## main exons

September 5, 2021

## 1 eQTL boxplot

This is script ported from python to fix unknown plotting error.

```
[1]: suppressPackageStartupMessages({
    library(tidyverse)
    library(ggpubr)
})
```

#### 1.1 Functions

```
[2]: feature = "exons"
```

#### 1.1.1 Cached functions

```
[3]: get_eqtl_df <- function(){</pre>
        eGenes_file = pasteO('/ceph/projects/v4_phase3_paper/analysis/eqtl_analysis/
      →all/',
                             feature, '/expression_gct/prepare_expression/

¬fastqtl_permutation/',
                             '_m/Brainseq_LIBD.genes.txt.gz')
        eGenes = data.table::fread(eGenes_file) %>%
             select(gene_id, variant_id, maf, slope, slope_se, pval_nominal, qval)__
      →%>%
             arrange(qval)
        return(eGenes)
     memEQTL <- memoise::memoise(get_eqtl_df)</pre>
     get_residualized_df <- function(){</pre>
         expr_file = paste0("/ceph/projects/v4_phase3_paper/analysis/eqtl_analysis/
     ⇔all/",
                           feature, "/expression_gct/covariates/
     feature, "_residualized_expression.csv")
        return(data.table::fread(expr_file) %>% column to rownames("gene_id"))
     memRES <- memoise::memoise(get_residualized_df)</pre>
```

#### 1.1.2 Simple functions

```
[4]: feature map <- function(feature){
         return(list("genes"="Gene", "transcripts"= "Transcript",
                     "exons"= "Exon", "junctions"= "Junction")[[feature]])
     }
     get_geno_annot <- function(){</pre>
         return(memSNPs() %>% select(CHR, SNP, POS, COUNTED, ALT))
     }
     get_snps_df <- function(){</pre>
         return(memSNPs() %>% select("SNP", starts_with("Br")))
     }
     letter_snp <- function(number, a0, a1){</pre>
         if(is.na(number)){ return(NA) }
         if( length(a0) == 1 & length(a1) == 1){
             seps = ""; collapse=""
         } else {
             seps = " "; collapse=NULL
         return(paste(paste0(rep(a0, number), collapse = collapse),
                      pasteO(rep(a1, (2-number)), collapse = collapse), sep=seps))
     }
     get_snp_df <- function(variant_id, gene_id){</pre>
         zz = get_geno_annot() %>% filter(SNP == variant_id)
         xx = get_snps_df() %>% filter(SNP == variant_id) %>%
             column_to_rownames("SNP") %>% t %>% as.data.frame %>%
             rownames_to_column("BrNum") %>% mutate(COUNTED=zz$COUNTED, ALT=zz$ALT)_
      →%>%
             rename("SNP"=all_of(variant_id))
         yy = memRES()[gene_id, ] %>% t %>% as.data.frame %>%
             rownames_to_column("BrNum")
         ## Annotated SNPs
         letters = c()
```

```
for(ii in seq_along(xx$COUNTED)){
        a0 = xx$COUNTED[ii]; a1 = xx$ALT[ii]; number = xx$SNP[ii]
        letters <- append(letters, letter_snp(number, a0, a1))</pre>
    xx = xx %>% mutate(LETTER=letters, ID=paste(SNP, LETTER, sep="\n"))
    df = inner_join(xx, yy, by="BrNum") %>% mutate_if(is.character, as.factor)
    return(df)
memDF <- memoise::memoise(get snp df)</pre>
save_ggplots <- function(fn, p, w, h){</pre>
    for(ext in c('.pdf', '.png', '.svg')){
        ggsave(paste0(fn, ext), plot=p, width=w, height=h)
    }
}
get_biomart_df <- function(){</pre>
    biomart = data.table::fread("../_h/biomart.csv")
memMART <- memoise::memoise(get_biomart_df)</pre>
get_gene_symbol <- function(gene_id){</pre>
    ensemblID = gsub("\\..*", "", gene_id)
    geneid = memMART() %>% filter(ensembl_gene_id == gsub("\\..*", "", gene_id))
    if(dim(geneid)[1] == 0){
        return("")
    } else {
        return(geneid$external gene name)
    }
}
plot_simple eqtl <- function(fn, gene_id, variant_id, eqtl_annot, prefix){</pre>
    bxp = memDF(variant_id, gene_id) %>%
        ggboxplot(x="ID", y=gene_id, fill="red", add="jitter", xlab="",
                  ylab="Residualized Expression", outlier.shape=NA,
                  add.params=list(alpha=0.5), alpha=0.4,
                  ggtheme=theme_pubr(base_size=20, border=TRUE)) +
        font("xy.title", face="bold") +
        ggtitle(paste(prefix, eqtl annot, sep='\n')) +
        theme(plot.title = element_text(hjust = 0.5, face="bold"))
    print(bxp)
    save_ggplots(fn, bxp, 7, 7)
}
```

