main

December 10, 2020

1 Boxplot of top five up- and downregulated genes

```
[1]: suppressMessages({library(ggpubr)
                       library(tidyverse)
                       library(data.table)})
[2]: save_ggplots <- function(p, fn, w=7, h=7){
         for(ext in c('.svg', '.png', '.pdf')){
             ggsave(p, filename=paste0(fn, ext), width=w, height=h)
         }
     }
     add_symnum <- function(res){</pre>
         symnum.args < -list(cutpoints = c(0, 0.0001, 0.001, 0.01, 0.05, 1),
                              symbols = c("****", "***", "**", "ns"))
         symnum.args$x <- res$adj.P.Val</pre>
         pvalue.signif <- do.call(stats::symnum, symnum.args) %>%
             as.character()
         pvalue.format <- format.pval(res$adj.P.Val, digits = 2)</pre>
         res <- res %>%
             dplyr::ungroup() %>%
             mutate(FDR = pvalue.format, p.signif = pvalue.signif)
         return(res)
     }
```

1.1 Select most significant DEGs, up- and downregulated

```
[3]: genes = fread('../../metrics_summary/_m/female_specific_DE_genes.txt') %>%
    filter(Type == 'gene') %>% add_symnum()

up_genes = genes %>% filter(t > 0) %>% mutate('Direction'='Upregulated') %>%
    →head(5)

down_genes = genes %>% filter(t < 0) %>% mutate('Direction'='Downregulated')

→%>% head(5)

sig_genes = bind_rows(up_genes, down_genes) %>% mutate_at(vars(Direction), as.

→factor)
```

```
Feature
                                        gencodeID
                                                             Symbol
                                                                        ensemblID
                                                                                          Chron
                   <chr>
                                        <chr>
                                                             <chr>
                                                                        <chr>
                                                                                           <chr>
                   ENSG00000153132.12
                                        ENSG00000153132.12
                                                            CLGN
                                                                        ENSG00000153132
                                                                                          chr4
                   ENSG00000179083.6
                                        ENSG00000179083.6
                                                            FAM133A
                                                                        ENSG00000179083
                                                                                          chrX
                   ENSG00000165733.7
                                        ENSG00000165733.7
                                                            BMS1
                                                                        ENSG00000165733
                                                                                          chr10
                   ENSG00000183023.18
                                        ENSG00000183023.18
                                                            SLC8A1
                                                                        ENSG00000183023
                                                                                          chr2
A data.table: 10 \times 15
                   ENSG00000236268.5
                                        ENSG00000236268.5
                                                            LINC01361
                                                                        ENSG00000236268
                                                                                          chr1
                   ENSG00000268049.1
                                        ENSG00000268049.1
                                                                        ENSG00000268049
                                                                                          chr19
                   ENSG00000083812.11
                                        ENSG00000083812.11
                                                            ZNF324
                                                                        ENSG00000083812
                                                                                          chr19
                   ENSG00000118960.12
                                        ENSG00000118960.12
                                                            HS1BP3
                                                                        ENSG00000118960
                                                                                          chr2
                   ENSG00000114554.11
                                        ENSG00000114554.11
                                                            PLXNA1
                                                                        ENSG00000114554
                                                                                          chr3
                   ENSG00000173714.7
                                        ENSG00000173714.7
                                                             WFIKKN2
                                                                        ENSG00000173714
                                                                                          chr17
```

1.2 Load phenotypes

```
Individual ID
                                  SampleID
                                                               Dx
                                                                      Sex
                 <fct>
                                  <fct>
                                                               <fct>
                                                                      <fct>
A data.table: 2 \times 4
                 CMC HBCC 003
                                  CMC HBCC RNA PFC 3158
                                                              CTL
                                                                      Male
                 CMC HBCC 005
                                  CMC HBCC RNA PFC 3152
                                                              CTL
                                                                     Male
```

1.3 Load residualized expression

```
[5]: res_file = '../../../../interaction_sex_sz/cmc_dlpfc/_m/genes/

→residualized_expression.tsv'

resdf0 = fread(res_file) %>%

filter(V1 %in% sig_genes$gencodeID) %>%

column_to_rownames(var="V1") %>% t %>%

as.data.frame %>% rownames_to_column() %>%
```

```
rename(SampleID=rowname) %>%
  mutate_at("SampleID", as.factor)
resdf0[1:2, 1:5]
```

Warning message in fread(res_file):

"Detected 858 column names but the data has 859 columns (i.e. invalid file). Added 1 extra default column name for the first column which is guessed to be row names or an index. Use setnames() afterwards if this guess is not correct, or fix the file write command that created the file to create a valid file."

