polyDFE User Manual, BETA Version

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polyDFE infers the distribution of fitness effects (DFE) of new mutations from polymorphism, and, if available, can also incorporate divergence data obtained from an outgroup. The polymorphism data is provided as an unfolded site frequency spectrum (SFS). Note that, in order to obtain an unfolded SFS, at least one outgroup is needed. However, including the divergence data when inferring the DFE can lead to bias in the estimated parameters (see ? for further discussion).

Once the DFE is obtained, polyDFE also calculates α , the rate of adaptive molecular evolution, commonly defined as the proportion of fixed adaptive mutations among all non-synonymous substitutions. This code implements the program described in ?.

Contents

1 Authors

The polyDFE has been developed and implemented by Paula Tataru and Marco A.P. Franco.

2 License

polyDFE is open source, distributed under the GNU General Public License, version 3. See LICENSE.txt for details.

3 Requirements

polyDFE is implemented in C and uses the GSL library. This should be downloaded from https://www.gnu.org/software/gsl/ and compiled accordingly. polyDFE has been tested using GSL-1.16.

4 Installation

The simplest way to compile polyDFE is

- 1. cd to the directory containing the makefile and type make all to compile the code.
- 2. You can remove the program binaries and object files from the source code directory by typing make clean.

If the GSL library is not installed in the default directory, than the installation path needs to be specified in the makefile by updating USR_INC and USR_LIB and uncommenting the lines at the top of the makefile.

5 Running polyDFE

Running polyDFE with the argument -h will print out the usage (required and optional arguments)

polyDFE runs in two modes: -s, used for simulating data under the hierarchical probabilistic model, and -o, used for estimating the DFE and α . If neither -s or -o are given, then -o bfgs is used by default (see argument -o on page ?? for details). If both -s and -o are given, polyDFE will terminate with an error. Optional arguments are given between [].

5.1 Shared arguments

A few arguments are shared between the two modes of polyDFE.

Regardless of the mode polyDFE is ran in, the following argument is required

• -d data_file: path to the data file.

data_file is where polyDFE will write the simulated data, if -s is used, or will read the input data, if -o is used.

The file can contain any number of empty lines and comment lines that start with #. Excluding these, the structure of data_file is

$$m_{\text{neut}} \quad m_{\text{sel}} \quad n$$

$$m_{\text{neut}} \text{ lines} \begin{cases} p_{\text{neut}}(1) & p_{\text{neut}}(2) & \dots & p_{\text{neut}}(n-1) & l_{\text{neut}} & d_{\text{neut}} & l_{\text{neut}}^d \\ \vdots & \vdots & \ddots & \vdots & \vdots & \vdots \\ m_{\text{sel}} \text{ lines} \end{cases} \begin{cases} p_{\text{sel}}(1) & p_{\text{sel}}(2) & \dots & p_{\text{sel}}(n-1) & l_{\text{sel}} & d_{\text{sel}} & l_{\text{sel}}^d \\ \vdots & \vdots & \ddots & \vdots & \vdots & \vdots \end{cases}$$

where

- \circ m_{neut} and m_{sel} are the number of fragments that are assumed to be evolving neutrally or under selection, respectively;
- \circ *n* is the ingroup sample size (number of sequences);
- o $p_z(i)$ is the i^{th} entry in the SFS within the fragment, for sites that are assumed to be evolving neutrally, z = neut, or under selection, z = sel;
- \circ l_z is the total number of sites within the fragment used for the SFS (number of successfully sequenced sites within the ingroup) that are assumed to be evolving neutrally, z = neut, or under selection, z = sel;
- o d_z is the total number of fixed mutations in the ingroup relative to the outgroup (divergence counts) within the fragment, that are assumed to be evolving neutrally, z = neut, or under selection, z = sel;
- o l_z^d is the total number of sites within the fragment used for the divergence counts (number of successfully sequenced sites within the ingroup and outgroup) that are assumed to be evolving neutrally, z = neut, or under selection, z = sel; l_z and l_z^d , for a given fragment, could be different if certain sites are successfully sequenced in the ingroup, but not in the outgroup.

The divergence data (last two columns, d_z and l_z^d) can be absent from the file. If it is absent, it should be absent for all fragments.

polyDFE uses data divided into fragments in order to model variability in mutation rates, as described in ?. If only one neutral and one selected fragments are provided, than variability in mutation rates is not modeled.