#### 1.1.3 GWAS plots

```
[5]: get_gwas_snps <- function(){</pre>
         gwas_snp_file = paste0('../../summary_table/_m/Brainseq_LIBD_caudate',
                                  '_4features_PGC2.signifpairs.txt.gz')
         gwas_df = data.table::fread(gwas_snp_file) %>% filter(Type ==_
      →feature_map(feature)) %>%
             select(c("variant_id", "gene_id", "rsid", "hg38chrc", "OR", "SE", "P", 
      \hookrightarrow "A1".
                       "A2", "pgc2_a1_same_as_our_counted", "is_index_snp")) %>%
             distinct() %>% arrange(P)
         return(gwas_df)
     memGWAS <- memoise::memoise(get_gwas_snps)</pre>
     get_gwas_snp <- function(variant){</pre>
         return(memGWAS() %>% filter(variant id == variant))
     }
     get_risk_allele <- function(variant){</pre>
         gwas_snp = get_gwas_snp(variant)
         if(gwas_snp$OR > 1){
             ra = gwas_snp$A1
         }else{
             ra = gwas_snp$A2
         return(ra)
     }
     get_eqtl_gwas_df <- function(){</pre>
         return(memEQTL() %>% inner_join(memGWAS(), by="variant_id"))
     }
     get_gwas_ordered_snp_df <- function(variant_id, gene_id,__
      →pgc2_a1_same_as_our_counted, OR){
         df = memDF(variant_id, gene_id)
         if(pgc2_a1_same_as_our_counted){
             if(OR < 1) { df = df %>% mutate(SNP = 2-SNP, ID=paste(SNP, LETTER, I
      →sep="\n")) }
         } else {
             if(OR > 1){ df = df %>% mutate(SNP = 2-SNP, ID=paste(SNP, LETTER,__
      →sep="\n")) }
         return(df)
     }
     plot_gwas_eqtl <- function(fn, gene_id, variant_id, eqtl_annot,</pre>
```

## 1.2 Plot eQTL

#### 1.2.1 DRD2 plot

```
[7]: drd2_short = "ENST00000346454.7"; drd2_long = "ENST00000362072.7"

exons_df = data.table::fread("../../../DE_gwas_eqtl_integration/_h/exons.

→csv") %>%

filter(transcript_id %in% c(drd2_short, drd2_long)) %>%

select(gene_id, gene_name, exon_id, brainseq_exon_id) %>%

distinct(brainseq_exon_id, .keep_all=TRUE)

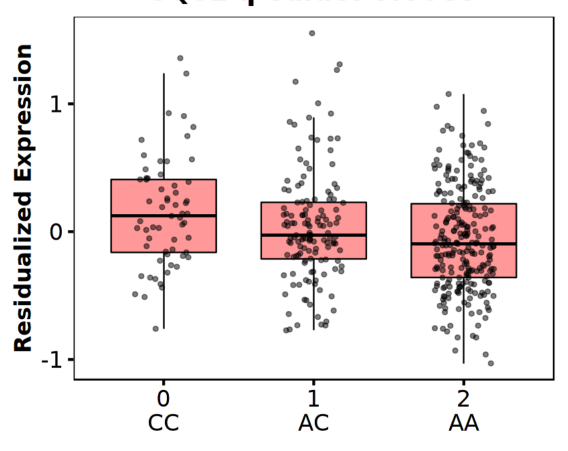
all_drd2_exons = exons_df$brainseq_exon_id

drd2_df = memEQTL() %>% filter(gene_id %in% all_drd2_exons)

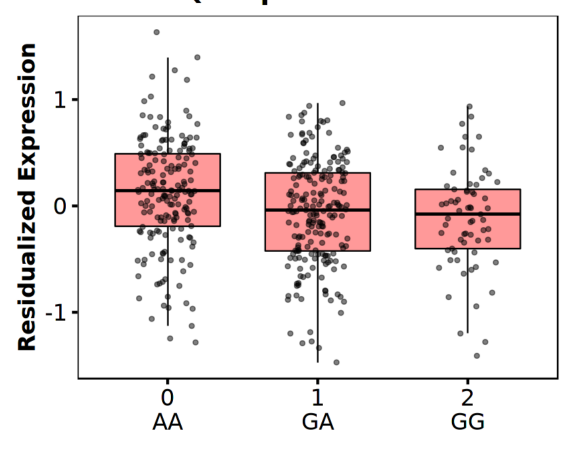
drd2_df
```

```
gene id variant id
                                                                                          pval nominal
                                                           maf
                                                                     slope
                                                                               slope se
                       <chr>
                                <chr>
                                                           <dbl>
                                                                     <dbl>
                                                                                <dbl>
                                                                                          <dbl>
                      e667157
                                chr11:113336077:A:C
                                                           0.291196
                                                                     0.162631
                                                                               0.0355340
                                                                                          6.28127e-06
    A data.table: 4 \times 7
                      e667159
                                chr11:113532227:G:A
                                                           0.374718
                                                                     0.146535
                                                                               0.0380270
                                                                                          1.35279e-04
                      e667153
                                chr11:113512525:AGTTT:A
                                                           0.198646
                                                                     -0.110192
                                                                               0.0341356
                                                                                          1.34726e-03
                      e667158
                                chr11:113281068:C:T
                                                           0.118510
                                                                    0.138094
                                                                               0.0432721
                                                                                          1.52628e-03
[8]: for(x in seq_along(drd2_df$gene_id)){
         anno = get_drd2_exon_annotation(drd2_df$gene_id[x])
         en = gsub(" ", "_", anno)
         ids = exons_df %>% filter(brainseq_exon_id == drd2_df$gene_id[x]) %>%_
      ⇔select("exon_id")
         fn = paste("drd2_eqtl", en, sep="_")
         eqtl_annot = paste("eQTL q-value:", signif(drd2_df$qval[x], 2))
         prefix = paste(anno, ids$exon_id, sep='\n')
         plot_simple_eqtl(fn, drd2_df$gene_id[x], drd2_df$variant_id[x], eqtl_annot,__
      \rightarrowprefix)
         #print(eqtl_annot)
```

