forward search
35
                            query, instead of the full sequences. (default: False)
36
     --all_hits
                            Write all forward hits listed in the input csv file.
37
                            (default: False)
38
     --split_by_query_title
39
                            Write sequences to files according to the query title
40
                            of the query which retrieved them in a similarity
41
                            search. (default: False)
42
     --split_by_top_rev_srch_hit SPLIT_BY_TOP_REV_SRCH_HIT
43
                            Write sequences to files according to the top hit that
44
                            they retrieve in a reverse search, for each sequence
45
                            that meets the reverse search criteria. (Provide the
46
                            reverse search identifier, eg,
47
```

```
"rev_srch_20180924122402-1") (default: None)
1
     --split_to_query_fastas
2
                           Write sequences to separate files with filenames that
3
                           can be easily parsed for loading the the files as
                           queries using the add_to_queries command. (default:
                           False)
   3.28
          amoebae check depend
   usage: amoebae [-h]
   Check that all the dependencies (other than python modules) are properly
   installed and useable.
   optional arguments:
13
     -h, --help show this help message and exit
14
   3.29
          amoebae check imports
  usage: amoebae [-h]
   Check that all the import statements used in the AMOEBAE repository run
18
   without error.
19
20
   optional arguments:
21
     -h, --help show this help message and exit
22
   3.30
          amoebae regen genome info
   usage: amoebae [-h] data_dir_path
25
  Write a new genome info spreadsheet (O_genome_info.csv) file using filenames
26
   from the Genomes directory.
28
   positional arguments:
     data_dir_path Specify the full path to an existing AMOEBAE data directory,
30
                    which contains a 'Genomes' subdirectory. The new genome info
31
                    file will be added to this subdirectory.
32
33
```

$_{\scriptscriptstyle 6}$ 4 Miscellaneous scripts

optional arguments:

-h, --help

34

35

Several scripts of less general applicablity than the amoebae commands descibed above are included in the AMOEBAE toolkit. See the amoebae/misc_scripts directory (https:

show this help message and exit

- 1 //github.com/laelbarlow/amoebae/tree/master/misc_scripts). Most scripts have in-
- ² formation regarding usage in the files themselves. More detailed information regarding some
- of these scripts may be added to this documentation in the future.

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