Figure 1 for Deleterious Alleles paper

To reproduce the figures, we should set pvpDiallel as the current path, i.e. setwd("~/Documents/Github/pvpDiallel/"). And then use knitr package to knit a pdf file. Or simplely click Knit PDF icon in RStudio. Note, to generate each panel into seperate pdf files, we should turn getpdf into TRUE (i.e. getpdf=TRUE) when calling the plotting functions.

First of all, determine fond size:

```
fs <- 1.3 # times bigger than default
getpdf <- TRUE # get figures in seperated pdf [TRUE] or not [FALSE]
```

Figure 1a

```
plotH2 <- function(getpdf=TRUE, outfile, ...){</pre>
 H2 <- read.csv("data/DIalleleHeritability.csv")</pre>
 h <- read.csv("cache/loh_pBPHmax_median.csv")</pre>
 H2 <- merge(H2, h, by.x="Traits", by.y="trait")
 H2 <- H2[order(H2$h), ]</pre>
  barplot(H2[,4], names.arg = H2$Traits, ...)
  if(getpdf == TRUE){
    pdf(outfile, width=6, height=6)
    barplot(H2[,4], names.arg = H2$Traits, ...)
    box()
    dev.off()
 }
}
plotH2(getpdf=getpdf, outfile="graphs/Fig1a_v3.pdf",
       ylim=c(0, 1), col="antiquewhite3", ylab="Heritability",
          cex.axis=fs, cex.names=fs, cex.lab=fs)
```

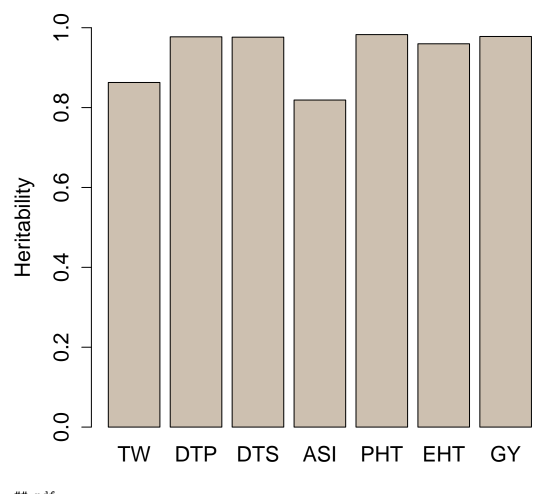


Figure 1b

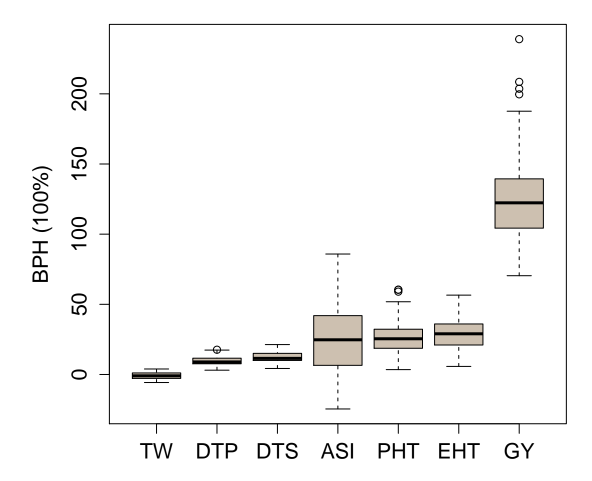


Figure 1c

```
plotReg <- function(getpdf=TRUE, outfile, ...){</pre>
  snptab <- read.csv("cache/daf_gerp2.csv")</pre>
  snptab <- snptab[order(snptab$GERP2),]</pre>
 plx <- predict(loess(snptab$meandaf ~ snptab$GERP2), se=T)</pre>
  x <- snptab$GERP2
 y <- snptab$meandaf
 plot(x, y, ...)
  lines(snptab$GERP2, plx$fit, col="cornflowerblue", lwd=2)
  lines(snptab$GERP2, plx$fit - qt(0.975,plx$df)*plx$se, lty=2, lwd=2, col="black")
  lines(snptab$GERP2, plx$fit + qt(0.975,plx$df)*plx$se, lty=2, lwd=2, col="black")
  if(getpdf == TRUE){
    pdf(outfile, width=6, height=6)
    plot(x, y, ...)
    lines(snptab$GERP2, plx$fit, col="cornflowerblue", lwd=2)
    lines(snptab$GERP2, plx$fit - qt(0.975,plx$df)*plx$se, lty=2, lwd=2, col="black")
    lines(snptab$GERP2, plx$fit + qt(0.975,plx$df)*plx$se, lty=2, lwd=2, col="black")
```

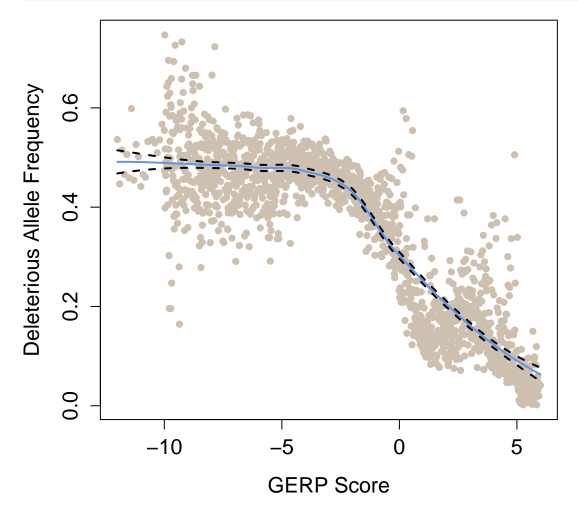


Figure 1d

To run this chunk, we need to install beanplot package.

```
library("beanplot")
plotbeans <- function(getpdf=TRUE, outfile, ...){

res <- read.csv("cache/mgerp_cm.csv")
  cutoff <- quantile(res$gen, c(0.25, 0.5, 0.75))
  res$sq <- 0</pre>
```

```
res[res$gen < cutoff[1], ]$sq <- 1</pre>
  res[res$gen >= cutoff[1] & res$gen < cutoff[2], ]$sq <- 2
  res[res$gen >= cutoff[2] & res$gen < cutoff[3], ]$sq <- 3</pre>
  res[res$gen >= cutoff[3], ]$sq <- 4
  beanplot(mgerp ~ sq, data = res, kernel="cosine", ll = 0.04, cex=fs, side = "no", cut=10, ylim=c(0.5,
         border = NA, col=list("#cd5b45", "antiquewhite3", "antiquewhite3", "antiquewhite3"),
         xaxt="n", ...)
  axis(side = 1, at = 1:4, labels = c("<25\", "25-50\", "50-75\", ">75\"), cex.axis=fs)
  if(getpdf == TRUE){
    pdf(outfile, width=6, height=6)
    beanplot(mgerp ~ sq, data = res, kernel="cosine", ll = 0.04, cex=1.5, side = "no", cut=10, ylim=c(0
         border = NA, col=list("#cd5b45", "antiquewhite3", "antiquewhite3", "antiquewhite3"),
         xaxt="n", ...)
    axis(side =1, at =1:4, labels =c("<25%", "25-50%", "50-75%", ">75%"), cex.axis=1.3)
    dev.off()
  }
}
plotbeans(getpdf=getpdf, outfile="graphs/Fig1d_v3.pdf",
          ylab="GERP Score", xlab="Quantiles of Recombination Rate (cM/Mb)",
          cex.axis=fs, cex.lab=fs)
```

log="y" selected
log="y" selected

