LCA Command

Description

Computes the least common ancestor of a group of nodes in a tree. The tree must be specified in the Rich Newick Format.

Usage

```
LCA tree_ident ({taxon1, taxon2, [, taxon3...]} [{taxon4, taxon5, [, taxon6...]}]) [result output file]
```

tree_ident	The name of the network to analyse.	mandator y
({taxon1, taxon2, [, taxon3]} [,{taxon4, taxon5, [, taxon6]}])	A list of sets of taxa in the specified tree.	mandator y
result output file	Optional file destination for command output.	optional

The tool prints the given tree with all internal nodes labeled. If a node is not labeled in the original tree, then the node receives a generated name. Then the tool reads sets of nodes. For each node set read, the command prints their least common ancestor.

Examples

```
#NEXUS
BEGIN TREES;
Tree tree1 = ((1,4,3),2);
Tree tree2 = ((1,2),(4,3));

END;

BEGIN PHYLONET;
LCA tree2 ({1,4}, {2,3});
END;
```

See Also

• List of PhyloNet Commands