Simulated tree - Feb 2016

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```
library(ape)
library(ggplot2)
```

I. LOAD STUFF

Import simulation outputs

Import ExaML tree 0

```
t_uk <- read.tree(file = "../phylo-uk/data/ExaML_result.subUKogC_noDRM.finaltree.000")
## drop OG
og <- c("Ref1", "Ref2", "Ref3", "Ref4", "Ref5", "Ref6", "HXB2")
t_uk <- drop.tip(t_uk, og )
t_uk

Phylogenetic tree with 12164 tips and 12163 internal nodes.

Tip labels:
34695, 21677, 72292, 81292, 85197, 53538, ...

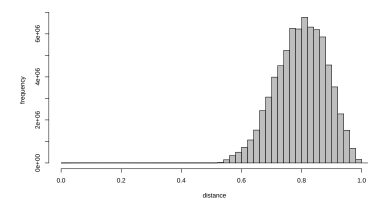
Rooted; includes branch lengths.</pre>
```

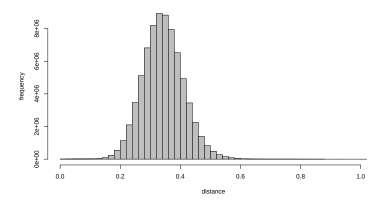
II. TREE DISTANCES

Patristic distances?

```
#### cluster size to real data. Need to have same number of clusters ?
## get distances
```

```
##- matrix first into distances
#- sim tree
if (file.exists("data/simtree_dist.rds")){
 dsimtree <- readRDS("data/simtree_dist.rds")</pre>
} else {
dsimtree <- as.dist(cophenetic.phylo(tree))</pre>
saveRDS(dsimtree, file = "data/simtree_dist.rds")
}
# uk tree
if (file.exists("data/uktree_dist.rds")){
 duktree <- readRDS("data/uktree_dist.rds")</pre>
 duktree <- as.dist(cophenetic.phylo(t_uk))</pre>
  saveRDS(duktree, file = "data/uktree_dist.rds")
head(dsimtree)
[1] 24930 24930 24930 24930 24930 24930
head(duktree)
[1] 2.000001e-06 8.505254e-02 7.145780e-02 1.257209e-01 1.010726e-01 1.104545e-01
simx <- dsimtree / (max(dsimtree) - min(dsimtree))</pre>
ukx <- duktree / (max(duktree) - min(duktree))</pre>
# rm(simtree, uktree)
\#\#- histogram distances
# summary(x)
hist(simx, breaks = 50, xlab = "distance", ylab = "frequency", main = "", col = "grey")
hist(ukx, breaks = 50, xlab = "distance", ylab = "frequency", main = "", col = "grey")
```





III. CLUSTERING

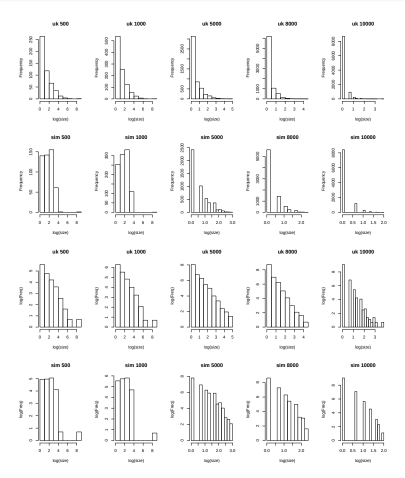
Clustering with fixed number of groups?

