## PhyloNetworks and SNaQ Tutorial

Claudia Solís-Lemus

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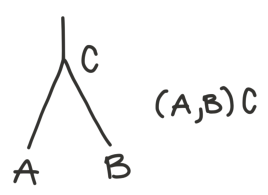
## PhyloNetworks

- Julia package for phylogenetic networks
- www.github.com/crs14/PhyloNetworks

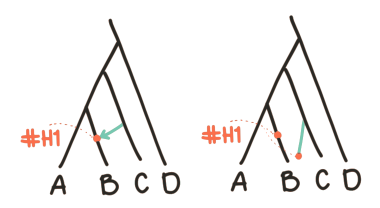


- Read/write networks in parenthetical format
- Plot networks
- Root networks
- Estimate max pseudolikelihood network, bootstrap: SNaQ
- Future: trait evolution models on networks, better parallelization

Recall that internal nodes can have name:

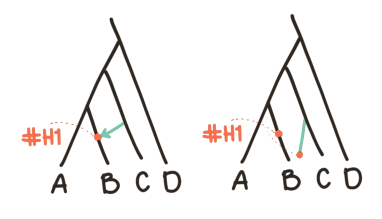


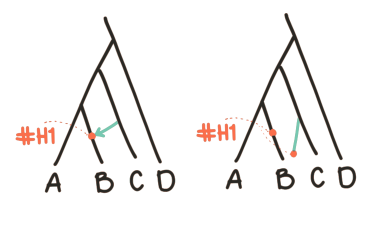
Split the hybrid node into two nodes with the same name:

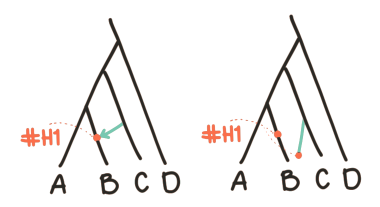


#### By convention:

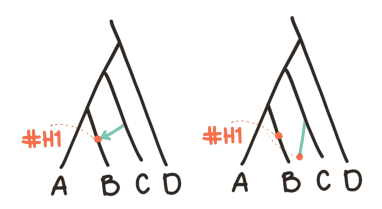
- hybrid name: # + H,LGT,R + number
- minor hybrid edge leads to leaf







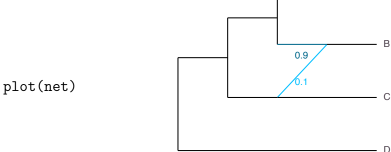
(C, #H1):branch length:bootstrap support:gamma



## Read/write networks in parenthetical format

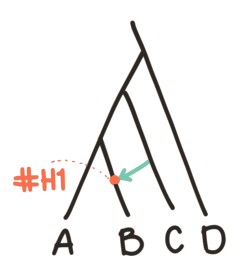
net is a Julia object of type HybridNetwork. It can be a tree.

### Plot networks



### Root networks

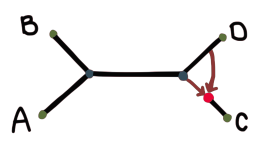
Careful with root position



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### Root networks

Careful with root position: semi-directed networks



```
rootonedge!(net,edge#)
rootatnode!(net,node#)

plot(net, showEdgeNumber=true, showEdgeLength=false)
plot(net, showNodeNumber=true, showEdgeLength=false)

Also: rootatnode!(net,outgroup)
Julia convention: ! in function means it modifies arguments
```

## Estimate max pseudolik network with SNaQ

#### You need:

- List of gene trees or
  - Table of CF
- 2 Starting topology for the search

## Input: list of gene trees

Plain text file: one tree in parenthetical format per line

```
treefile.txt (6:3.500, (1:2.236, (5:1.850, (4:0.660, (2:0.632,3:0.632):0.028):1.190):0.386):1.264); (6:2.569, ((1:0.818, (2:0.677, 4:0.677):0.141):1.268, (3:1.105,5:1.105):0.982):0.483); ((3:0.676, (1:0.104,2:0.104):0.572):2.362, (6:2.790, (4:0.551,5:0.551):2.238):0.248); (6:2.728, ((3:0.655,5:0.655):1.202, (1:0.881, (2:0.783,4:0.783):0.098):0.976):0.871); (6:4.714, (5:2.433, (3:1.318, (4:0.896, (1:0.506,2:0.506):0.391):0.422):1.114):2.281); (6:2.210, ((3:0.694,5:0.694):1.302, (4:0.726, (1:0.351,2:0.351):0.376):1.270):0.214); (6:3.077, (5:2.349, ((1:0.499,2:0.499):0.764, (3:0.730,4:0.730):0.473):1.146):0.728); (((1:0.211,2:0.211):1.629, (3:0.754,4:0.754):1.086):0.801, (5:2.280,6:2.280):0.361); (6:2.181, ((4:0.875,5:0.875):1.131, (2:1.304, (1:0.821,3:0.821):0.484):0.701):0.176); (6:2.249, (1:2.092, ((2:0.617,4:0.617):1.413, (3:0.749,5:0.749):1.281):0.062):0.157);
```

RAxML or MrBayes will put gene trees in separate files, you need to combine these files into one (PhyloNetwork does not do this!)

