# e-Finder

A tool to find multigene elements in assembled sequences using profile HMMs

# A Quick Guide

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# e-Finder - A Quick Guide

#### 1 Introduction

e-Finder is a generic tool for detecting and extracting multigene elements from assembled genomes using profile HMMs. e-Finder executes hmmsearch program (HMMER package) to run similarity searches using profile HMMs as queries against translated sequences of the assembled genomes. Any region containing a cluster of genes can be detected, such as transposons, CRISPR-Cas systems, prophages, operons, etc.

#### 2 Main features

- Fully configurable
- FASTA files containing finished or partially assembled genomes can be used as input
- Previously run hmmsearch results can be reprocessed with new sets of parameters
- Score and e-value cutoff values can be used to set the stringency of the search

#### 3 Before using e-Finder

#### 3.1 System requirements

e-Finder was developed in Perl language and can be used in any POSIX compliant operating system such as UNIX and Linux distributions with an installed Perl interpreter (<a href="http://www.perl.org">http://www.perl.org</a>).

### 3.2 Third-party programs

e-Finder requires the following programs and databases:

- transeq (EMBOSS package <a href="http://emboss.sourceforge.net/">http://emboss.sourceforge.net/</a>). This program is used to translate assembled nucleotide sequences into the six possible frames.
- hmmsearch (HMMER3 package <a href="http://hmmer.org/">http://hmmer.org/</a>; Eddy, 2011). This program is used to run similarity searches of profile HMMs against metagenomic datasets. The program must be located in a directory listed in the PATH of the operating system.

tblastn - (BLAST package - <a href="http://blast.ncbi/nlm/nih/gov">http://blast.ncbi/nlm/nih/gov</a>); Altschul *et al.*,
 1997). The program is used to detect the coding regions corresponding to the positive HMMs in the assembled sequences.

#### 3.3 How to cite

• If you use this program for your publication, please cite e-Finder it as: e-Finder program (developed by Liliane S. Oliveira and Arthur Gruber, University of São Paulo, Brazil, unpublished).

#### 4 Understanding e-Finder's workflow

#### 4.1 Input data

e-Finder uses two types of input data: a set of profile HMMs and one or more FASTA files containing assembled sequences (Figure 1A). In addition, a tabular file containing a list of dataset identifiers and the respective organism names can optionally be used. This information, if provided, is used to generate a final CSV report.

#### 4.1.1 profile HMMs

Profile HMMs are used to identify the corresponding genes in the assembled sequences. Single or multiple protein markers can be used to detect the elements, each one represented by one or more profile HMMs. For instance, casposons are mobile genetic elements composed of a variable number of genes (Krupovic *et al.*, 2014) but always presenting *cas1* (Cas1 endonuclease) and *pol8* (PolB-like RNA polymerase) genes. Thus, one can use profile HMMs constructed from proteins coded by these genes to fish casposon elements in bacterial and archaeal genomes. Also, if multiple profile HMMs are available for each protein, they can be used in combination. This strategy may increase the sensitivity of detection for more remote elements, as different protein domains may present variable rates of divergence and very often such rates are not known *a priori*. e-Finder can identify cutoff score tags inserted in the profile HMMs and use their values as filters in the similarity search step (see below for more details).

# 4.1.2 Assembled sequence datasets

As input, e-Finder can use FASTA files containing sequences derived from either single or multiple genomes. Alternatively, e-Finder can also use a directory containing multiple subdirectories, each one specific for a given genome, which in turn contains the corresponding FASTA files. This input option is particularly convenient when using genomic data from PATRIC (<a href="https://www.patricbrc.org/">https://www.patricbrc.org/</a> — Wattam *et al.*, 2017), the Bacterial Bioinformatics Resource Center, a comprehensive repository of assembled genomes that employs this data storage structure. Other genomic and metagenomic datasets can also be used, provided that they are previously assembled.

# 4.2 Detection of candidate sequences

e-Finder uses profile HMMs as a query in similarity searches against assembled sequences (Figure 1B). Since hmmsearch does not automatically translate nucleotide sequences (like tblastn does), transeq program is invoked to translate all sequences into the six possible reading frames. Once this step is performed, e-Finder then executes hmmsearch to run the searches with its default parameter -S 10, which corresponds to an e-value of 10, a low-stringency condition. While this choice ensures a high sensitivity of detection, the user can still define an alternative score or e-value to be used as threshold for the sequence selection step. Different secondary thresholds can be used in multiple e-Finder executions to arbitrarily increase specificity, with no need to rerun the hmmsearch search. In fact, if the e-Finder detects a previous similarity search file, it will only analyze that file to retrieve the results that fall within the newly established threshold. This feature allows to skip the slow similarity search execution, saving precious time. Alternatively, threshold values can also be specifically hardcoded in each profile HMMs as cutoff score tags (e.g. CUTOFF SCORE 45.3). e-Finder automatically identifies this tag in all models, running hmmsearch with parameter -T (45.3 in this example), thus allowing custom cutoffs to be used for each corresponding model. The CUTOFF SCORE is a proprietary tag that has no effect on hmmsearch program itself but allows e-Finder to invoke it with the most appropriate threshold for each profile HMM. All models from the Viral MinionDB (<u>http://www.bioinfovir.icb.usp.br/miniondb</u>) are provided with the appropriate cutoff scores. E-Finder executes hmmsearch with the -T option using the respective cutoff value.