1.3.1 Merge and melt dataframe

1.8580 2.19

```
SampleID
                                            Sex
                                                   Dx
                                                           gencodeID
                                                                               Res
                                                                                          Featu
              <fct>
                                            <fct>
                                                   <fct>
                                                           <fct>
                                                                               <dbl>
                                                                                          <chr
A tibble: 2 \times 10
              CMC HBCC RNA PFC 3158
                                                   CTL
                                                                                          ENSO
                                            Male
                                                           ENSG00000083812.11
                                                                               -0.4536485
              CMC_HBCC_RNA_PFC_3158
                                                   CTL
                                                                                          ENSO
                                            Male
                                                           ENSG00000114554.11
                                                                               0.2280048
```

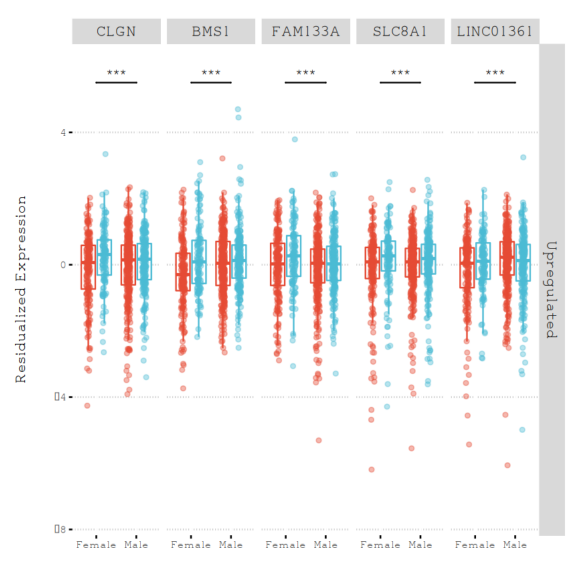
1.3.2 Initial ggplot with ggpubr

```
tmp = sig_genes %>%
    mutate(group1='Female', group2='Male', y_pos=5.5) %>%
    filter(Direction == 'Upregulated') %>%
    mutate_if(is.character, as.factor)

bxp_up <- df %>% filter(Direction=="Upregulated") %>%
        ggboxplot(x="Sex", y="Res", facet.by=c("Direction", "New_ID"),
        add='jitter', color="Dx", xlab='', palette="npg",
        panel.labs.font=list(face='bold', size = 14),
        ylab='Residualized Expression', ylim=c(-7.5, 6),
        add.params=list(alpha=0.4), legend="",
        ggtheme=theme_pubclean(), outlier.shape=NA) +
```

Warning message:

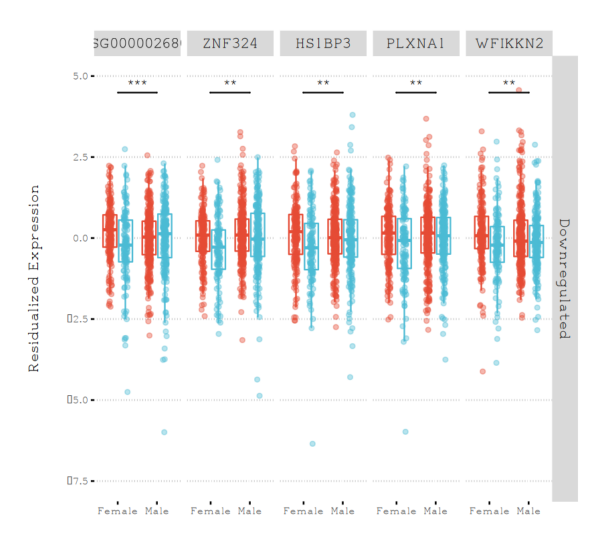
"Ignoring unknown aesthetics: xmin, xmax, annotations, y_position"



```
[8]: tmp = sig_genes %>%
         mutate(group1='Female', group2='Male', y_pos=4.5) %>%
         filter(Direction == 'Downregulated') %>%
         mutate_if(is.character, as.factor)
     bxp_down <- df %>% filter(Direction=="Downregulated") %>%
             ggboxplot(x="Sex", y="Res", facet.by=c("Direction", "New_ID"),
                       add='jitter', color="Dx", xlab='', palette="npg",
                       panel.labs.font=list(face='bold', size = 14),
                       ylab='Residualized Expression', ylim=c(-7.5, 5),
                       add.params=list(alpha=0.4), legend="bottom",
                       ggtheme=theme_pubclean(), outlier.shape=NA) +
              geom_signif(data=tmp, tip_length = 0,
                          aes(xmin=group1, xmax=group2, annotations=p.signif,__
      \rightarrowy_position = y_pos),
                          manual=TRUE) +
              font("xy.title", size=14, face="bold") + font("legend.title", size=16, __
      →face="bold") +
              font("legend.text", size=16)
     bxp_down
```