The provided files input/example_1, input/example_2, input/example_3, input/example_4 and input/example_5 are examples of data_file.

Two optional arguments are shared between the the modes of polyDFE

- -m model(A, B, C, D) [K]: assumed DFE model. polyDFE assumes that the DFE ϕ takes one of four specific functional forms, encoded A, B, C, and D, described below
 - A: the DFE is given by a reflected displaced Gamma distribution, parameterized by \bar{S} , b and S_{\max} , with density

$$\phi_A(S;\bar{S},b,S_{\max}) = \left\{ \begin{array}{ll} f_{\Gamma}(S_{\max}-S;S_{\max}-\bar{S},b) & \text{if } S \leq S_{\max} \\ 0 & \text{otherwise} \end{array} \right.$$

where \bar{S} is the mean of the DFE, b is the shape of the Gamma distribution, S_{\max} is the maximum value that S can take, and $f_{\Gamma}(x; m, b)$ is the density of the Gamma distribution with mean m and shape b.

 \circ B: the DFE is given by a mixture of a Gamma and discrete distributions, parameterized by S_d , b, p_b and S_b , where

$$\phi_B(S; S_d, b, p_b, S_b) = \begin{cases} (1 - p_b) f_{\Gamma}(-S; -S_d, b) & \text{if } S \leq 0 \\ p_b & \text{if } S = S_b \\ 0 & \text{otherwise} \end{cases}$$

where S_d is the mean of the DFE for $S \leq 0$, b is the shape of the Gamma distribution, p_b is the probability that S > 0, and S_b is the shared selection coefficient of all positively selected mutations, and $f_{\Gamma}(x; m, b)$ is the density of the Gamma distribution with mean m and shape b.

 \circ C: the DFE is given by a mixture of a Gamma and Exponential distributions, parameterized by S_d , b, p_b and S_b , where

$$\phi_C(S; S_d, b, p_b, S_b) = \begin{cases} (1 - p_b) f_{\Gamma}(-S; -S_d, b) & \text{if } S \le 0 \\ p_b f_e(S; S_b) & \text{if } S > 0 \end{cases}$$

where S_d is the mean of the DFE for $S \leq 0$, b is the shape of the Gamma distribution, p_b is the probability that S > 0, S_b is the mean of the DFE for S > 0, and $f_{\Gamma}(x; m, b)$ is the density of the Gamma distribution with mean m and shape b, while $f_e(x; m)$ is the density of the Exponential distribution with mean m.

 \circ D: the DFE is given as a discrete DFE, where the selection coefficients can take one of S_i distinct values, $1 \le i \le K$, where each value S_i has probability p_i , with

$$\sum_{i=0}^{K} p_i = 1$$

The value of K needs to be provided in the command line for model D.

If -m is not given, model C is used by default.

• -t: the total running time of polyDFE will be measured and printed on the standard output at the end.

-i is also common to the two modes. However, -i is required when simulating data (mode -s), while when inferring the parameters (mode -o), it is an optional argument for DFE models A, B and C. Note though that if using model D, then -i has to be given, as it contains the information about the selection coefficients S_i .

• -i init_file ID: path to a file containing the values of the DFE and other parameters, and id.
init_file can contain any number of empty lines and comment lines that start with #. Excluding these,
the structure of each line is

id
$$[0|1] \epsilon_{\text{an}} \quad [0|1] \epsilon_{\text{cont}} \quad [0|1] \lambda \quad [0|1] \bar{\theta} \quad [0|1] a \quad < \text{DFE parameters with flags} > \quad [0|1] r_2 r_3 \dots r_n$$

Each line starts with a positive numerical id. This is used such that multiple configurations of the parameters can be stored in the same file. When running polyDFE, the desired line for initializing the parameters is specified through the id by setting ID to the right value.

The id is followed by the values of the parameters, in the order $\epsilon_{\rm an}$, $\epsilon_{\rm cont}$, λ , $\bar{\theta}$, a, the DFE parameters (as detailed for argument -m above), and the r_i , $2 \le i \le n$, parameters, where n is the ingroup sample size.

All parameters are preceded by a flag, which is either 0 or 1. In the \neg s mode, the flag is not used, but this becomes important for the \neg o mode, where the flag indicates if a parameter should be estimated (flag 0) or should be kept fixed to the value provided in init_file (flag 1). All r_i parameters share one flag. The r_i parameters can also be provided for i > n, but those values will be ignored. This is useful if the same line in init_file is used for multiple data sets that have different sample sizes.

