## InferNetwork\_ML

## Description

Infers a species network(s) with a specified number of reticulation nodes using maximum likelihood. The returned species network(s) will have inferred branch lengths and inheritance probabilities. During the search, branch lengths and inheritance probabilities of a proposed species network can be either sampled or optimized. For the first case, after the search, users can ask the program to further optimize those parameters of the inferred network. To optimize the branch lengths and inheritance probabilities to obtain the maximum likelihood for that species network, we use Richard Brent's algorithm (from his book "Algorithms for Minimization without Derivatives", p. 79). The species network and gene trees must be specified in the Rich Newick Format.

The inference can be made using only topologies of gene trees, or using both topologies and branch lengths of gene trees. The latter one requires the input gene trees to be ultrametric.

The input gene trees can be gene tree distributions inferred from Bayesian methods like MrBayes. See the second example below.

#### Usage

InferNetwork\_ML geneTreeList numReticulations [-a taxa map] [-bl] [-b threshold] [-s startingNetwork] [-n
numNetReturned] [-h {s1 [,s2...]}] [-w (w1,...,w7)] [-x numRuns] [-m maxNetExamined] [-md moveDiameter] [-rd
reticulationDiameter] [-f maxFailure] [-o] [-po] [-p (rel,abs)] [-r maxRounds] [-t maxTryPerBr] [-i
improveThreshold] [-l maxBL] [-pl numProcessors] [-di] [resultOutputFile]

geneTreeList	Comma delimited list of gene tree identifiers or comma delimited list of sets of gene tree identifiers. See details.	mand atory
numReticulatio ns	Maximum number of reticulations to added.	mand atory
-b threshold	Gene trees bootstrap threshold. Edges in the gene trees that have support lower than threshold will be contracted.	option
-а tаха тар	Gene tree / species tree taxa association.	option
-bl	Use the branch lengths of the gene trees for the inference.	option
-s startingNetw ork	Specify the network to start search. Default value is the optimal MDC tree.	optior al
-n numNetRetu rned	Number of optimal networks to return. Default value is 1.	option al
-h {s1 [, s2]}	A set of specified hybrid species.	option
-w (w1,, w7)	The weights of operations for network arrangement during the network search. Default value is (0.1,0.15,0.55,0.15,0.15,2.8).	option al
-x numRuns	The number of runs of the search. Default value is 5.	option
-m maxNetExa mined	Maximum number of network topologies to examined. Default value is infinity.	option
-md <i>moveDiam</i> eter	Maximum diameter to make an arrangement during network search. Default value is infinity.	option
-rd reticulation Diameter	Maximum diameter for a reticulation event (the distance between two parents of a reticulation node). Default value is infinity.	option
-f maxFailure	Maximum consecutive number of failures for hill climbing. Default value is 100.	option al
-0	If specified, during the search, for every proposed species network, its branch lengths and inheritance probabilities will be optimized to compute its likelihood. Default value is false.	option al
-ро	If specified, after the search the returned species networks will be optimized for their branch lengths and inheritance probabilities. Default value is false.	option al
-p <i>(rel, abs)</i>	The original stopping criterion of Brent's algorithm. Default value is (0.01, 0.001).	option al

-r maxRound	Maximum number of rounds to optimize branch lengths for a network topology. Default value is 100.	option al
-t maxTryPerBr	Maximum number of trial per branch in one round to optimize branch lengths for a network topology. Default value is 100.	option al
-i improveThres hold	Minimum threshold of improvement to continue the next round of optimization of branch lengths. Default value is 0.001.	option al
-l maxBL	Maximum branch lengths considered. Default value is 6.	option al
-pl numProcess ors	Number of processors if you want the computation to be done in parallel. Default value is 1.	option al
-di	Output the Rich Newick string of the inferred network that can be read by Dendroscope .	option al
resultOutputFile	Optional file destination for command output.	option al

It is mandatory to specify the number of reticulation nodes to added to the starting network. By default, the inference uses only the topologies of gene trees, however, users can also use both topologies and branch lengths of the gene trees to do the inference, by specifying option -bl. By default, it is assumed that only one individual is sampled per species in gene trees. However, the option [-a taxa map] allows multiple alleles to be sampled. If users have a prior knowledge of the hybrid species, they can specify them using option -h.