```
simhc <- hclust(simx, method = "average") # UPGMA</pre>
ukhc <- hclust(ukx, method = "average")</pre>
##- cut based on height
if (F){
#- function of heights
nheights <- 10 # number of threshold
up <- round(mean(simx), 1) # cut up to the mean
# breaks \leftarrow seq(1/nbreaks, 1-(1/nbreaks), by = 1/nbreaks)
simclus \leftarrow cutree(simhc, h = seq(up / nheights, up, by = up / nheights) ) # h = breaks
up <- round(mean(ukx),1) # cut up to the mean
ukclus <- cutree(ukhc, h = seq(up / nheights, up, by = up /nheights) )
# rm(simx, ukx)
}
#- cut as function of k groups
kgroups <- c(500, 1000, 5000, 8000, 10000)
simclus <- cutree(simhc, k = kgroups ) # h = breaks</pre>
ukclus <- cutree(ukhc, k = kgroups )</pre>
\#\ colnames(simclus)\ \leftarrow\ paste("k",colnames(simclus),sep='')
\# colnames(ukclus) <- paste("k",colnames(ukclus),sep='')
head(simclus)
```

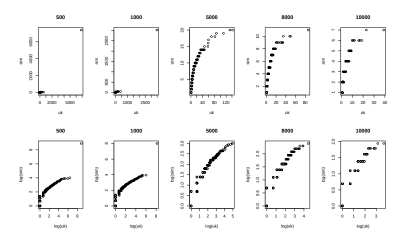
```
500 1000 5000 8000 10000
7
    1
       1 1
                  1
8
    2
         2
             2
                  2
9
    3
         3
             3
                  3
                       3
10
        4
             4
                  4
                       4
    4
      5 5 5
                       5
11 5
      6
12
   6
           6
                6
                       6
head(ukclus)
     500 1000 5000 8000 10000
         1 1 1
34695
      1
            1
                1
21677
       1
           1
                2
72292
       1
                     2
81292
       1
           1
                1
                     3
85197
     1
           1
                3
                     4
                          4
53538 1
          1
                4
                     5
                          5
##- Calculate size(=Freq) of each cluster across different threshold
simfreqClust <- apply(simclus, 2, function(x) as.data.frame(table(x))) # list</pre>
ukfreqClust <- apply(ukclus, 2, function(x) as.data.frame(table(x)))</pre>
# str(simfreqClust)
# head(simfreqClust[[1]])
##- number of different clusters by threshold # if number varies !
\# sapply(simfreqClust, function(x) dim(x)[1])
# sapply(ukfreqClust, function(x) dim(x)[1])
##- cluster size
#- sim
sapply(simfreqClust, function(x) summary(x$Freq))
          500
               1000 5000 8000 10000
         1.00 1.00 1.000 1.00 1.000
Min.
1st Qu.
        2.00 2.00 1.000 1.00 1.000
Median
         6.00 6.00 2.000 1.00 1.000
       24.33 12.16 2.433 1.52 1.216
3rd Qu. 13.00 13.00 3.000 2.00 1.000
      7395.00 3019.00 20.000 11.00 7.000
Max.
sapply(ukfreqClust, function(x) summary(x$Freq))
               1000
                       5000 8000 10000
          500
         1.00
               1.00 1.000 1.00 1.000
Min.
        1.00 1.00 1.000 1.00 1.000
1st Qu.
Median
         2.00 2.00 1.000 1.00 1.000
Mean
         24.33 12.16
                       2.433 1.52 1.216
3rd Qu. 7.00 6.00
                       2.000 1.00 1.000
      6868.00 3486.00 142.000 86.00 39.000
##- percentiles
# sapply(freqClust, function(x) round(quantile(x$Freq,
\# probs = c(0.01, 0.05, 0.1, 0.25, 0.5, 0.75, 0.95, 0.99, 1))))
```

Distribution of sizes - semitransformed, transformed

```
##- distr of cluster sizes: log(x) and Y untransformed
par(mfcol=c(2, length(kgroups)))
for (i in 1:length(kgroups)){
 h <- hist(log(ukfreqClust[[i]]$Freq),</pre>
       main = paste("uk", names(ukfreqClust)[i]),
       xlab = "log(size)")
  hist(log(simfreqClust[[i]]$Freq),
       main = paste("sim", names(simfreqClust)[i]),
       xlab = "log(size)")
}
##- distr of cluster sizes: log(x) and log(y)
par(mfcol=c(2, length(kgroups)))
for (i in 1:length(kgroups)){
 h <- hist(log(ukfreqClust[[i]]$Freq), plot = F)</pre>
 h$counts <- log1p(h$counts) # log(y)
  plot(h, ylab = "log(Freq)",
       main = paste("uk", names(ukfreqClust)[i]),
       xlab = "log(size)")
  h <- hist(log(simfreqClust[[i]]$Freq), plot = F)</pre>
  h$counts <- log1p(h$counts) # log(y)
  plot(h, ylab = "log(Freq)",
          main = paste("sim", names(simfreqClust)[i]),
          xlab = "log(size)")
```



QQ plot



IV. ADD PATIENTS DATA

Add data from sample states and time