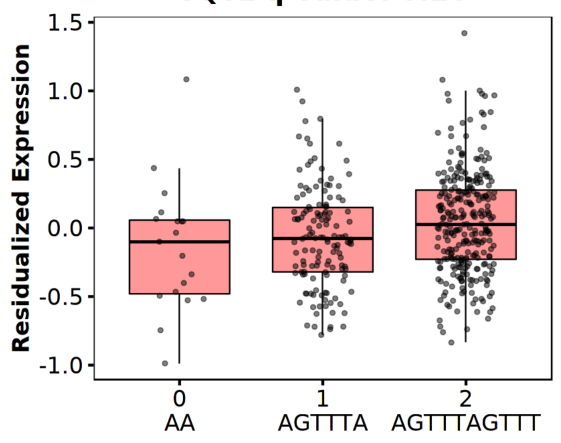
DRD2 exon 7 ENSE00001247012.2 eQTL q-value: 0.0089



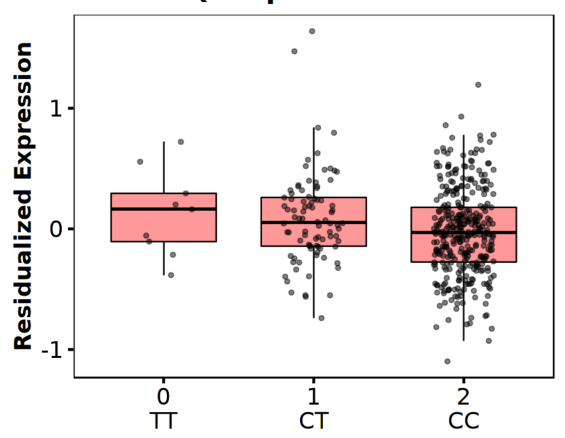
DRD2 exon 1L ENSE00002282098.1 eQTL q-value: 0.087



DRD2 exon 2 ENSE00003665261.1 eQTL q-value: 0.26



# DRD2 exon 8 ENSE00002210277.1 eQTL q-value: 0.27



#### 1.2.2 GWAS association

```
[9]: memGWAS() %>% filter(gene_id %in% all_drd2_exons)
                                                                  OR
                                                                          SE
                                                                                   Ρ
                                                                                           A1
                                     gene_id
                                                       hg38chrc
                                                                                                    A2
                         variant_id
                                              rsid
    A data.table: 0 \times 11
                         <chr>
                                     <chr>
                                               <chr>
                                                       <chr>
                                                                  <dbl>
                                                                          <dbl>
                                                                                   < dbl >
                                                                                            <chr>
                                                                                                    <chr>
```