Warning message:

[&]quot;Ignoring unknown aesthetics: xmin, xmax, annotations, y_position"



Dx 🖨 CTL 🖨 SZ

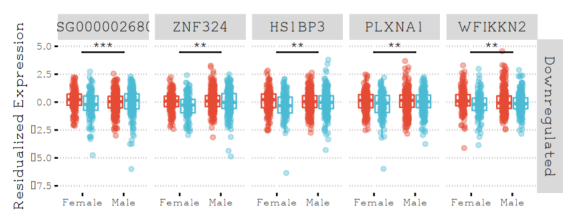
1.3.3 Annotate figure

```
[9]: figure <- ggarrange(bxp_up, bxp_down, ncol = 1, nrow = 2)
figure <- annotate_figure(figure, top=text_grob("CMC DLPFC", size=20,_u

face='bold'))
figure
```

CMC DLPFC





Dx 🖨 CTL 🖨 SZ

```
[10]: save_ggplots(figure, 'femaleSpecific_SZ', 12, 10)
```

1.4 Session Info

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

[1] "2020-12-10 15:07:19 EST"

user system elapsed 11.040 0.451 13.723

Session info setting value

version R version 4.0.2 (2020-06-22)

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 2020-12-10

Packages

1 acrages							
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		1.2.0	2020-11-02	[1]	CRAN	(R	4.0.2)
base64enc		0.1-3	2015-07-28	[1]	CRAN	(R	4.0.2)
broom		0.7.2	2020-10-20	[1]	CRAN	(R	4.0.2)
Cairo		1.5-12.2	2020-07-07	[1]	CRAN	(R	4.0.2)
car		3.0-10	2020-09-29	[1]	CRAN	(R	4.0.2)
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		2.2.0	2020-11-20	[1]	CRAN	(R	4.0.2)
colorspace		2.0-0	2020-11-11	[1]	CRAN	(R	4.0.2)
cowplot		1.1.0	2020-09-08	[1]	CRAN	(R	4.0.2)
crayon		1.3.4	2017-09-16	[1]	CRAN	(R	4.0.2)
curl		4.3	2019-12-02	[1]	CRAN	(R	4.0.2)
data.table	*	1.13.4	2020-12-08	[1]	CRAN	(R	4.0.2)
DBI		1.1.0	2019-12-15	[1]	CRAN	(R	4.0.2)
dbplyr		2.0.0	2020-11-03	[1]	CRAN	(R	4.0.2)
digest		0.6.27	2020-10-24	[1]	CRAN	(R	4.0.2)
dplyr	*	1.0.2	2020-08-18	[1]	CRAN	(R	4.0.2)
ellipsis		0.3.1	2020-05-15	[1]	CRAN	(R	4.0.2)
evaluate		0.14	2019-05-28	[1]	CRAN	(R	4.0.2)
fansi		0.4.1	2020-01-08	[1]	CRAN	(R	4.0.2)
farver		2.0.3	2020-01-16	[1]	CRAN	(R	4.0.2)
forcats	*	0.5.0	2020-03-01	[1]	CRAN	(R	4.0.2)
foreign		0.8-80	2020-05-24	[2]	CRAN	(R	4.0.2)
fs		1.5.0	2020-07-31	[1]	CRAN	(R	4.0.2)
gdtools	*	0.2.2	2020-04-03	[1]	CRAN	(R	4.0.2)
generics		0.1.0	2020-10-31	[1]	CRAN	(R	4.0.2)
ggplot2	*	3.3.2	2020-06-19	[1]	CRAN		4.0.2)
ggpubr	*	0.4.0	2020-06-27	[1]	CRAN		4.0.2)
ggsci		2.9	2018-05-14	[1]	CRAN		4.0.2)
ggsignif		0.6.0	2019-08-08	[1]	CRAN		4.0.2)
55 5				- -			,

```
1.4.2
                        2020-08-27 [1] CRAN (R 4.0.2)
glue
gridExtra
              2.3
                        2017-09-09 [1] CRAN (R 4.0.2)
gtable
              0.3.0
                        2019-03-25 [1] CRAN (R 4.0.2)
haven
              2.3.1
                        2020-06-01 [1] CRAN (R 4.0.2)
hms
              0.5.3
                        2020-01-08 [1] CRAN (R 4.0.2)
                        2020-06-16 [1] CRAN (R 4.0.2)
htmltools
              0.5.0
httr
              1.4.2
                        2020-07-20 [1] CRAN (R 4.0.2)
IRdisplay
              0.7.0
                        2018-11-29 [1] CRAN (R 4.0.2)
                        2020-07-20 [1] CRAN (R 4.0.2)
IRkernel
              1.1.1
jsonlite
              1.7.1
                        2020-09-07 [1] CRAN (R 4.0.2)
              0.4.2
                        2020-10-20 [1] CRAN (R 4.0.2)
labeling
              0.2.0
                        2020-03-06 [1] CRAN (R 4.0.2)