```
##- converting sample states in table of co-variates ?
demes <- as.vector(read.csv(file = "demes.csv")$x)</pre>
sampleTimes <- scan( file = 'sampleTimes' )</pre>
ss <- matrix( scan( file = 'sampleStates' ) ,</pre>
                byrow = TRUE,
                ncol = length(demes))
colnames(ss) <- demes</pre>
dim(ss)
[1] 12164
            121
max(ss[,121]) # nothing on source
[1] 0
demo <- data.frame()</pre>
for (i in 1:dim(ss)[1]){ # dim(ss)[1]
  deme <- names(which(ss[i,] == 1)) # name of column which has value 1
  patient <- i
  time <- sampleTimes[i]</pre>
  age <- as.numeric( regmatches( deme,</pre>
                 regexec( "\\.age([0-9])", deme) )[[1]][2] )
```

```
care <- as.numeric( regmatches( deme,</pre>
                regexec( "care([0-9])", deme) )[[1]][2] )
  stage <- as.numeric( regmatches( deme,</pre>
                regexec( "stage([0-9])", deme) )[[1]][2] )
 risk <- as.numeric( regmatches( deme,</pre>
                regexec( "riskLevel([0-9])", deme) )[[1]][2] )
 demo <- rbind(demo, cbind(</pre>
    patient, time, age, care, stage, risk))
str(demo)
'data.frame': 12164 obs. of 6 variables:
$ patient: num 1 2 3 4 5 6 7 8 9 10 ...
 $ time : num 11245 8627 6679 8446 8900 ...
        : num 3 3 4 4 4 3 4 3 3 4 ...
 $ age
 $ care : num 1 1 1 1 1 1 1 1 1 1 ...
 $ stage : num 1 1 1 5 4 2 5 2 3 5 ...
         : num 1 1 1 1 1 1 1 1 2 1 ...
 $ risk
##- date of diagnosis ?
date0 <- as.Date('1979-01-01')</pre>
demo$datediag <- date0 + demo$time</pre>
min(demo$datediag)
[1] "1996-11-15"
max(demo$datediag)
[1] "2013-02-15"
```

Add cluster size by patient

```
##- function to calculate both numclus and sizeclus for each sequindex into a LIST
##- with same variable names
  ##- in list
  1 <- list()
  for (i in 1:length(kgroups)) {
  #- cluster number
  numclus <- as.data.frame(simclus[, i])</pre>
  numclus <- cbind(rownames(numclus), numclus)</pre>
  colnames(numclus) <- c("id", "num")</pre>
  row.names(numclus) <- NULL</pre>
  # head(numclus)
  #- size of cluster
  a <- merge(x = numclus, y = simfreqClust[[i]],
             by.x = "num", by.y = "x",
             all.x = TRUE, sort = FALSE)
  #- binary clustering variable
  a$Clus <- ifelse(a$Freq > 1, 1, 0)
  #- colnames
  colnames(a) [which(colnames(a) =="Freq")] <- "size"</pre>
  colnames(a) [which(colnames(a) == "Clus")] <- "clus"</pre>
  l[[i]] <- a
  names(1)[i] <- names(simfreqClust[i])</pre>
  }
 rm(a, numclus)
```

```
# str(l)
##-proportion in or out clusters
sapply(1, function(x) round(prop.table(table(x$clus)),2))
   500 1000 5000 8000 10000
0 0.01 0.01 0.2 0.46 0.69
1 0.99 0.99 0.8 0.54 0.31
##- cluster sizes
sapply(1, function(x) summary(x$size))
        500
             1000 5000 8000 10000
Min. 1 1.0 1.000 1.000 1.000 1st Qu. 22 11.0 2.000 1.000 1.000
             21.0 4.000 2.000 1.000
Median 7395
Mean 4504 762.7 4.425 2.223 1.488
3rd Qu. 7395 52.0 6.000 3.000 2.000
Max. 7395 3019.0 20.000 11.000 7.000
```

V. REGRESSIONS

A. Logistic

