dmeav02 Usage

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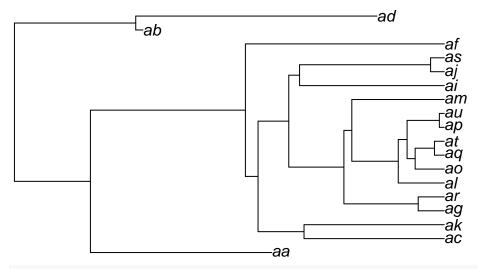
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tree simulation and consistency

functions

- 1. sim.tree to simulate trees
- 2. phylo2vectors to convert a phylo into the set of vectors characterizing the tree

```
s = sim.tree(ct=6)
plot(s$phylo)
```



s\$tree

```
## $wt
   [1] 1.34229643 0.82351820 0.49113791 0.07833963 1.11018263 0.13703041
   [7] 0.14772407 0.18520556 0.11678888 0.04850796 0.43201880 0.08572596
## [13] 0.27286554 0.23045126 0.09587227 0.08744999 0.03298648 0.13411852
  [19] 0.04211276 0.05302846 0.05263828
##
##
## $E
##
   ##
## $n
                4 3 4 5 4 5
                                6
                                   7 8 9 8 9 10 11 12 13 14 15
##
   Г17
       1 2
##
   [1] "aa" "aa" "ab" "ab" "ac" "ac" "aa" "ag" "ai" "ac" "ag" "al" "ad" "al"
  [15] "ao" "ao" "ag" "aj" "aq" "ap"
##
##
## $br
##
   [1] 1.342296 2.165815 2.656953 2.735292 3.845475 3.982505 4.130229
  [8] 4.315435 4.432224 4.480732 4.912750 4.998476 5.271342 5.501793
## [15] 5.597666 5.685116 5.718102 5.852221 5.894333 5.947362 6.000000
```

```
phylo2vectors(s$phylo)
## $wt
  [1] 1.34229643 0.82351820 0.49113791 0.07833963 1.11018263 0.13703041
##
## [7] 0.14772407 0.18520556 0.11678888 0.04850796 0.43201880 0.08572596
## [13] 0.27286554 0.23045126 0.09587227 0.08744999 0.03298648 0.13411852
## [19] 0.04211276 0.05302846 0.05263828
##
## $E
   ##
##
## $n
## [1] 1 2 3 4 3 4 5 4 5 6 7 8 9 8 9 10 11 12 13 14 15
all.equal(s$tree$wt,phylo2vectors(s$phylo)$wt)
## [1] TRUE
all.equal(s$tree$E,phylo2vectors(s$phylo)$E)
## [1] TRUE
all.equal(s$tree$n,phylo2vectors(s$phylo)$n)
## [1] TRUE
to do:
  1. phylo2vectorsdoes not give the vector S corresponding to topology. add it
  2. remove the root on vector outputs, keep consistency
  3. arenge the vectors2phylo function and make a lot simpler the sim.tree using it
  4. add the log in an log output file
Tree manipulation
s = sim.tree(ct=2, seed=5)
plot(s$phylo)
                                                              ·ab
                                                              ac
```

s2=phylo2vectors(s\$phylo)
s2

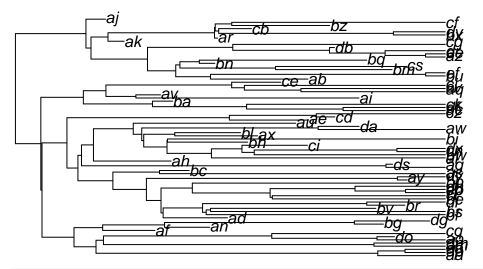
·aa

```
## $wt
## [1] 1.0954824 0.4297802 0.4747374
##
## $E
## [1] 1 1
##
## $n
## [1] 1 2 3
s$tree
## $wt
## [1] 1.0954824 0.4297802 0.4747374
## $E
## [1] 1 1
##
## $n
## [1] 1 2 3
##
## $S
## [1] "aa" "aa"
##
## $br
## [1] 1.095482 1.525263 2.000000
up = update.tree(s2, 0.5, 1.5)
up
## $wt
## [1] 0.50000000 0.59548244 0.40451756 0.02526264 0.47473736
## $E
## [1] 1 1 0 1
##
## $n
## [1] 1 2 3 2 3
```

