# main

October 1, 2021

# 1 Exploring the eigen values and correlation with phenotypes

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

# 1.1 Prepare data

## 1.1.1 Load phenotypes data

```
[2]: pheno = data.table::fread("../../../input/phenotypes/merged/_m/

→merged_phenotypes.csv")

pheno %>% head(2)
```

```
[3]: ancestry = data.table::fread("../../../input/ancestry_structure/structure.

out_ancestry_proportion_raceDemo_compare")
ancestry %>% head(2)
```

```
Afr
                                       Eur
                     id
                                                group
                     <chr>
                              <dbl>
                                       <dbl>
                                                <chr>
A data.table: 2 \times 4
                     Br2374
                              0.007
                                       0.993
                                                CAUC
                     Br1857
                              0.001
                                       0.999
                                                CAUC
```

# 1.1.2 Load eigen values

```
[4]: eigen = data.table::fread("../../_m/eigengenes.csv")
modules = eigen %>% select(-V1) %>% colnames
eigen %>% dim
eigen[1:2, 1:5]
```

1. 230 2. 38

```
MEblack
                                            MEblue
                                                         MEbrown
                    V1
                                                                      MEcyan
                             < dbl >
                                            <dbl>
                                                         <dbl>
                                                                       < dbl >
                    < chr >
A data.table: 2 \times 5
                    R12872
                             -0.040931317
                                           0.02966489
                                                         0.09745435
                                                                      -0.06366120
                                                                      0.03820389
                    R12873 -0.003001924
                                           -0.03115660
                                                         -0.01174513
```

#### 1.1.3 Merge data

```
[5]: dt = eigen %>% left_join(pheno, by="V1") %>%
         inner_join(ancestry, by=c("BrNum"="id"))
    dt %>% dim
    dt[1:2, 1:5]
```

#### 1. 230 2. 52

```
V1
                            MEblack
                                          MEblue
                                                       MEbrown
                                                                    MEcyan
                                                                    <dbl>
                   <chr>
                            <dbl>
                                          <dbl>
                                                       <dbl>
A data.table: 2 \times 5
                   R12872 -0.040931317
                                          0.02966489
                                                       0.09745435
                                                                    -0.06366120
                   R12873
                           -0.003001924
                                          -0.03115660
                                                       -0.01174513
                                                                    0.03820389
```

## 1.2 Examine correlation of modules with ancestry

#### 1.2.1 Linear model

```
[6]: pvals = c()
for(mod in modules){
    model = paste0("Eur ~ ", mod)
    res = anova(lm(model, data=dt))
    pvals = c(pvals, res[mod, "Pr(>F)"])
}
fdr <- p.adjust(pvals, method="fdr")
df1 = data.frame("Modules"=modules, "Pvalue"=pvals, "FDR"=fdr)
df1 %>% filter(`Pvalue` < 0.05)</pre>
```

```
[7]: df1 %>% mutate(Tissue="Caudate") %>% data.table::fwrite("eigen_correlation_ancestry.tsv", sep='\t')
```

#### 1.2.2 Pearson correlation

```
[8]: pvals = c(); est = c()
for(mod in modules){
    res = cor.test(dt[["Eur"]], dt[[mod]], method="pearson")
    pvals = c(pvals, res$p.value)
    est = c(est, res$estimate[[1]])
}
```

```
fdr <- p.adjust(pvals, method="fdr")
df2 = data.frame("Modules"=modules, "Rho"=est, "Pvalue"=pvals, "FDR"=fdr)
df2 %>% filter(Pvalue < 0.05)</pre>
```

```
FDR
                    Modules
                                 Rho
                                             Pvalue
                    <chr>
                                 <dbl>
                                             <dbl>
                                                           <dbl>
A data.frame: 3 \times 4 MEmagenta
                                             0.0425120843 0.38830793
                                 -0.1338802
                    MEskyblue
                                 -0.1593326
                                             0.0155758706 0.28815361
                    MEviolet
                                 -0.2376683
                                            0.0002757979 \quad 0.01020452
```