## 1.3 Session Info

```
[10]: Sys.time()
   proc.time()
   options(width = 120)
   sessioninfo::session_info()
```

## [1] "2021-09-05 15:54:12 EDT"

user system elapsed 1702.624 615.663 336.513

Session info setting value

version R version 4.0.3 (2020-10-10)

os Arch Linux

system x86\_64, linux-gnu

ui X11 language (EN)

collate en\_US.UTF-8
ctype en\_US.UTF-8
tz America/New\_York

date 2021-09-05

## Packages

| package    | * | version  | date       | lib | sourc | е  |        |
|------------|---|----------|------------|-----|-------|----|--------|
| abind      |   | 1.4-5    | 2016-07-21 | [1] | CRAN  | (R | 4.0.2) |
| assertthat |   | 0.2.1    | 2019-03-21 | [1] | CRAN  | (R | 4.0.2) |
| backports  |   | 1.2.1    | 2020-12-09 | [1] | CRAN  | (R | 4.0.2) |
| base64enc  |   | 0.1-3    | 2015-07-28 | [1] | CRAN  | (R | 4.0.2) |
| broom      |   | 0.7.9    | 2021-07-27 | [1] | CRAN  | (R | 4.0.3) |
| cachem     |   | 1.0.6    | 2021-08-19 | [1] | CRAN  | (R | 4.0.3) |
| Cairo      |   | 1.5-12.2 | 2020-07-07 | [1] | CRAN  | (R | 4.0.2) |
| car        |   | 3.0-11   | 2021-06-27 | [1] | CRAN  | (R | 4.0.3) |
| carData    |   | 3.0-4    | 2020-05-22 | [1] | CRAN  | (R | 4.0.2) |
| cellranger |   | 1.1.0    | 2016-07-27 | [1] | CRAN  | (R | 4.0.2) |
| cli        |   | 3.0.1    | 2021-07-17 | [1] | CRAN  | (R | 4.0.3) |
| colorspace |   | 2.0-2    | 2021-06-24 | [1] | CRAN  | (R | 4.0.3) |
| crayon     |   | 1.4.1    | 2021-02-08 | [1] | CRAN  | (R | 4.0.3) |
| curl       |   | 4.3.2    | 2021-06-23 | [1] | CRAN  | (R | 4.0.3) |
| data.table |   | 1.14.0   | 2021-02-21 | [1] | CRAN  | (R | 4.0.3) |
| DBI        |   | 1.1.1    | 2021-01-15 | [1] | CRAN  | (R | 4.0.2) |
| dbplyr     |   | 2.1.1    | 2021-04-06 | [1] | CRAN  | (R | 4.0.3) |
| digest     |   | 0.6.27   | 2020-10-24 | [1] | CRAN  | (R | 4.0.2) |
| dplyr      | * | 1.0.7    | 2021-06-18 | [1] | CRAN  | (R | 4.0.3) |
| ellipsis   |   | 0.3.2    | 2021-04-29 | [1] | CRAN  | (R | 4.0.3) |
| evaluate   |   | 0.14     | 2019-05-28 | [1] | CRAN  | (R | 4.0.2) |
| fansi      |   | 0.5.0    | 2021-05-25 | [1] | CRAN  | (R | 4.0.3) |
| farver     |   | 2.1.0    | 2021-02-28 | [1] | CRAN  | (R | 4.0.3) |
| fastmap    |   | 1.1.0    | 2021-01-25 | [1] | CRAN  | (R | 4.0.2) |
| forcats    | * | 0.5.1    | 2021-01-27 | [1] | CRAN  | (R | 4.0.2) |
| foreign    |   | 0.8-80   | 2020-05-24 | [2] | CRAN  | (R | 4.0.3) |
| fs         |   | 1.5.0    | 2020-07-31 | [1] | CRAN  | (R | 4.0.2) |
| generics   |   | 0.1.0    | 2020-10-31 | [1] | CRAN  | (R | 4.0.2) |
| ggplot2    | * | 3.3.5    | 2021-06-25 | [1] | CRAN  | (R | 4.0.3) |
| ggpubr     | * | 0.4.0    | 2020-06-27 | [1] | CRAN  | (R | 4.0.2) |

```
0.6.2
                        2021-06-14 [1] CRAN (R 4.0.3)
ggsignif
glue
               1.4.2
                        2020-08-27 [1] CRAN (R 4.0.2)
gtable
              0.3.0
                        2019-03-25 [1] CRAN (R 4.0.2)
haven
              2.4.3
                        2021-08-04 [1] CRAN (R 4.0.3)
hms
              1.1.0
                        2021-05-17 [1] CRAN (R 4.0.3)
              0.5.2
                        2021-08-25 [1] CRAN (R 4.0.3)
htmltools
httr
               1.4.2
                        2020-07-20 [1] CRAN (R 4.0.2)
IRdisplay
               1.0
                        2021-01-20 [1] CRAN (R 4.0.2)
                        2021-05-11 [1] CRAN (R 4.0.3)
IRkernel
              1.2
jsonlite
              1.7.2
                        2020-12-09 [1] CRAN (R 4.0.2)