The search: Option -m allows users to specify the maximum number of networks examined during the search. Users can specify the weights of seven operations for network arrangement through option -w. The seven weights correspond to adding a reticulation node, deleting a reticulation node, relocating the head of a reticulation edge, relocating the tail of an edge, reversing the direction of a reticulation edge, replacing a reticulation edge and changing branch lengths and inheritance probabilities, respectively. Furthermore, users can use option -md to specify the maximum move diameter of an operation for network rearrangement, like what local-SPR does. Also, users can use option -rd to specify the maximum reticulation diameter which is the distance (the number of branches) between the two parents of a reticulation node. In order to avoid getting stuck at some local optimum, it is recommended to performed the search multiple times, which users can specify by option -x. The -s option allows the users to specify a starting network (can be a tree) for network search. If the starting network is not specified, the optimal tree under MDC (command infer\_ST\_MDC) will be used. If it is not binary, a random resolution will be used. By default, only the first optimal species network will be returned. However, users can use -n option to ask for multiple optimal networks

During the search, by default, simulated annealing is used (See *Salter* and *Pearl* 2001 for details of settings), where the branch lengths and inheritance probabilities are sampled. In this case, through option *-po*, as a post-processing, users can optimize the branch lengths and inheritance probabilities of the species networks returned by the search. If the dataset is not large and a large a mount of memory is available, users can use option *-o* to optimize the branch lengths and inheritance probabilities of every proposed network during the search. In this case, simple hill climbing will be used, and only the first 5 operations for network arrangement will be used. If branch lengths of the gene trees are used (option *-bl*), the latter case will be applied.

To optimize the branch lengths and inheritance probabilities of a species network, we use Richard Brent's algorithm (from his book "Algorithms for Minimization without Derivatives", p. 79). Users can use different options to control this process. Option -p allows users to specify the original stopping criterion of Brent's algorithm. More precisely, abs and rel define a tolerance tol = rel |x| + abs. We optimize the branch lengths one by one. For every branch, it terminates when either maxTryPerBr (option -t) trials have been made or the Brent's algorithm suggests so. Users can put an upper bound of the branch lengths through option -t. Optimization of all branch lengths consists of a round. After every round, if the improvement in terms of likelihood score is greater than that from last round by at least improveThreshold (option -t), we starts next round. A maximum of maxRound (option -t) rounds will be tried.

If users want to run the computation in parallel (in terms of the gene trees). Please specify the number of processors through option -pl.

## Examples

```
#NEXUS
BEGIN TREES;
Tree geneTree1 = ((C,((B,D),A)),E);
Tree geneTree2 = (B,(D,(C,(A,E))));
Tree geneTree3 = (D,(B,((C,E),A)));
Tree geneTree4 = (D,((B,E),(C,A)));
END;

BEGIN PHYLONET;
InferNetwork_ML (geneTree1,geneTree2,geneTree3,geneTree4) 1;
END;
```

```
#NEXUS
BEGIN TREES;
Tree geneTree1 = [&W 0.9] ((C,((B,D),A)),E);
Tree geneTree2 = [&W 0.1] (B,(D,(C,(A,E))));
Tree geneTree3 = [&W 0.6] (D,(B,((C,E),A)));
Tree geneTree4 = [&W 0.4] (D,((B,E),(C,A)));
END;

BEGIN PHYLONET;
InferNetwork_ML (geneTree1,geneTree2,geneTree3,geneTree4) 1;
END;
```

```
#NEXUS
BEGIN TREES;

Tree geneTree1 = ((C:3,((B:1,D:1):1,A:2):1):1,E:4);
Tree geneTree2 = (B:4,(D:3,(C:2,(A:1,E:1):1):1):1);
Tree geneTree3 = (D:4,(B:3,((C:1,E:1):1,A:2):1):1);
Tree geneTree4 = (D:3,((B:1,E:1):1,(C:1,A:1):1):1);
END;

BEGIN PHYLONET;
InferNetwork_ML (geneTree1,geneTree2,geneTree3,geneTree4) 1 -bl;
END;
```

#### **Command References**

- Y. Yu, N. Ristic and L. Nakhleh. Fast algorithms and Heuristics for Phylogenomics under hybridization and incomplete lineage sorting. BMC Bioinformatics, vol. 14, no. Suppl 15, p. S6, 2013.
- Y. Yu, J. Dong, K. Liu, and L. Nakhleh, Maximum Likelihood Inference of Reticulate Evolutionary Histories, Proceedings of the National Academy
  of Sciences, vol. 111, no. 46, pp. 16448-16453, 2014.

# See Also

• List of PhyloNet Commands