Package 'Triples'

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Title Test your phylog	geny		
Version 1.0			
Author Ambuj Kuma	ır		
Maintainer Ambuj Kumar <ambuj@ufl.edu> Description Calculates resolution of triples from the set of phylogeny trees Depends ape (>= 2.0)</ambuj@ufl.edu>			
		License GPL (>= 2)	
		URL https://github.co	om/Ambuj-UF/Triples
R topics documented:			
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triples	SuperMatrix Rooted Triple (SMRT)		
Description			
Calculates triples	resolution from the input phylogeny tree file.		
Usage			
triples(inpFil	e = "", authorityFile = "", outgroup = "", output)		
Arguments			
authorityFile	File that contains list of taxa to be included for calculating triples resolution. If authorityFile = "" (the default) then the program terminates with message		
inpFile	a file name specified by either a variable of mode character, or a double-quoted string; if inpFile = "" (the default) then the program terminates with message		
outgroup	Tree outgroup taxa name. If outgroup = "" (the default) then the program terminates with message		
output	a file name specified by either a variable of mode character, or a double-quoted string; if output = "" (the default) programs generates output in Triples.txt file		

2 triples

Author(s)

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References

DeGiorgio M, Degnan JH. Fast and Consistent Estimation of Species Trees Using Supermatrix Rooted Triples. Mol Biol Evol. 2010 Mar;27(3):552-69. http://mbe.oxfordjournals.org/content/27/3/552.long

Examples

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triples(inpFile = "", authorityFile = "", outgroup = "", output)
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