# 1.3 Clustering modules

#### 1.3.1 Eigengene clustering

**png:** 2

#### 1.3.2 clustering within significant module violet

```
[10]: load("../../ m/01.RData", verbose=TRUE)
      MEviolet = data.table::fread("../../_m/modules.csv", header=TRUE) %>%
          filter(module == "violet")
      violet.mat = datExpr %>% as.data.frame %>%
          select(any_of(MEviolet$V1)) %>% as.matrix
     Loading objects:
       datExpr
       sample_table
       datTraits
[11]: biomart_file = "../../../input/biomart/biomart.csv"
      biomart = data.table::fread(biomart_file)
      annot = data.frame("Geneid"=colnames(violet.mat),
                          "ensembl_gene_id"=gsub("\\..*", "", colnames(violet.mat)))__

√/<sub>0</sub>>%

          left_join(biomart, by="ensembl_gene_id")
      annot$external_gene_name %>% is.na %>% sum
[12]: colnames(violet.mat) <- annot$external_gene_name
```

**png:** 2

```
1.4 Reproducibility Information
[14]: Sys.time()
      proc.time()
      options(width = 120)
      sessioninfo::session_info()
     [1] "2021-10-01 13:00:44 EDT"
              system elapsed
      19.337
               1.607 21.291
      Session info
      setting value
      version R version 4.0.3 (2020-10-10)
               Arch Linux
      system
               x86_64, linux-gnu
               X11
      ui
      language (EN)
               en_US.UTF-8
      collate
               en_US.UTF-8
      ctype
      tz
               America/New_York
               2021-10-01
      date
      Packages
                                           lib source
      package
                     * version date
      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)
                       1.2.1
                                2020-12-09 [1] CRAN (R 4.0.2)
      backports
      base64enc
                       0.1-3
                                2015-07-28 [1] CRAN (R 4.0.2)
                       0.36.1
      BiocGenerics
                                2021-04-16 [1] Bioconductor
      broom
                       0.7.9
                                2021-07-27 [1] CRAN (R 4.0.3)
      Cairo
                       1.5-12.2 2020-07-07 [1] CRAN (R 4.0.2)
                                2021-06-27 [1] CRAN (R 4.0.3)
      car
                       3.0-11
      carData
                       3.0-4
                                2020-05-22 [1] CRAN (R 4.0.2)
                                2016-07-27 [1] CRAN (R 4.0.2)
      cellranger
                       1.1.0
                       0.4.13
                                2021-06-09 [1] CRAN (R 4.0.3)
      circlize
```

```
cli
                  3.0.1
                           2021-07-17 [1] CRAN (R 4.0.3)
                           2021-04-16 [1] CRAN (R 4.0.3)
clue
                  0.3 - 59
cluster
                  2.1.0
                           2019-06-19 [2] CRAN (R 4.0.3)
                  2.0-2
                           2021-06-24 [1] CRAN (R 4.0.3)
colorspace
ComplexHeatmap * 2.6.2
                           2020-11-12 [1] Bioconductor
                           2021-02-08 [1] CRAN (R 4.0.3)
crayon
                  1.4.1
curl
                  4.3.2
                           2021-06-23 [1] CRAN (R 4.0.3)
data.table
                  1.14.2
                           2021-09-27 [1] CRAN (R 4.0.3)
DBI
                           2021-01-15 [1] CRAN (R 4.0.2)
                  1.1.1
                           2021-04-06 [1] CRAN (R 4.0.3)
dbplyr
                  2.1.1
                  0.6.28
                           2021-09-23 [1] CRAN (R 4.0.3)
digest
                * 1.0.7
                           2021-06-18 [1] CRAN (R 4.0.3)
dplyr
                  0.3.2
                           2021-04-29 [1] CRAN (R 4.0.3)
ellipsis
                  0.14
                           2019-05-28 [1] CRAN (R 4.0.2)
evaluate
fansi
                  0.5.0
                           2021-05-25 [1] CRAN (R 4.0.3)
                  1.1.0
                           2021-01-25 [1] CRAN (R 4.0.2)
fastmap
forcats
                * 0.5.1
                           2021-01-27 [1] CRAN (R 4.0.2)
                  0.8-80
                           2020-05-24 [2] CRAN (R 4.0.3)
foreign
                  1.5.0
                           2020-07-31 [1] CRAN (R 4.0.2)
fs
                  0.1.0
                           2020-10-31 [1] CRAN (R 4.0.2)
generics
                           2020-12-15 [1] CRAN (R 4.0.2)
GetoptLong
                  1.0.5
                           2021-06-25 [1] CRAN (R 4.0.3)
ggplot2
                * 3.3.5
ggpubr
                * 0.4.0
                           2020-06-27 [1] CRAN (R 4.0.2)
                  0.6.3
                           2021-09-09 [1] CRAN (R 4.0.3)
ggsignif
GlobalOptions
                  0.1.2
                           2020-06-10 [1] CRAN (R 4.0.2)
                           2020-08-27 [1] CRAN (R 4.0.2)
glue
                  1.4.2
                  0.3.0
                           2019-03-25 [1] CRAN (R 4.0.2)
gtable
haven
                  2.4.3
                           2021-08-04 [1] CRAN (R 4.0.3)
                           2021-09-26 [1] CRAN (R 4.0.3)
hms
                  1.1.1
htmltools
                  0.5.2
                           2021-08-25 [1] CRAN (R 4.0.3)
                  1.4.2
                           2020-07-20 [1] CRAN (R 4.0.2)
httr
IRanges
                  2.24.1
                           2020-12-12 [1] Bioconductor
IRdisplay
                  1.0
                           2021-01-20 [1] CRAN (R 4.0.2)
IRkernel
                  1.2
                           2021-05-11 [1] CRAN (R 4.0.3)
                           2020-12-09 [1] CRAN (R 4.0.2)
jsonlite
                  1.7.2
lifecycle
                  1.0.1
                           2021-09-24 [1] CRAN (R 4.0.3)
                           2021-02-26 [1] CRAN (R 4.0.3)
lubridate
                  1.7.10
                  2.0.1
                           2020-11-17 [1] CRAN (R 4.0.2)
magrittr
                  0.61.0
                           2021-09-17 [1] CRAN (R 4.0.3)
matrixStats
                           2020-05-19 [1] CRAN (R 4.0.2)
modelr
                  0.1.8
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
                           2021-09-26 [1] CRAN (R 4.0.3)
pillar
                  1.6.3
                  2.0.3
                           2019-09-22 [1] CRAN (R 4.0.2)
pkgconfig
                  0.1 - 7
                           2013-12-03 [1] CRAN (R 4.0.2)
png
               * 0.3.4
                           2020-04-17 [1] CRAN (R 4.0.2)
purrr
R6
                  2.5.1
                           2021-08-19 [1] CRAN (R 4.0.3)
```