lifecycle
lubridate
              1.7.9.2
                        2020-11-13 [1] CRAN (R 4.0.2)
magrittr
              2.0.1
                        2020-11-17 [1] CRAN (R 4.0.2)
modelr
              0.1.8
                        2020-05-19 [1] CRAN (R 4.0.2)
munsell
              0.5.0
                        2018-06-12 [1] CRAN (R 4.0.2)
openxlsx
              4.2.3
                        2020-10-27 [1] CRAN (R 4.0.2)
              0.3 - 3.1
                        2020-10-14 [1] CRAN (R 4.0.2)
pbdZMQ
              1.4.7
                        2020-11-20 [1] CRAN (R 4.0.2)
pillar
              2.0.3
                        2019-09-22 [1] CRAN (R 4.0.2)
pkgconfig
                        2020-12-05 [1] CRAN (R 4.0.2)
              1.5.0
ps
                        2020-04-17 [1] CRAN (R 4.0.2)
purrr
            * 0.3.4
R6
              2.5.0
                        2020-10-28 [1] CRAN (R 4.0.2)
                        2020-07-06 [1] CRAN (R 4.0.2)
Rcpp
              1.0.5
readr
            * 1.4.0
                        2020-10-05 [1] CRAN (R 4.0.2)
              1.3.1
                        2019-03-13 [1] CRAN (R 4.0.2)
readxl
                        2020-01-28 [1] CRAN (R 4.0.2)
              1.1.0
repr
                        2019-05-16 [1] CRAN (R 4.0.2)
reprex
              0.3.0
                        2018-11-26 [1] CRAN (R 4.0.2)
rio
              0.5.16
              0.4.9
                        2020-11-26 [1] CRAN (R 4.0.2)
rlang
              0.6.0
                        2020-06-18 [1] CRAN (R 4.0.2)
rstatix
rstudioapi
              0.13
                        2020-11-12 [1] CRAN (R 4.0.2)
rvest
              0.3.6
                        2020-07-25 [1] CRAN (R 4.0.2)
              1.1.1
                        2020-05-11 [1] CRAN (R 4.0.2)
scales
sessioninfo
              1.1.1
                        2018-11-05 [1] CRAN (R 4.0.2)
stringi
              1.5.3
                        2020-09-09 [1] CRAN (R 4.0.2)
stringr
            * 1.4.0
                        2019-02-10 [1] CRAN (R 4.0.2)
svglite
              1.2.3.2
                        2020-07-07 [1] CRAN (R 4.0.2)
              0.3.2
                        2020-09-29 [1] CRAN (R 4.0.2)
systemfonts
tibble
            * 3.0.4
                        2020-10-12 [1] CRAN (R 4.0.2)
            * 1.1.2
                        2020-08-27 [1] CRAN (R 4.0.2)
tidyr
                        2020-05-11 [1] CRAN (R 4.0.2)
tidyselect
              1.1.0
tidyverse
            * 1.3.0
                        2019-11-21 [1] CRAN (R 4.0.2)
uuid
              0.1 - 4
                        2020-02-26 [1] CRAN (R 4.0.2)
vctrs
              0.3.5
                        2020-11-17 [1] CRAN (R 4.0.2)
withr
              2.3.0
                        2020-09-22 [1] CRAN (R 4.0.2)
xml2
              1.3.2
                        2020-04-23 [1] CRAN (R 4.0.2)
              2.1.1
                        2020-08-27 [1] CRAN (R 4.0.2)
zip
```

- [1] /home/jbenja13/R/x86_64-pc-linux-gnu-library/4.0
 [2] /usr/lib/R/library