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## Input: table of CF

```
Plain text file, comma separated (or ;), 7 columns:
```

- 1 Taxon 1
- 2 Taxon 2
- 3 Taxon 3
- 4 Taxon 4
- $CF_{12|34}$
- 6  $CF_{13|24}$
- **7**  $CF_{14|23}$

```
ableCF.txt >
"t1","t2","t3","t4","CF12 34","CF13 24","CF14 23"
"6","1","5","4",0.4,0.6,0.0
"6","1","5","2",0.2,0.8,0.0
"6","1","5","3",0.5,0.5,0.0
"6","1","4","2",0.4,0.6,0.0
"6","1","4","3",0.4,0.2,0.4
"6","1","2","3",0.2,0.1,0.7
"6","5","4","2",0.8,0.0,0.2
"6","5","4","3",0.4,0.4,0.2
"6","5","2","3",0.6,0.4,0.0
"6","4","2","3",0.3,0.2,0.5
"1", "5", "4", "2", 0.4, 0.0, 0.6
"1","5","4","3",0.3,0.5,0.2
"1","5","2","3",0.1,0.8,0.1
"1","4","2","3",0.1,0.5,0.4
"5","4","2","3",0.3,0.2,0.5
```

This involves running BUCKy on every 4-taxon subset of the taxa, and get the CF from the three possible splits from the .concordance file into a table (PhyloNetworks does not do this!)

## Input: starting topology

Plain text file: tree (or networks )in parenthetical format

```
startTree.txt ~
(6:3.500, (1:2.236, (5:1.850, (4:0.660, (2:0.632, 3:0.632):0.028):1.190):0.386):1.264);
```

BUCKy can give you a primary concordance tree (but recall that you ran BUCKy analyses on subsets of taxa before, not all taxa)

networks April 7th, 2016 13 / 27 Input: list of gene trees or table of CF

There is a lot of pre-processing before running SNaQ

## Input: list of gene trees or table of CF

There is a lot of pre-processing before running SNaQ

Luckily for us: https://github.com/nstenz/TICR

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## Input: table of CF and starting tree with TICR

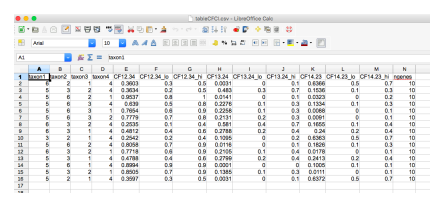
https://github.com/nstenz/TICR

- MDL to delimit loci
- MrBayes to perform individual gene analyses
- BUCKy to estimate quartet concordance factors
- Quartet MaxCut to estimate a binary population tree

TICR gives us everything we need!

## Input: table of CF with TICR

#### https://github.com/nstenz/TICR



```
d = readTableCF("tableCFCI.csv")
    T = readTopologyLevel1("startTree.txt")
Estimate best network with h_{max} = 1, runs = 10 (default):
               estNet1 = snaq!(T,d)
                  plot(estNet1)
  You can also change the number of independent runs:
          estNet1 = snaq!(T,d, runs=5)
   Suggestion for big datasets: Start with runs=1
```

#### snaq.out

Default rootname: snaq (can be set with the option filename=)

-Pseudolik: the smaller, the better!

You can read the best network overall, and the best network per run with:

nets = readInputTrees("snaq.out")
and plot one of them: plot(nets[2])