For DFE model D, only the probabilities p_i are estimated, while the selection coefficients S_i are fixed to the given values.

For details of the meaning of each parameter, please see? The provided files input/init_model_A.txt, input/init_model_BandC.txt and input/init_model_D.txt give examples of the structure of the init_file for the different DFE models.

Note that the a parameter is directly linked to variability of mutation rates. If it is wished for mutation rates to be assumed constant, than a can be fixed to -1. If both m_{neut} and m_{sel} provided in the input

datafile (through argument -d) are equal to 1 (i.e. the data is not divided into fragments), then the mutation variability is not estimated by default. When mutation variability is allowed, than the observed data (counts) are assumed to follow a negative binomial distribution, otherwise they are assumed to follow a Poisson distribution. See ? for more details on mutation variability.

The parameter $\epsilon_{\rm cont}$ allows for modeling the SFS for the sites that are assumed to be evolving neutrally as a mixture of neutral and selected sites. Note that this parameter is not identifiable and it is therefore recommended that it's flag is always 1. However, this parameter can be used to test the robustness of the method at the assumption of neutrality: one can check how much the estimated DFE and α would change if $\epsilon_{\rm cont}$ was set to some value different than 0.

The provided files input/init_model_A.txt, input/init_model_BandC.txt and input/init_model_D.txt are examples of init_file.

5.2 Simulating data

The simulation mode (-s) also requires the following argument

• -s m_neut L_neut m_sel L_sel n: simulate data.

When using -s, simulated data will be printed in data_file with $m_{\text{neut}} = \texttt{m_neut}$, $m_{\text{sel}} = \texttt{m_sel}$, $l_{\text{neut}} = l_{\text{neut}}^d = \texttt{L_neut}$ and $l_{\text{sel}} = l_{\text{sel}}^d = \texttt{L_sel}$, for all simulated fragments. For details on m_z and l_z , with $z \in \{\text{neut}, \text{sel}\}$, see argument -d on page ??.

5.3 Estimating DFE and α

The estimation (optimization) mode (-o) allows for the estimation of DFE and α . For this, next to the arguments described above, it allows for a series of optional arguments, which control the optimization, both which and how the parameters are estimated, but also when a set of parameters found is considered to be optimal.

- -o (bfgs, conj_pr, conj_fr): optimization method to use for estimating the parameters.
 - polyDFE uses GSL to run a local optimization algorithm and find estimates of the parameters. GSL implements multiple optimization algorithms. Generally, optimization is more efficient if it also uses the derivative of the function to be optimized. Therefore, in polyDE we have chosen to use three of the algorithms offered by GSL, which all rely on the function (in this case, the data likelihood) derivatives. These are:
 - o bfgs: the Broyden-Fletcher-Goldfarb-Shanno (BFGS) algorithm.
 - o conj_pr: the Polak-Ribiere conjugate gradient algorithm;
 - conj_fr: the Fletcher-Reeves conjugate gradient algorithm.
 For details on the different optimization algorithms, please see the GSL manual. When -o is not given, -o BFGS is used by default. polyDFE has been tested using the BFGS algorithm.
- -r range_file ID: path to a file containing the allowed range of all parameters, and id. range_file can contain any number of empty lines and comment lines that start with #. Excluding these, the structure of each line is
- id $k = \epsilon_{\text{an}}^{\min} \epsilon_{\text{an}}^{\max} = \epsilon_{\text{cont}}^{\min} \epsilon_{\text{cont}}^{\max} = \lambda^{\min} \lambda^{\max} = \bar{\theta}^{\min} \bar{\theta}^{\max} = a^{\min} a^{\max} = < \text{ranges for DFE parameters} > -r^{\min} r^{\max}$

Each line starts with a positive numerical id. This is used such that multiple configurations of the parameters can be stored in the same file. When running polyDFE, the desired line for the range of the parameters is specified through the id by setting ID to the right value.