# 4.3 Selection of positive sequences

Once the positive sequences have been detected, e-Finder checks whether the user-defined criteria for the synthetic context have been met (Figure 1C). Thus, each sequence must contain a minimum number of markers and the appropriate intergenic distances, including or not the possibility of overlapping genes. Beside gene composition, the user can also specify whether or not a strict gene order must be found. Regions containing the multiple gene markers are extracted together with size-defined 5'and 3' flanking regions and submitted to size filtering. By using a selected genetic code, ORFs coding for the protein sequences detected by the models are then identified, extracted and conceptually translated into full-length protein sequences.

#### 4.4 Sequence extraction and report generation

In the final phase (Figure 1D), e-Finder stores the nucleotide and protein sequences of each selected region. Also, the program generates and stores a CSV spreadsheet file listing all elements found in each assembled sequence and their corresponding markers, coordinates and intergenic distances, among other features.

#### 5 Understanding e-Finder parameters

# 5.1 Mandatory parameters:

e-Finder has two mandatory parameters that must be specified by the user at the command line: the dataset directory (or file) and the profile HMM file.

-dd|dataset\_directory <directory name> - Directory that contains
multiple subdirectories, each one with a FASTA file of the nucleotide
sequence(s) that will be used by e-Finder. This is the typical data structure
obtained from the PATRIC repository (<a href="https://www.patricbrc.org/">https://www.patricbrc.org/</a>).

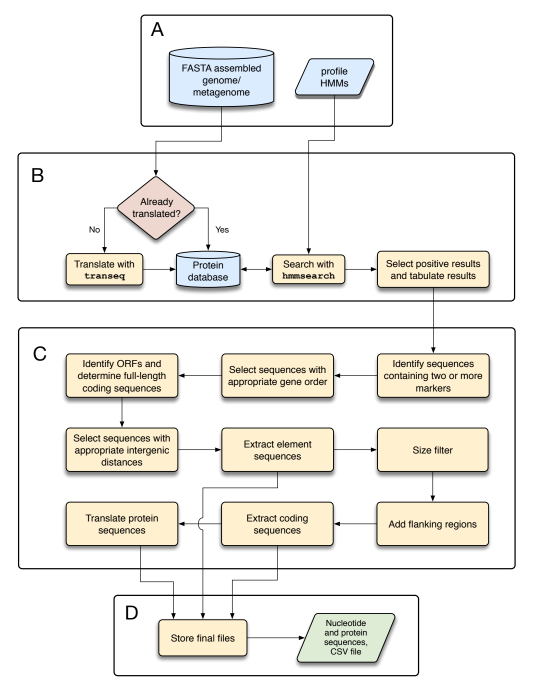


Figure 1 – Workflow of e-Finder. Two types of input data are used by e-Finder: a set of profile HMMs and one or more FASTA files containing assembled sequences (A). In the similarity-based detection phase (B), e-Finder invokes transeq to translate the assembled sequences into the six possible open reading frames. Profile HMMs are then used as queries in similarity searches against the translated dataset using hmmsearch. e-Finder selects all sequences containing positive results, according to user-defined threshold values. In the next phase (C), e-Finder determines whether user-specified markers are present in the appropriate syntenic context in each of selected sequences. Regions containing the defined markers are extracted together with size-defined flanking regions and submitted to size filtering. ORFs coding for sequences detected by the models are then identified, extracted and translated into full-length protein sequences. In the final phase (D), e-Finder stores all element sequences and a CSV spreadsheet file listing the elements found in each assembled sequence and their corresponding markers, among other features.

- -df|dataset\_file <file name> Dataset file in FASTA format. This file
   contains the nucleotide sequence(s) that is used by e-Finder.
- -i|input\_files <file name> Single or multiple files containing one or more profile HMMs each. For each marker, a distinct profile HMM file must be provided with a prefix identifier, followed by an underscore character and additional letters, or directly by an extension (e.g. prot1.hmm, prot2.hmm, etc.). If using multiple files, their names must be separated by commas (e.g. -i file1.hmm, file2.hmm, file3.hmm). Marker prefixes can be used to define syntenic context (see parameter -synteny).