```
##- model: clus ~ age + stage + time + risk
##- care = 1 for all at diagnosis
## ex.
logit_model = "clus ~ age + stage + time + risk"
logit_model_std = "clus ~ scale(age) + scale(stage) + scale(time) + scale(risk)"
lapply(listclus, function(x) summary(glm(formula = logit_model_std,
                               data = x,
                               family = binomial(link = "logit"))))
$`500`
glm(formula = logit_model_std, family = binomial(link = "logit"),
   data = x)
Deviance Residuals:
   Min 1Q Median
                             3Q
                                      Max
-3.2932 0.1035 0.1089 0.1179 0.1598
Coefficients:
   Estimate Std. Error z value Pr(>|z|)
```

```
(Intercept) 5.06851 0.11629 43.585 <2e-16 ***
scale(age) 0.07370 0.11120 0.663 0.5075
scale(stage) 0.01809 0.11472 0.158 0.8747
scale(time) -0.09012 0.11632 -0.775 0.4385
scale(risk) -0.18421 0.10107 -1.823 0.0684.
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 943.23 on 12163 degrees of freedom
Residual deviance: 939.00 on 12159 degrees of freedom
AIC: 949
Number of Fisher Scoring iterations: 8
$`1000`
Call:
glm(formula = logit_model_std, family = binomial(link = "logit"),
   data = x
Deviance Residuals:
  Min 1Q Median 3Q
                                 Max
-3.2188 0.1310 0.1442 0.1589 0.2238
Coefficients:
          Estimate Std. Error z value Pr(>|z|)
(Intercept) 4.55488 0.09111 49.994 <2e-16 ***
scale(age) 0.05994 0.08537 0.702 0.4826
scale(stage) 0.02410 0.08834 0.273 0.7850
scale(risk) -0.16634
                   0.07873 -2.113 0.0346 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 1456.7 on 12163 degrees of freedom
Residual deviance: 1445.4 on 12159 degrees of freedom
AIC: 1455.4
Number of Fisher Scoring iterations: 7
$`5000`
Call:
glm(formula = logit_model_std, family = binomial(link = "logit"),
   data = x
Deviance Residuals:
  Min 1Q Median 3Q
-1.8353 0.6524 0.6639 0.6708 0.6828
Coefficients:
```

```
Estimate Std. Error z value Pr(>|z|)
(Intercept) 1.393642 0.022721 61.338 <2e-16 ***
scale(age) 0.001913 0.022860 0.084 0.933
scale(stage) 0.002938 0.022910 0.128 0.898
scale(time) -0.024301 0.022878 -1.062 0.288
scale(risk) 0.013449 0.022864 0.588 0.556
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 12135 on 12163 degrees of freedom
Residual deviance: 12134 on 12159 degrees of freedom
AIC: 12144
Number of Fisher Scoring iterations: 4
$`8000`
Call:
glm(formula = logit_model_std, family = binomial(link = "logit"),
   data = x)
Deviance Residuals:
 Min 1Q Median 3Q
                              Max
-1.265 -1.245 1.095 1.111 1.142
Coefficients:
             Estimate Std. Error z value Pr(>|z|)
(Intercept) 0.1585314 0.0181926 8.714 <2e-16 ***
scale(age) -0.0006118 0.0183335 -0.033 0.973
scale(stage) 0.0180774 0.0183458 0.985 0.324
scale(time) -0.0027158 0.0182654 -0.149
                                        0.882
scale(risk) -0.0204569 0.0181770 -1.125
                                        0.260
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 16787 on 12163 degrees of freedom
Residual deviance: 16784 on 12159 degrees of freedom
AIC: 16794
Number of Fisher Scoring iterations: 3
$`10000`
Call:
glm(formula = logit_model_std, family = binomial(link = "logit"),
   data = x)
Deviance Residuals:
   Min 1Q Median
                           3Q
-0.9038 -0.8656 -0.8471 1.5116 1.6187
```

```
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -0.80789 0.01964 -41.132 <2e-16 ***
scale(age) 0.02309 0.01985 1.163
                                          0.245
scale(stage) 0.03683 0.01981 1.859
                                        0.063 .
scale(time) 0.02224 0.01975 1.126 0.260
scale(risk) -0.01789 0.01974 -0.906 0.365
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 15031 on 12163 degrees of freedom
Residual deviance: 15024 on 12159 degrees of freedom
AIC: 15034
Number of Fisher Scoring iterations: 4
# logistic <- function(x, m = logit_model){</pre>
# fit \leftarrow glm(m, data = x,
#
                    family = binomial(link = "logit"))
# co <- coef(summary(fit))</pre>
# ## odds ratios and 95% CI
# # or <- exp(cbind(OR = coef(fit), confint(fit)))
# # return(list(co, or))
   return(cbind(co[,c(1,4)]))
#
# }
#
# ##- test 1 level
# c <- listclus[[1]]
\# logistic(x = c, m = logit_model_std)
# ##- all levels
# lapply(listclus, function(x) logistic(x, m = logit_model_std))
```

B. Linear