The id is followed by k, which controls the transformation of the parameters (see below) and the range within each parameter should be estimated. Some parameters are intrinsically constrained (for example, p_b for DFE models B and C is, by construction, constrained to be between 0 and 1), while many parameters have to be positive (or negative, for example, S_d for DFE models B and C). In order to treat all parameters in the same way and to also allow the user to narrow the range in which the optimum parameters are searched, all parameters are then constrained to the range provided by the user through

the range_file. However, the optimization algorithms polyDFE uses (see argument -o on page ?? for details) are unconstrained, in that the parameters can take any value in the interval $(-\infty, +\infty)$. In order to change the original constrained optimization problem to an unconstrained one, polyDFE transforms the parameters from the range provided by the user to the interval $(-\infty, +\infty)$. For this, a generalized logistic function is used. This function is parameterized by a growth rate (or steepness of curve) k, which controls the spread of the transformation: the smaller the k, the more quicker the transformed values move from $-\infty$ to $+\infty$. polyDFE has been tested with a value of k=0.01. If the optimization algorithm cannot find a good solution (i.e. the gradient is small, see argument -p on page ?? for details), the k parameter can be changed in the hope of better performance.

Note that, in order for the optimization algorithm to successfully find the optimum parameters (the parameters that have the height likelihood), the ranges should be large enough to ensure that they cover the optimum values.

If initial values of the parameters provided through the -i argument or as calculated when the argument -e is provided are not within the ranges, the program will automatically update the ranges.

The provided files input/range_model_A.txt, input/range_model_BandC.txt and input/range_model_D.txt are examples of range_file.

• -e: automatically estimate the initial value of the parameters.

The optimization algorithms require an initial value for the parameters to be optimized. This can either be provided using $-\mathbf{i}$ if the user has a good idea for such values, or can, by using the argument $-\mathbf{e}$. If $-\mathbf{e}$ is used without $-\mathbf{i}$, then all the parameters (expect for $\epsilon_{\rm cont}$, which, by default, is not estimated and set to 0) will be automatically initialized and then optimized. However, if $-\mathbf{i}$ and $-\mathbf{e}$ are used together, the initial value of the parameters is estimated automatically only for those parameters that are flagged with 0 and are consequently optimized.

The parameters $\bar{\theta}$, a, λ and r_i can be estimated deterministically. The rest of the parameters are estimated using a grid search approach. However, for the DFE model D, the probabilities p_i are set to be uniform when $-\mathbf{e}$ is used.

If -i is not used, -e is used by default.

• -j: use jointly both the provided initial value and automatically estimated value of parameters.

When the argument -j is used together with -i, then the optimization algorithm is ran twice, once where the initial value of the parameters is the one given in the init_file, and a second time where the automatic estimation from -e is used. At the end, the parameters that have the best likelihood, found from one of the two runs, will be reported.

• -w: do not use divergence data.

When the argument -w is used, then the DFE and other parameters are estimated without using divergence data, as described in (?). If the divergence data is absent from the data_file, -w will be used by default.

• -g grouping_file ID: path to a file containing grouping information for the r_i parameters, and id. grouping_file can contain any number of empty lines and comment lines that start with #. Excluding these, the structure of each line is

id
$$G$$
 $i_1 i_2 \dots i_G$

Each line starts with a positive numerical id. This is used such that multiple configurations of the grouping can be stored in the same file. When running polyDFE, the desired line for the range of the parameters is specified through the id by setting ID to the right value.

The grouping_file is used to provide information on grouping the r_i parameters. By default, each entry $1 \le i < n$ in the SFS, and the divergence count, have one value of r_i associated to it. If n is very large, this leads to a lot of parameters that need to be estimated. One could expect that the distortion that is incurred on the counts that are modeled using the r_i parameters could be similar for neighboring values (for example, i-1, i and i+1). Then only one r value can be estimated for those counts that share a similar distortion.

The grouping given in $grouping_file$ will result in r values corresponding to the SFS entries as follows (where one interval represents one range for the SFS entries):

$$\begin{bmatrix} r_1 & r_2 & r_3 & \dots & r_{G+1} \\ [1] & [2, i_1] & [i_1 + 1, i_2] & \dots & [i_G, n'] \end{bmatrix}$$

where n' is either n, if divergence data is used in the estimation (see argument $-\mathbf{w}$ on page ?? for details), or n-1 otherwise. For identifiability reasons, r_1 is always set to 1 and not estimated.

The provided file input/params.grouping is an example of a grouping_file.

• -p optim_file ID: path to a file containing parameters that control the optimization algorithm, and id.

optim_file can contain any number of empty lines and comment lines that start with #. Excluding these, the structure of each line is

id $\epsilon_{\rm abs}$ step size tol max iter

Each line starts with a positive numerical id. This is used such that multiple configurations for the optimization algorithms can be stored in the same file. When running polyDFE, the desired line for the range of the parameters is specified through the id by setting ID to the right value.