#### snaq.log

```
snag.log ~
optimization of topology. BL and inheritance probabilities using:
 tolerance parameters: ftolRel=1.0e-5, ftolAbs=1.0e-6,
                       xtolAbs=0.0001, xtolRel=0.001.
 max number of failed proposals = 100, multiplier M = 10000.
Outgroup: none (for rooting at the final step)
rootname for files: snag
BEGIN: 10 runs on starting tree (2.3.(4.(5.(1.6):0.386):1.19):0.028):
Wed Apr 6 18:19:08 2016
main seed 38429
seed: 38429 for run 1
 BEGIN SNaO for run 1, seed 38429 and hmax 1 changed starting topology by NNI move
Begins heuristic optimization of network-----
found best network, now we re-optimize branch lengths and gamma more precisely
STOPPED for not having more moves to propose: movesfail [0,25,25,4,0,61], Nmov [217,25,25,4,10000,61]
END optTopLevel: found minimizer topology at step 128 (failures: 54) with -loglik=113.59691 and
ht min=[0.18096.7.49654.0.35354.2.97594.0.60066.5.03393.0.0]
PERFORMANCE: total number of moves (proposed, successful, accepted) in general, and to fix gamma=0.0,t=0.0 cases
         -----moves general-----
                                                                            -moves gamma.t----
         Num. Proposed
move
                          Num.Successful Num.Accepted
                                                              Num. Proposed
                                                                             Num.Successful
                                                                                              Num. Accepted
add
                                                                     NΑ
mvorigin 29
mytarget 29
                          29
chdir
                  5
                                   5
delete
nni
                  63
Total
                  127
                                   65
                                                                                                       0.8
Proportion
FINISHED SNaQ, typeof best PhyloNetworks, HybridNetwork, -loglik of best 113.59691264573962
((6, (5, #H7:5.034::0.181):7.497):2.976, (1,2):0.601, ((4,3):0.0) #H7:0.354::0.819);
seed: 58439 for run 2
```

## SNaQ output files: .err file

snaq.err



Total errors: 0 in seeds Int64[]

#### Example:

Total errors: 1 in seeds [4545]

You need to run the following function with the same settings that caused the error:

snaqDebug(T,d,seed=4545)

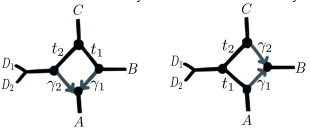
**Suggestion 1:** Start the estimation for  $h_{max} = h$  in the best network at  $h_{max} = h - 1$ 

Suggestion 2: Keep in mind the level-1 restriction, maybe no more hybridizations can be added if the number of taxa is small!

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# Troubleshooting: my network does not make sense!

- Recall that the root is meaningless, use root functions
- Recall the identifiability issue with direction of hybrid edges:



use topologyMaxQPseudolik!(newNet,d)

- Start the estimation for  $h_{max} = h$  in the best network at  $h_{max} = h 1$
- Recall **level-1** assumption: maybe no more hybridizations can be added
- Recall that the bigger the network, the more gene trees are needed

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## Bugs and errors

- Use PhyloNetworks users google group (link in Github)
- Use "Issues" in Github
- Send me an email claudia@stat.wisc.edu, specially with output from snaqDebug

## Getting help

- Use PhyloNetworks users google group (link in Github)
- README file in Github has step-by-step commands
- ?plot inside Julia
- Soon: http://phylonetworks.readthedocs.org/en/latest/
- Send me an email claudia@stat.wisc.edu

```
using DataFrames

df = readtable("tableCFCI.csv")

bootNet = bootsnaq(T,df,hmax=1,nrep=10,bestNet=estNet1,
runs=1)

nrep is the number of bootstrap replicates
bestNet is the estimated network with original data
bootNet is a vector of size nrep with estimated network per replicate
Output files: .out, .log, .err per bootstrap replicate, and bootsnaq.out,
bootsnaq.log
```

#### Bootstrap: summary

- df0,tree0 = treeEdgesBootstrap(bootNet,estNet1) will calculate the bootstrap support of tree edges by edge number in df0, and the main underlying tree in tree0
- plot(tree0, showEdgeNumber=true, showEdgeLength=false)

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#### Bootstrap: summary

HFmat,discTrees = hybridDetection(bootNet,estNet1,outgroup) to summarize bootstrap on hybrid nodes: HFmat will have one row per bootstrap network, and number of columns depending on number of hybrids in estNet1: for example, if estNet1 had 3 hybrids for example, HFmat will have 6 columns:

- first 3 columns indicate the presence (1.0) or absense (0.0) of each hybrid (column) for each bootstrap network (row)
- the last 3 columns indicate the estimated gamma in the bootstrap network if the hybrid was found (0.0 if not found)

### Bootstrap: summary

Only makes sense to summarize networks that have the same underlying tree: discTrees has different trees to the underlying tree in estNet1 length(discTrees)

dfhyb = summarizeHFdf(HFmat)
dfhyb has one row per hybrid, and 5 columns:

- hybrid index
- number of trees that match the underlying tree in estNet1 (same for all hybrids)
- number of networks with that hybrid
- mean estimated gamma among networks with the hybrid
- sd estimated gamma among networks with the hybrid

Note that last row contains in 3er column the number of networks that have all same hybrids as estNet1 (hybrid index, mean gamma and sd gamma are meaningless)