```
0.453
scale(stage) -6.856e-03 9.144e-03 -0.750
scale(time) 4.858e-03 9.104e-03 0.534 0.594
scale(risk) -2.027e-03 9.070e-03 -0.224
                                           0.823
Residual standard error: 1 on 12159 degrees of freedom
Multiple R-squared: 8.078e-05, Adjusted R-squared: -0.0002482
F-statistic: 0.2456 on 4 and 12159 DF, p-value: 0.9125
$`1000`
Call:
lm(formula = lm_model_std, data = x)
Residuals:
           1Q Median
                          3Q
  Min
                                  Max
-0.6277 -0.5808 -0.5659 -0.5336 1.7610
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) -2.208e-14 9.067e-03
                                 0.000 1.000
scale(age) -1.427e-02 9.138e-03 -1.562
                                           0.118
                                          0.831
scale(stage) -1.947e-03 9.144e-03 -0.213
scale(time) 1.686e-03 9.103e-03 0.185
                                          0.853
scale(risk) 1.506e-03 9.070e-03 0.166
                                           0.868
Residual standard error: 1 on 12159 degrees of freedom
Multiple R-squared: 0.0002211, Adjusted R-squared: -0.0001078
F-statistic: 0.6724 on 4 and 12159 DF, p-value: 0.6111
$`5000`
Call:
lm(formula = lm_model_std, data = x)
Residuals:
           1Q Median
                           3Q
   Min
-1.0397 -0.6975 -0.1676 0.4660 4.5356
Coefficients:
              Estimate Std. Error t value Pr(>|t|)
             2.744e-16 9.067e-03 0.000 1.0000
(Intercept)
                                 0.091
            8.351e-04 9.137e-03
                                          0.9272
scale(age)
scale(stage) -2.079e-04 9.143e-03 -0.023
                                          0.9819
                                         0.7079
scale(time) 3.411e-03 9.103e-03 0.375
scale(risk) 1.969e-02 9.069e-03 2.171 0.0299 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 1 on 12159 degrees of freedom
Multiple R-squared: 0.0004, Adjusted R-squared: 7.114e-05
F-statistic: 1.216 on 4 and 12159 DF, p-value: 0.3015
$`8000`
```

```
Call:
lm(formula = lm model std, data = x)
Residuals:
   Min
          1Q Median 3Q
-0.7774 -0.7385 -0.1467 0.4663 5.3336
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) 2.257e-15 9.068e-03 0.000 1.000
scale(age) -5.272e-03 9.138e-03 -0.577 0.564
scale(stage) 8.204e-03 9.144e-03 0.897 0.370
scale(time) 7.313e-03 9.104e-03 0.803 0.422
scale(risk) -1.942e-03 9.070e-03 -0.214 0.830
Residual standard error: 1 on 12159 degrees of freedom
Multiple R-squared: 0.000139, Adjusted R-squared: -0.0001899
F-statistic: 0.4227 on 4 and 12159 DF, p-value: 0.7924
$`10000`
lm(formula = lm_model_std, data = x)
Residuals:
  Min
          1Q Median
                        3Q
                               Max
-0.6097 -0.5563 -0.5292 0.5587 6.2095
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) -1.373e-15 9.065e-03 0.000 1.0000
scale(age) 9.973e-03 9.136e-03 1.092 0.2750
scale(stage) 2.187e-02 9.142e-03 2.392 0.0168 *
scale(time) 9.734e-03 9.101e-03 1.070 0.2848
scale(risk) 2.170e-03 9.067e-03 0.239 0.8109
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 0.9998 on 12159 degrees of freedom
Multiple R-squared: 0.0006847, Adjusted R-squared: 0.000356
F-statistic: 2.083 on 4 and 12159 DF, p-value: 0.08024
```