The optim_file is used to control the behavior of the optimization algorithms from GSL.

- \circ $\epsilon_{\rm abs}$: this the absolute tolerance value that the gradient of the function to be optimized (here, the likelihood) is tested against. When the gradient is smaller than this values, than GSL considers that it has successfully found an optimum. If this value is too large, it will lead to an early termination of the algorithm. If the value is too small, it might lead to a longer running time of the algorithm without, potentially, much improvement on the function. However, it is always best to have an absolute tolerance value that is too small than too large. If a set of values found is a true optimum, than the gradient should be 0. Default value: 0.00001.
- step size: the GSL optimization algorithms use a line search and the step size determines the size of the first step performed in the search. Default value: 2.
- tol: specifies the accuracy of the line search, with the precise meaning depending on the algorithm used. Default value: 0.1.
- max iter: the GSL optimization algorithms are iterative. If the number of performed iterations reaches max iter, than the algorithm is stopped, regardless of the gradient value. Default value: 1500.

For more details on the above, please consult the GSL manual.

The provided file input/params.optim is an example of an optim_file.

• -b [basinhop_file ID]: path to a file containing parameters that control the basin-hopping algorithm, and id.

basinhop_file can contain any number of empty lines and comment lines that start with #. Excluding these, the structure of each line is

id max same max iter temp step accept rate interval factor

Each line starts with a positive numerical id. This is used such that multiple configurations for the basin-hopping algorithm can be stored in the same file. When running polyDFE, the desired line for the range of the parameters is specified through the id by setting ID to the right value.

By default, polyDFE runs the optimization algorithm once (or twice, see argument -j on page ?? for details), and reports the results. However, the optimization algorithms are local, in that they only find an optimum that is in the neighborhood of the initial values of the parameters. To look more thoroughly for a global optimum, polyDFE allows the user to run the basin-hopping algorithm (???). polyDFE contains a reimplementation of the basin-hopping algorithm found in the Python library, scipy (??).

Basin-hopping attempts to find the global optimum by iterating the following steps

- 1. perturbating randomly the current value of the parameters;
- 2. running the local optimization algorithm (as chosen through -o, see page ?? for details) using the new values as initial values;
- 3. accepting or rejecting the found optimum based on the likelihood.

Basin-hopping has been shown to be very efficient for a wide variety of problems in physics and chemistry. Unfortunately, there is no way to determine if the true global optimum has actually been found, and therefore it is left to the user to ensure that. The algorithm is controlled through

- max same: If after max same iterations, the basin-hopping algorithm does not find an improved set of parameters, than the algorithm stops. Default value: 50.
- \circ max iter: The basin-hopping algorithm is ran for a maximum of max iter iterations. Default value: 500.
- temp: The temp is the temperature used in the metropolis criterion when accepting or rejecting the new values. For best results, the temperature should be comparable to the typical difference in likelihood between local optima. Default value: 1.
- o step: The step controls how far away from the current value the perturbated value is. This is crucial for the algorithm's performance. Ideally, it should be comparable to the typical separation between local optima of the likelihood. The algorithm implemented in polyDFE will automatically adjust the step, but it make take many iterations to find an optimal value. Default value: 50.
- accept rate: The target accept rate (percentage of new values that are accepted in step 3) for when adjusting the step. Default value: 0.5.
- o interval: Every interval iterations, the basin-hopping algorithm adjusts the step. Default value: 10.
- o factor: When the step is adjusted, it is done with this factor. Default value: 0.9.

The argument -b can be ran without basinhop_file ID. If only -b is given, then polyDFE runs basinhopping with the default parameter values, given above.

The provided file input/params.basinhop is an example of a basinhop_file.

• -1 min: limit running time to min.

To control the running time of polyDFE, one run of the optimization algorithm is stopped if it has used more than min. By default, min is set to 60. Note that if min is set too low, there is a high chance that the optimization algorithm will terminate without finding good optimal parameters.

• -v verbose(0, 1, frequency): verbose frequency.

