## Figure 2 of the GERP paper

```
knitr::opts_knit$set(root.dir=normalizePath('../'))
```

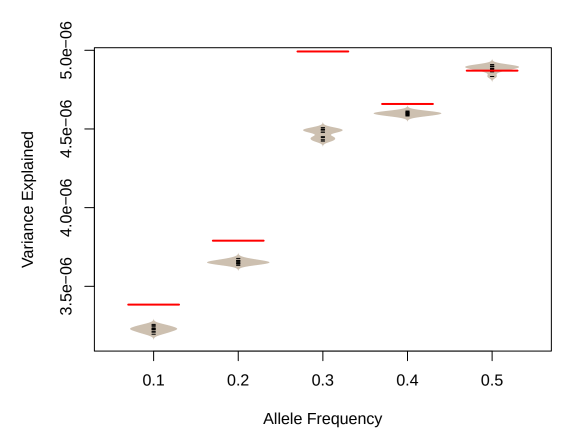
To reproduce the figures, we should set GERP-diallel/ as the root path, i.e. setwd("~/Documents/Github/GERP-diallel/"). And then use knitr package to get a pdf file. Or simplely click Knit PDF icon on RStudio. Note, to produce panels into seperate pdf files, we should set getpdf into TRUE (i.e. getpdf=TRUE) when calling the plotting functions.

First of all, determine fond size and set the getpdf option:

```
#par(mar=c(5,4,4,2))
par(font=2, font.lab=2, font.axis=2)
fs = 1.6 # times bigger than default
fsize = 16 # absolute font size
getpdf = TRUE # get figures in seperated pdf [TRUE] or not [FALSE]
```

## Figure 2a

```
plot_fig2a <- function(outfile, getpdf){</pre>
  library("beamplot")
  res <- read.csv("cache/persnp_var_perse.csv")</pre>
  myd <- subset(res, sample != 0)</pre>
  myd0 <- subset(res, sample == 0)
  #### for trait perse
  beamplot(nvar ~ frq1, data = myd, ll = 0.04, cex=1.5, border = NA,
         what=c(0, 1, 0, 1), col=c(c("antiquewhite3", "black")), cex=fs,
         xlab="Allele Frequency", ylab="Variance Explained" )
  for(i in 1:nrow(myd0)){
    lines(x=c(i-0.3, i+0.3), y=c(myd0$nvar[i], myd0$nvar[i]), lwd=2, col="red")
  }
  if(getpdf == TRUE){
    pdf(outfile, width=5, height=5)
    #### for trait perse
    par(font=2, font.lab=2, font.axis=2, cex.lab=fs)
    beamplot(nvar ~ frq1, data = myd, ll = 0.04, border = NA,
            what=c(0, 1, 0, 1), col=c(c("antiquewhite3", "black")),
            xlab="Allele Frequency", ylab="Variance Explained" )
    for(i in 1:nrow(myd0)){
      lines(x=c(i-0.3, i+0.3), y=c(myd0$nvar[i], myd0$nvar[i]), lwd=2, col="red")
    }
    dev.off()
  }
}
#######
plot_fig2a(outfile="graphs/Fig2a_del_snps.pdf", getpdf)
```



## pdf ## 2

## Figure 2b

```
library(ggplot2)

## Warning: package 'ggplot2' was built under R version 3.2.3

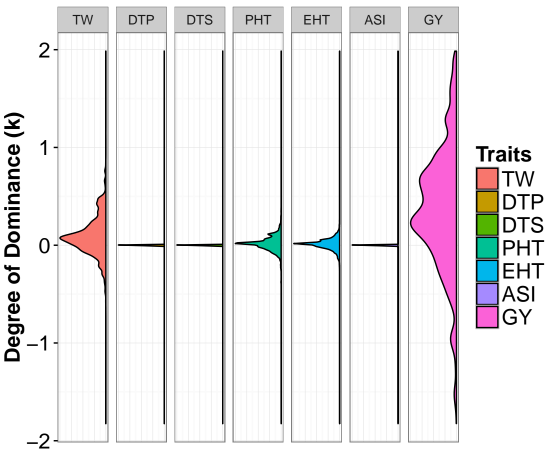
library(reshape2)
source("lib/multiplot.R")

plot_eff <- function(outfile, getpdf){
    res1 <- read.csv("cache/gblup_var_updated.csv")
    res2 <- read.csv("cache/kval_perse_5x.csv")
    med <- read.csv("cache/loh_pMPH_median.csv")
    bymed <- med[order(med$h),]

cols <- c("#f6546a", "#daa520", "#00ff00", "#66cdaa", "#3b5998", "#8a2be2", "#ff00ff")

#######
theme_set(theme_grey(base_size = 18))

res2$trait <- toupper(res2$trait)
    res2$trait <- factor(res2$trait, levels=bymed$trait)
    p2 <- ggplot(data=res2) +</pre>
```



