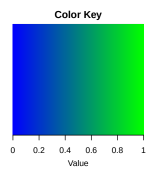
uk3

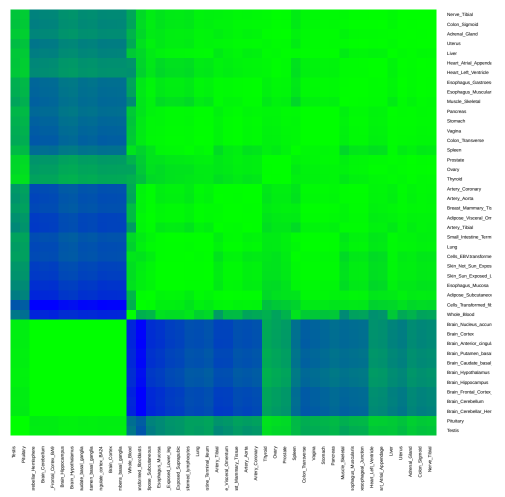
Here we plot the correlation matrix and the first 3 eigenvectors of Uk3.

library('knitr')

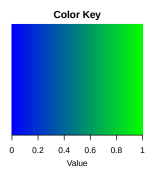
```
knitr::opts_chunk$set(cache=TRUE)
opts_chunk$set(fig.path = "/Users/sarahurbut/Dropbox/PaperEdits/Paper/Figures/")
covmat=readRDS("../Data/covmatAug13withED.rds")
z.stat=read.table("../Data/maxz.txt")
names=colnames(z.stat)
pis=readRDS("../Data/pisAug13withED.rds")$pihat
pi.mat=matrix(pis,ncol=54,nrow=22,byrow = T)
library(gplots)
library(ggplot2)
library('colorRamps')
#install.packages("fields")
library(fields)
k=3
hclust.2=function (d, method = "average", members = NULL) {hclust(d, method, members)}
      x=cov2cor(covmat[[k]])
x[x<0]=0
      colnames(x)=names
      rownames(x)=names
\# h=heatmap.2((x),symm=TRUE,dendrogram="none",density="none",trace="none",col=blue2green(256),main=past
\# heatmap.2((x[h\$rowInd, h\$rowInd]), Colv=FALSE, Rowv=FALSE, symm=TRUE, dendrogram="none", density="none", training for the strength of th
h=heatmap.2((x), #symm=TRUE,
                                       dendrogram="none", density="none", trace="none", col=blue2green(256), main=paste0("Cov2CorU", k,
```



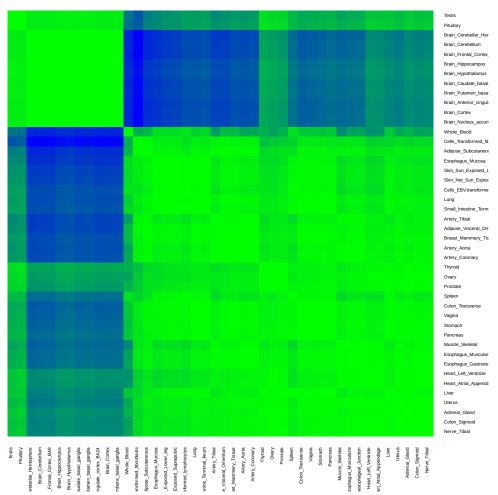
Cov2CorU3pihat=0.67



heatmap.2((x[h\$rowInd,h\$rowInd]),Colv=FALSE,Rowv=FALSE,symm=TRUE,dendrogram="none",density="none",trace



Cov2CorU3pihat=0.67



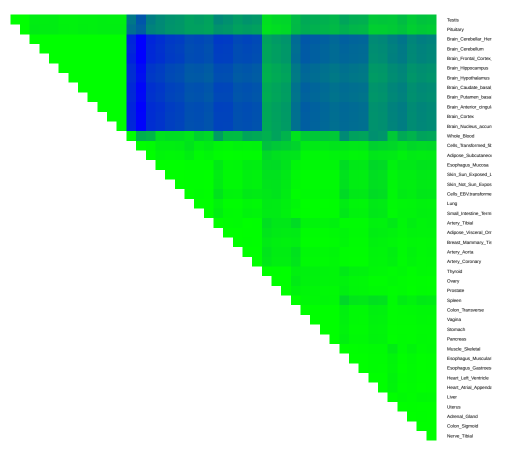
#write.table(h\$rowInd, "uk3rowindices.txt")

heatmap:

```
col=blue2green(256),
main=paste0("Cov2CorUk3"),
cexRow=0.5,cexCol=0.5,cex.main=0.5,labCol="")
```