              0.4.2
                        2020-10-20 [1] CRAN (R 4.0.2)
labeling
              1.0.0
                        2021-02-15 [1] CRAN (R 4.0.3)
lifecycle
lubridate
              1.7.10
                        2021-02-26 [1] CRAN (R 4.0.3)
magrittr
              2.0.1
                        2020-11-17 [1] CRAN (R 4.0.2)
memoise
              2.0.0
                        2021-01-26 [1] CRAN (R 4.0.2)
              0.1.8
modelr
                        2020-05-19 [1] CRAN (R 4.0.2)
munsell
              0.5.0
                        2018-06-12 [1] CRAN (R 4.0.2)
              4.2.4
                        2021-06-16 [1] CRAN (R 4.0.3)
openxlsx
              0.3 - 5
                        2021-02-10 [1] CRAN (R 4.0.3)
pbdZMQ
               1.6.2
                        2021-07-29 [1] CRAN (R 4.0.3)
pillar
pkgconfig
              2.0.3
                        2019-09-22 [1] CRAN (R 4.0.2)
            * 0.3.4
                        2020-04-17 [1] CRAN (R 4.0.2)
purrr
R.methodsS3
              1.8.1
                        2020-08-26 [1] CRAN (R 4.0.3)
              1.24.0
                        2020-08-26 [1] CRAN (R 4.0.3)
R.oo
R.utils
              2.10.1
                        2020-08-26 [1] CRAN (R 4.0.3)
              2.5.1
R6
                        2021-08-19 [1] CRAN (R 4.0.3)
               1.0.7
                        2021-07-07 [1] CRAN (R 4.0.3)
Rcpp
readr
            * 2.0.1
                        2021-08-10 [1] CRAN (R 4.0.3)
                        2019-03-13 [1] CRAN (R 4.0.2)
readxl
               1.3.1
              1.1.3
                        2021-01-21 [1] CRAN (R 4.0.2)
repr
              2.0.1
                        2021-08-05 [1] CRAN (R 4.0.3)
reprex
rio
              0.5.27
                        2021-06-21 [1] CRAN (R 4.0.3)
rlang
              0.4.11
                        2021-04-30 [1] CRAN (R 4.0.3)
              0.7.0
                        2021-02-13 [1] CRAN (R 4.0.3)
rstatix
              0.13
                        2020-11-12 [1] CRAN (R 4.0.2)
rstudioapi
rvest
               1.0.1
                        2021-07-26 [1] CRAN (R 4.0.3)
scales
               1.1.1
                        2020-05-11 [1] CRAN (R 4.0.2)
sessioninfo
              1.1.1
                        2018-11-05 [1] CRAN (R 4.0.2)
                        2021-08-25 [1] CRAN (R 4.0.3)
stringi
              1.7.4
stringr
            * 1.4.0
                        2019-02-10 [1] CRAN (R 4.0.2)
              2.0.0
                        2021-02-20 [1] CRAN (R 4.0.3)
svglite
                        2021-05-11 [1] CRAN (R 4.0.3)
systemfonts
               1.0.2
tibble
            * 3.1.4
                        2021-08-25 [1] CRAN (R 4.0.3)
tidyr
            * 1.1.3
                        2021-03-03 [1] CRAN (R 4.0.3)
tidyselect
               1.1.1
                        2021-04-30 [1] CRAN (R 4.0.3)
tidyverse
            * 1.3.1
                        2021-04-15 [1] CRAN (R 4.0.3)
tzdb
              0.1.2
                        2021-07-20 [1] CRAN (R 4.0.3)
utf8
               1.2.2
                        2021-07-24 [1] CRAN (R 4.0.3)
```

| uuid  | 0.1-4 | 2020-02-26 [1] CRAN (R 4.0.2) |
|-------|-------|-------------------------------|
| vctrs | 0.3.8 | 2021-04-29 [1] CRAN (R 4.0.3) |
| withr | 2.4.2 | 2021-04-18 [1] CRAN (R 4.0.3) |
| xml2  | 1.3.2 | 2020-04-23 [1] CRAN (R 4.0.2) |
| zip   | 2.2.0 | 2021-05-31 [1] CRAN (R 4.0.3) |

- [1] /home/jbenja13/R/x86\_64-pc-linux-gnu-library/4.0 [2] /usr/lib/R/library