RColorBrewer		1.1-2	2014-12-07	[1]	CRAN	(R 4.0.2)
Rcpp		1.0.7	2021-07-07	[1]	CRAN	(R 4.0.3)
readr	*	2.0.2	2021-09-27	[1]	CRAN	(R 4.0.3)
readxl		1.3.1	2019-03-13	[1]	CRAN	(R 4.0.2)
repr		1.1.3	2021-01-21	[1]	CRAN	(R 4.0.2)
reprex		2.0.1	2021-08-05	[1]	CRAN	(R 4.0.3)
rio		0.5.27	2021-06-21	[1]	CRAN	(R 4.0.3)
rjson		0.2.20	2018-06-08	[1]	CRAN	(R 4.0.2)
rlang		0.4.11	2021-04-30	[1]	CRAN	(R 4.0.3)
rstatix		0.7.0	2021-02-13	[1]	CRAN	(R 4.0.3)
rstudioapi		0.13	2020-11-12	[1]	CRAN	(R 4.0.2)
rvest		1.0.1	2021-07-26	[1]	CRAN	(R 4.0.3)
S4Vectors		0.28.1	2020-12-09	[1]	Bioco	nductor
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)
shape		1.4.6	2021-05-19	[1]	CRAN	(R 4.0.3)
stringi		1.7.4	2021-08-25	[1]	CRAN	(R 4.0.3)
stringr	*	1.4.0	2019-02-10	[1]	CRAN	(R 4.0.2)
tibble	*	3.1.4	2021-08-25	[1]	CRAN	(R 4.0.3)
tidyr	*	1.1.4	2021-09-27	[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)

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