#### 5.2 Optional parameters:

- -ce | circular <yes | no> Assume that the element is originally derived from a circular element (e.g. a circular phage genome). Default: no.
- -conf <configuration file> use a configuration file that lists all parameters for execution, overriding any parameter of the command line. An example of a configuration file follows below:

```
dd=test_directory
i=gene1.hmm,gene2.hmm
o=output_dir
ex=fna
id=40000
fs=10000
patric_list=genome_list.csv
ol=50
qc=0
```

- -cpu|cpu\_threads <integer> Number of threads to be used by hmmsearch. If not specified, e\_Finder determines the number of threads available in the multiprocessor server and uses by default half of this value.
- -e|e-value or -s|score <decimal> E-value (-e) or score (-s) threshold value. Report hmmsearch hits that present values equal to or lower than the E-value or equal to or larger than the score. Only one of these parameters and the respective value shall be provided (Default = -e 10).

- -ed|element\_distance <integer> Minimum distance between elements.
  Default = 5000.
- -ex|extension <fna|faa|fasta> Extension for input files. When using PATRIC database data, we suggest using fna, since this is the extension of the contig nucleotide sequence files provided in this resource. Default: fna.
- -fs|flanking\_size <integer> Size (in bp) of the 5' and 3' flanking regions that will be excised together with the multigene element (default = 5000). If a 0 (zero) value is used, e-Finder extracts the sequence comprised between the start codon of the first gene and the stop codon of the last gene.
- -gc <integer> Genetic code to define start codons and perform conceptual translation of the genes. e-Finder uses numbering codes defined on the NCBI Genetic Codes page and implemented on transeq (EMBOSS package): 0 (Standard); 1 (Standard with alternative initiation codons); 2 (Vertebrate Mitochondrial); 3 (Yeast Mitochondrial); 4 (Mold, Protozoan, Coelenterate Mitochondrial and Mycoplasma/Spiroplasma); 5 (Invertebrate Mitochondrial); 6 (Ciliate Macronuclear and Dasycladacean); 9 (Echinoderm Mitochondrial); 10 (Euplotid Nuclear); 11 (Bacterial); 12 (Alternative Yeast Nuclear); 13 (Ascidian Mitochondrial); 14 (Flatworm Mitochondrial); 15 (Blepharisma Macronuclear); 16 (Chlorophycean Mitochondrial); 21 (Trematode Mitochondrial); 22 (Scenedesmus obliquus); 23 (Thraustochytrium Mitochondrial). Default: 0.
- -h|help Display help screen.
- -ic|ignore\_cutoff Ignore cutoff scores in the profile HMMs and use a custom value defined by parameters -e or -s for all input models (default = yes). If -r no is used, e-Finder will use the cutoff scores specified in the respective CUTOFF SCORE tag of each profile HMM. For models not containing cutoff values, e-Finder will use the cutoff value specified by the parameter -e or -s. If none of these parameters are specified, the program will then use hmmsearch's default cutoff value (-E 10).
- -id|intergenic\_dist <integer> Maximum distance (in bp) between intergenic regions (default = 5000). This value is adopted for all intergenic

distances between any pair of genes. If the -synteny parameter is used, its values take precedence over those set for -id.

- -mg|min\_gene <integer> Minimum number of genes (default = 2). This parameter specifies the minimum number of genes that must be found for a sequence to be considered positive. For instance, if profile HMMs from four different proteins are used and the user specifies -mg 2, any sequence containing at least two of the four markers will be considered for downstream analysis of the remaining criteria.
- Output directory name (default: output\_dir). This is the directory where eFinder will store all output directories/file. All similarity search results are
  stored in the all\_results subdirectory. If the user specifies an output
  directory that already exists, e-Finder inspects the all\_results
  subdirectory and uses the hmmsearch results from the previous run. This
  feature saves processing time since it skips the relatively slow similarity search
  step. For each run, e-Finder creates a run\_# (e.g. run\_1, run\_2,
  run\_3, etc.) subdirectory where all output files are stored.
- -olloverlap <integer> Maximum allowed overlap distance (in bp) between open reading frames in the same coding strand. If a 0 (zero) value is used (default), no overlap is allowed.
- -pl|patric\_list <file name> Input file in PATRIC-like
   (https://www.patricbrc.org) tabular format. This two-column file lists accession
   codes and organism names, respectively, and provides information for e Finder to generate a final CSV file reporting all found multigene regions
   associated with the respective organism names. Accepted format:

```
genome_id genome_name
1309411.5 'Deinococcus soli'
1123738.3 'Echinacea purpurea'
551115.6 'Nostoc azollae'
1856298.3 'Osedax'
```

-sf|size\_filter <integer> - Minimum size (bp) of the excised element, not including user-defined flanking regions (parameter -fs). Default: 1000.