Color Key 0 0.2 0.4 0.6 0.8 1 Value

Cov2CorUk3



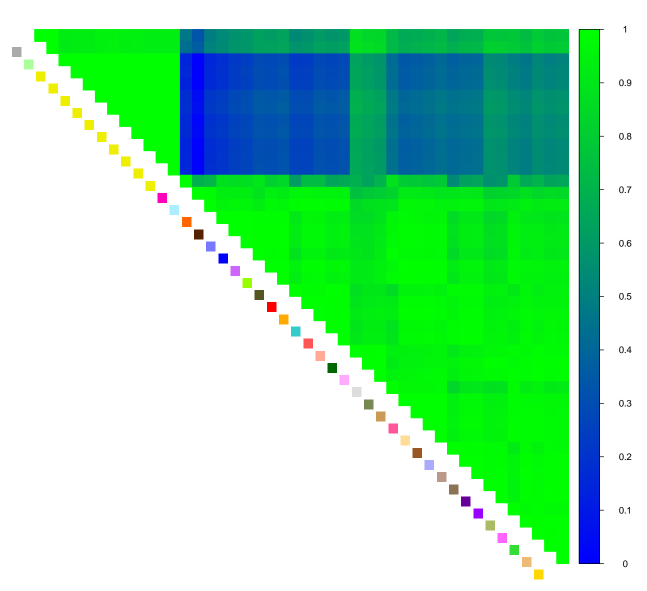
```
missing.tissues=c(7,8,19,20,24,25,31,34,37)
color.gtex=read.table("../Data/GTExColors.txt",sep = '\t', comment.char = '')[-missing.tissues,]
col = as.character(color.gtex[,2])
library('corrplot')

colnames(x)=NULL
rownames(x)=rep(".",44)
corrplot((x[h$rowInd,h$rowInd]),type="upper",cl.lim=c(0,1),tl.col=col[h$rowInd],tl.cex=8,method="color"
```

```
## Warning in ind1:ind2: numerical expression has 2 elements: only the first
## used
```

Warning in ind1:ind2: numerical expression has 2 elements: only the first
used





Let's try with different ordering:

```
#
# all.tissue.order=read.table("../../Dropbox/alltissueorder.txt")[,1]
#
# x=cov2cor(covmat[[k]])
# x[x<0]=0
#</pre>
```

```
#
              rownames(x) = names
 #
 #
\# smat=x[(all.tissue.order),(all.tissue.order)]
# smat[lower.tri(smat)] <- NA</pre>
#
# heatmap.2(smat, #symm=TRUE,
                                         Rowv=FALSE, Colv=FALSE,
#
#
                                          dendrogram="none", density="none", trace="none", #col=redblue,
 #
                                          col=blue2green(256),
                                          main=paste0("Cov2CortUk3"),
 #
 #
                                          cexRow=0.5, cexCol=0.5, cex.main=0.5)
 #
 #
 #
 #
 #
 #
 #
#
 #
#
#
# Eigenplots:
# ```{r eigenplots}
# missing.tissues=c(7,8,19,20,24,25,31,34,37)
\# color.gtex=read.table("../Data/GTExColors.txt",sep = '\t', comment.char = '')[-missing.tissues,]
# names=colnames(z.stat)
# vold=svd(covmat[[k]])$v;u=svd(covmat[[k]])$u
# v=vold[h$rowInd,]##shuffle so correct order
 # names=names[h$rowInd]
# color.qtex=color.qtex[h$rowInd,]
# for(j in 1:3){
\# \ barplot(v[,j]/v[,j][which.max(abs(v[,j]))], names="", cex.names=0.5, las=2, main=paste0("EigenVector",j,", one of the content of the co
```

#

colnames(x) = names

Note that the echo = FALSE parameter was added to the code chunk to prevent printing of the R code that generated the plot.