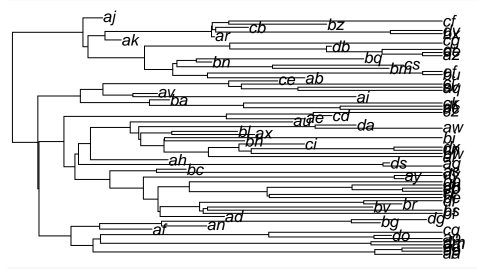
- 1. Doing update tree I realized that I should be more carefully in analysis when we lost crown time.
- 2. phylo2vectors really needs to take into account topology

more about consistency

```
s <- sim.tree()
plot(s$phylo)</pre>
```



plot(vectors2phylo(s\$tree))

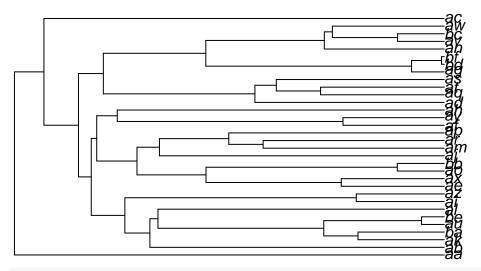


all.equal(s\$phylo,vectors2phylo(s\$tree))

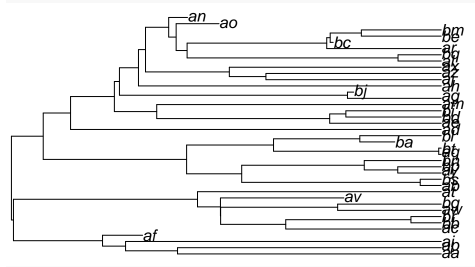
[1] TRUE

The reconstruction algorithm

```
s = sim.tree(ct=6)
plot(s$phylo.extant)
```



rec = rec.tree(tree=s\$tree.extant,pars=c(0.8,0.1,40))
plot(vectors2phylo(rec))



s\$tree.extant

```
## $wt
## [1] 0.520024562 0.376312474 0.440344992 0.162930297 0.069802563
## [6] 0.084344775 0.177608878 0.097022792 0.154247328 0.164701987
## [11] 0.102840481 0.019100064 0.590781785 0.002528972 0.290119065
## [16] 0.333377530 0.104880179 0.168239579 0.564265412 0.040787014
## [21] 0.006960564 0.102622237 0.113200338 0.022240266 0.169834369
## [26] 0.021267363 0.501251269 0.003713396 0.177518393 0.127974050
## [31] 0.254726892 0.034430135
##
## $E
   ##
## $n
  [1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23
##
## [24] 24 25 26 27 28 29 30 31 32
```

```
rec
## $wt
  [1] 0.082044932 0.028035271 0.409944359 0.376312474 0.440344992
## [6] 0.162930297 0.069802563 0.084344775 0.177608878 0.056818678
## [11] 0.040204115 0.154247328 0.164701987 0.102840481 0.019100064
## [16] 0.124804516 0.006977283 0.005780751 0.137490296 0.296151939
## [21] 0.019577001 0.002528972 0.120048496 0.170070569 0.333377530
## [26] 0.104880179 0.168239579 0.564265412 0.040787014 0.006960564
## [31] 0.041161701 0.061460536 0.085317081 0.027883257 0.022240266
## [36] 0.085197504 0.084636865 0.021267363 0.039924189 0.420559779
## [41] 0.040767300 0.003713396 0.177518393 0.127974050 0.254726892
## [46] 0.034430135
## $E
## [36] 0 1 1 1 0 1 1 1 1 1
##
## $n
## [1] 1 2 3 4 5 6 7 8 9 10 9 10 11 12 13 14 15 16 15 16 15 16 17
## [24] 18 19 20 21 22 23 24 25 24 25 24 25 26 25 26 27 28 27 28 29 30 31 32
## $prob
## [1] -35.05333
```

let's observe one estimation

```
s = sim.tree(seed = 1)
tree = s$tree.extant
st = sim.srt(tree,pars=c(0.8,0.1,40),n_trees=100)
mle.st(st)
## [1] 0.85475693 0.09726184 39.07543002
```

How good is the last iteration of the MCEM algorithm?

```
library(dmea)
n_sim = 537
n_trees = 10
MP = matrix(nrow=n_sim,ncol=3)
RP = matrix(nrow=n_sim,ncol=3)
p = proc.time()
for(i in 1:n_sim){
    est = sim.est(n_trees=n_trees,pars=c(0.8,0.1,40),seed=i)
    RP[i,] = est$real
    MP[i,] = est$est
}
print(proc.time()-p)
par_est_vis(P=MP,par=1,PR=RP)
par_est_vis(P=MP,par=2,PR=RP)
par_est_vis(P=MP,par=3,PR=RP)
```