The argument -v controls how often information about the status of the optimization algorithm is printed on the screen. By default, verbose is set to 0, and only the initial and optimum values of the parameters are printed. Otherwise, every verbose iterations of the optimization algorithm, information is printed about the current value of the parameters, the corresponding likelihood, gradient and status of the optimization. Apart from the possible status values that GSL uses, polyDFE has the additional values:

- -3: local optimization is stuck in the same parameter values (it cannot further improve on the current values).
- \circ -4: restarting the local optimization lead to the same parameter values.
- -5: the local optimization is being restarted; sometimes, when the local optimization cannot further improve on the current values, restarting the optimization can allow it to proceed further. If the local optimization is stuck in the same values, restarting is done at most 3 consecutive times.
- o -6: the local optimization used at least min minutes and will be stopped.
- -7: the likelihood evaluation returned NAN and the local optimization is therefore stopped. This can happen for certain values of the parameters, where the numerical integration over the DFE fails.
- -8: a maximum number of likelihood evaluations has been reached, and the local optimization has
 been forcefully stopped. If this status appears, is indicates an issue with the local optimization
 procedure and the obtained results are not reliable.

6 Post-processing using R

polyDFE is accompanied by a series for R functions that allow

- parsing of the output of polyDFE for easy manipulation in R
- creating init_file containing the best found parameters
- creating grouping for the r parameters
- calculating α using alternative definitions
- performing model testing

6.1 Parsing the output of polyDFE

The function

```
parseOutput(filename, init = FALSE)
```

allows the user to parse the output of polyDFE from multiple runs stored in filename. The function returns a list with entries for each separate run of polyDFE found, containing

- the DFE model used;
- the best likelihood found and the corresponding gradient;
- the values of all parameters (including those that have been fixed using the flag 1 in init_file);
- which parameters have been estimated;
- the value of n (the sample size of the data);
- expected SFS and, if relevant, divergence counts and missattributed polymorphism;
- estimates of α .

If init = TRUE, then the function returns the values for the local optimization where the parameter values found in init_file where used as initial values for the parameters during the optimization (see argument -e on page ?? for details), regardless if this returned the best likelihood or not.

6.2 Creating init_file

The function

```
createInitLines(estimates, outputfile, startingID, fix, groupingDiff = 0)
```

creates an init_file with the given values of the parameters.

estimates can either be a list returned by parseOutput, or a file name, as for parseOutput. For each run of polyDFE found, a new line is appended to <outputfile>_init containing the values of the best parameters found in the format described previously (see argument -i on page ?? for details). The IDs for each line are consecutive and start at startingID. The parameters given in fix are flagged with 1, while the rest of the parameters are flagged with 0.

Grouping of r parameters (see argument -g on page ?? for details) can be calculated automatically by setting groupingDiff. If the difference between r_i and r_{i+1} is smaller than groupingDiff, than they are placed in the same group. The grouping information is appended to <outputfile>_grouping.

6.3 Calculating α

The function

```
estimateAlpha(estimates, supLimit = 0, div = NULL, poly = FALSE)
```

Using the parameters found in estimates, which is one entry from the list returned by parseOutput, this function calculates both $\alpha_{\rm div}$ (when div = NULL) and $\alpha_{\rm dfe}$ (when div is given) (?).

? argues that mutations with positive selection coefficients that are not higher than a certain threshold should, effectively, be treated as neutral mutations when calculating α (see also ?). This threshold can be controlled here by setting supLimit.

When used, div should contain divergence data, which can be read from filename using the function

parseDivergenceData(filename)

As described in ?, divergence data can contain missatributed polymorphism. This can be corrected for by setting poly = TRUE.

6.4 Model testing

The function

```
compareModels(est1, est2 = NULL, nested = FALSE)
```

compares the models found in est1 and est2 by calculating the AIC and, where relevant, the LRT.

Both est1 and est2 can either be lists returned by parseOutput, or file names, as for parseOutput. If these contain more then one run of polyDFE, then run i from est1 is compared to run i from est2.

The function can automatically detect if models are nested for calculating the LRT, however nestedness can be enforced by setting nested = TRUE. Caution should be used when the r parameters have been grouped (see argument -g on page ?? for details), as in this case the function cannot correctly detect nestedness.

The function returns a list containing two entries

- AIC is a matrix containing the number of estimated parameters, the log likelihood and AIC for est1 in columns 1 3 and for est2 in columns 4 6, for all runs of polyDFE found;
- LRT is a matrix containing the degrees of freedom in column 1, the likelihood of est1 in column 2, the likelihood of est2 in column 3 and the p-value from the LRT test in column 4, for all runs of polyDFE found.

If est2 = NULL, only the AIC is calculated for est1.

6.5 Examples

References

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