-synteny <string> - Define gene order and maximum allowed distance (kb) between genes. Each marker is defined by a letter and the distances can be specified by decimals. Intergenic distances defined for parameter -synteny take precedence over those set for -id. If parameter -synteny is not specified, e-Finder will accept any sequences presenting the minimum number of genes (specified in parameter -mg), with any intergenic distances (up the maximum value defined by parameter -id).

If the region is assumed to be exogenously derived from a circular element (parameter -ce yes), then an additional distance between the last and the first marker listed must be given. In the example below, value 2.5 refers to the distance between markers e and a.

```
• Linear element: a, 2000, b, 1500, c, 3000, d, 3500, e
```

• Circular element: a,2000,b,1500,c,3000,d,3500,e,2500

-v | version - display program's version.

#### 6 Running e-Finder

To execute e-Finder, a configuration file can be used. For example:

```
e-finder.pl -conf configuration.cnf
```

An example of a configuration file follows below:

```
dd=/home/username/genomes
i=/home/username/hmm/gene1.hmm,/home/username/hmm/gene2.hmm,/home/user
name/hmm/gene3.hmm
o=results_dir
ic=yes
ex=fna
id=3000
fs=8000
patric_list=genome_list.csv
ol=50
```

• Alternatively, instead of running e-Finder with a configuration file, the program can be executed with all parameters specified in the command:

```
e-finder.pl -df <file> -i <file> -s|-e <decimal> <optional parameters>
```

• A command for the parameters specified in the configuration file above is:

```
e-finder.pl -dd /home/argruber/genomes -i /home/username/hmm/gene1.hmm,/home/username/hmm/gene2.hmm,/home/userna me/hmm/gene3.hmm -ic yes -ex fna -id 3000 -fs 8000 -patric_list genome list.csv -gc 11 -ol 50 -sf 1000 -o results dir
```

# 7 Inspecting e-Finder output files

Once e-Finder finishes the processing, an output directory is created with several files and subdirectories:

- logfile.txt: this file that reports all steps of the execution.
- all\_results: all similarity search results are stored in this directory. Since all subsequent results use the hmmsearch results as a starting point, it is possible to rerun e-Finder using different parameters (minimum number of genes, intergenic distances, etc.). In this case, the user just needs to execute e-Finder in a directory where all\_results folder already exists. If the user wants to use other profile HMMs, then a new directory must be created to execute a clean run.
- selected: this is the directory where e-Finder stores the results of all selected (positive) datasets, each one in a specific subdirectory. If a PATRIC-like data structure is used, the subdirectories are named with the corresponding identifier. Each subdirectory contains some files, including nucleotide sequences of the positive contig and ORFs of the proteins that were positive to the profile HMMs used in the search. Sequences of the corresponding proteins are also stored. In the root of the selected directory, e-Finder stores elements.fasta, a FASTA file that contains the nucleotide sequences of all elements found in the positive datasets. Additional files include protein sequences conceptually translated from the elements. These are the proteins detected by the profile HMMs.

- discarded: this is a directory containing the results of all positive datasets that did
  not fulfill all criteria to be selected. For instance, sequences that were positive for only
  marker, or whose intergenic length was out of the maximum defined distance.
- final\_report.csv: this is a tab-delimited result file that summarizes the results for all elements found. Stored information includes organism name, contig size, element coordinates in the contigs, gene coordinates and orientation, intergenic distances, etc.

#### 8 References

- Altschul, S.F., Madden, T.L., Schäffer, A.A., Zhang, J., Zhang, Z., Miller, W. & Lipman, D.J. (1997). "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs." Nucleic Acids Res. 25:3389-3402.
- Eddy SR. (2011). Accelerated Profile HMM Searches. PLoS Comput Biol. 7(10):e1002195.
- Krupovic M, Makarova KS, Forterre P, Prangishvili D, Koonin EV. (2014). Casposons: a new superfamily of self-synthesizing DNA transposons at the origin of prokaryotic CRISPR-Cas immunity. BMC Biol. 12:36.
- Wattam AR, Davis JJ, Assaf R, Boisvert S, Brettin T, Bun C, Conrad N, Dietrich EM, Disz T, Gabbard JL, Gerdes S, Henry CS, Kenyon RW, Machi D, Mao C, Nordberg EK, Olsen GJ, Murphy-Olson DE, Olson R, Overbeek R, Parrello B, Pusch GD, Shukla M, Vonstein V, Warren A, Xia F, Yoo H, Stevens RL. (2017). Improvements to PATRIC, the all-bacterial Bioinformatics Database and Analysis Resource Center. Nucleic Acids Res. 45(D1): D535-D542.