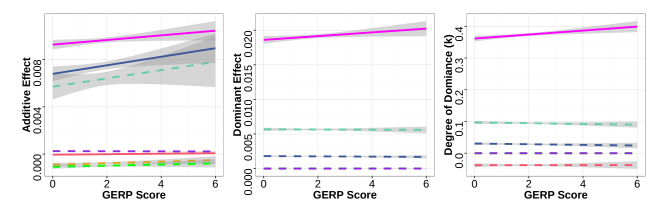
```
if(getpdf){
    pdf("graphs/Fig2b_var.pdf", width=8, height=4)
    p
    dev.off()
}
```

```
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## 2
```

## Figure 2c-e

```
#library(wesanderson)
library(data.table)
##
## Attaching package: 'data.table'
## The following objects are masked from 'package:reshape2':
##
##
       dcast, melt
source("lib/multiplot.R")
plot_adk_gerp <- function(outfile, getpdf){</pre>
  dat <- fread("largedata/gerpdat0x.csv", data.table=FALSE)</pre>
  dat$trait <- toupper(dat$trait)</pre>
  med <- read.csv("cache/loh_pMPH_median.csv")</pre>
  out <- read.csv("cache/eff_adk_0x.csv")</pre>
  out <- merge(med, out, by="trait")</pre>
  out <- out[order(out$h),]</pre>
  #cols <- wes_palette(7, name = "Zissou", type = "continuous")</pre>
  cols <- c("#f6546a", "#daa520", "#00ff00", "#66cdaa", "#3b5998", "#8a2be2", "#ff00ff")
  theme_set(theme_grey(base_size = 18))
  getlty <- function(df, eff, cutoff=0.05){</pre>
    df$1 <- 2
    if(nrow(df[df[, eff] < cutoff, ]) >0) df[df[, eff] < cutoff, ]$1 <- 1</pre>
  }
  lty1 <- getlty(df=out, eff="effa", cutoff=0.05)$1</pre>
  p1 <- ggplot(dat, aes(x=RS, y=Effect_A, colour=factor(trait, levels=out$trait),
                         linetype=factor(trait, levels=out$trait))) +
    labs(colour="Traits") +
    theme_bw() +
    xlab("GERP Score") +
    ylab("Additive Effect") +
    scale_color_manual(values=cols) +
    scale_linetype_manual(values=lty1) +
    guides(colour=FALSE, linetype=FALSE) +
    geom_smooth(method="gam", size=1.3) +
    theme(axis.text.y = element_text(angle = 90, hjust = 1),
          axis.text=element text(size=fsize),
          axis.title=element_text(size=fsize, face="bold"),
          legend.title = element_text(size=fsize, face="bold"),
          legend.text = element_text(size=fsize))
```

```
lty2 <- getlty(df=out, eff="effd", cutoff=0.05)$1</pre>
  p2 <- ggplot(dat, aes(x=RS, y=Effect_D, colour=factor(trait, levels=out$trait),</pre>
                        linetype=factor(trait, levels=out$trait))) +
   labs(colour="Traits") +
   theme bw() +
   xlab("GERP Score") +
   ylab("Dominant Effect") +
    scale color manual(values=cols) +
    scale linetype manual(values=lty2) +
    guides(colour=FALSE, linetype=FALSE) +
    geom_smooth(method="gam", size=1.3) + # Add linear regression line
    theme(axis.text.y = element_text(angle = 90, hjust = 1),
          axis.text=element_text(size=fsize),
          axis.title=element_text(size=fsize, face="bold"),
          legend.title = element_text(size=fsize, face="bold"),
          legend.text = element_text(size=fsize))
  lty3 <- getlty(df=out, eff="effk", cutoff=0.05)$1</pre>
  p3 <- ggplot(dat, aes(x=RS, y=k, colour=factor(trait, levels=out$trait),
                        linetype=factor(trait, levels=out$trait))) +
   labs(colour="Traits") +
   theme bw() +
   xlab("GERP Score") +
   ylab("Degree of Domiance (k)") +
    scale color manual(values=cols) +
   scale_linetype_manual(values=lty3) +
   guides(colour=FALSE, linetype=FALSE) +
   theme(axis.text.y = element_text(angle = 90, hjust = 1),
          axis.text=element_text(size=fsize),
          axis.title=element_text(size=fsize, face="bold"),
          legend.title = element_text(size=fsize, face="bold"),
          legend.text = element_text(size=fsize)) +
    geom_smooth(method="gam", size=1.3) # Add linear regression line
  multiplot(p1, p2, p3, cols=3)
  if(getpdf == TRUE){
   pdf(outfile, width=13, height=4)
   multiplot(p1, p2, p3, cols=3)
   dev.off()
 }
}
####
plot_adk_gerp(outfile="graphs/Fig2c-e_adk.pdf", getpdf)
##
Read 23.6% of 2376528 rows
Read 56.4% of 2376528 rows
Read 90.0% of 2376528 rows
Read 2376528 rows and 11 (of 11) columns from 0.265 GB file in 00:00:05
```



## pdf ## 2