main

August 4, 2021

1 Visualize results, local splicing

1.1 Load Libraries

```
[1]: library(tidyverse)
    library(ggplot2)
    library(DT)
    library(leafcutter)
    library(reshape2)
    library(gridExtra)
    library(intervals) # needed for pretty strand arrow placement
    library(foreach)
    library(grid)
    library(gtable)
    library(ggrepel)
```

tidyverse

```
1.3.1

ggplot2 3.3.5 purrr 0.3.4
tibble 3.1.2 dplyr 1.0.7
tidyr 1.1.3 stringr 1.4.0
readr 1.4.0 forcats 0.5.1
```

Conflicts tidyverse_conflicts()

Attaching packages

```
dplyr::filter() masks stats::filter()
dplyr::lag() masks stats::lag()
```

Loading required package: Rcpp

Attaching package: 'reshape2'

The following object is masked from 'package:tidyr':

smiths

```
The following object is masked from 'package:dplyr':
        combine
    Attaching package: 'intervals'
    The following object is masked from 'package:purrr':
        reduce
    The following object is masked from 'package:tidyr':
        expand
    Attaching package: 'foreach'
    The following objects are masked from 'package:purrr':
        accumulate, when
    1.2 Summary of results
[2]: lname = load('../../_m/leafviz.RData')
     lname
    1. 'introns' 2. 'clusters' 3. 'counts' 4. 'meta' 5. 'exons_table' 6. 'pca' 7. 'intron_summary' 8. 'clus-
    ter_summary' 9. 'introns_to_plot' 10. 'cluster_ids' 11. 'sample_table' 12. 'annotation_code'
    13. 'code'
[3]: levels(meta$group) <- c("Female", "Male")
     sample_table
```

Attaching package: 'gridExtra'

```
A data frame: 2 \times 2 \frac{\text{group count}}{<\text{chr}>} \frac{<\text{chr}>}{F} \frac{114}{M} \frac{245}{M}
```

[4]: cluster_summary

A data.frame: 3×2	Results	n
	<chr></chr>	<int $>$
	Number of differentially spliced clusters at $FDR = 0.05$	352
	Fully annotated	160
	Contain unannotated junctions	192

[5]: intron_summary

	Results	n
A data.frame: 5×2	<chr></chr>	<int $>$
	Number of fully annotated junctions	1249
	Number of junctions with cryptic 5' splice site	119
	Number of junctions with cryptic 3' splice site	145
	Number of junctions with two cryptic splice sites	38
	Number of novel junctions that connect two annotated splice sites	118

[6]: clusters['gene'] <- gsub("</i>", "", gsub("<i>", "", clusters\$gene))
head(clusters)

```
clusterID
                                                  coord
                                                                                                     FDR
                                                                             gene
                                                                                         annotation
                                          <dbl>
                         <chr>
                                                  <chr>
                                                                             <chr>
                                                                                         <chr>
                                                                                                     <dbl
                    352
                         clu_755_-
                                          3
                                                  chrX:53217966-53220839
                                                                             KDM5C
                                                                                         annotated
                                                                                                     2.27e
                         clu 744 -
                                                  chrX:53176622-53193437
                    350
                                          10
                                                                             KDM5C
                                                                                         cryptic
                                                                                                     3.64e
A data.frame: 6 \times 6
                    282
                         clu_109096_+
                                         9
                                                  chr6:72292046-72390598
                                                                             RIMS1
                                                                                         cryptic
                                                                                                     1.89e
                    291
                         clu 143938 -
                                          22
                                                  chr6:130863718-130885096
                                                                             EPB41L2
                                                                                         cryptic
                                                                                                     1.89e
                    340
                         clu_61504_+
                                          3
                                                  chrX:47199106-47199480
                                                                             UBA1
                                                                                         cryptic
                                                                                                     2.45e
                    227
                         clu_21090_-
                                          11
                                                  chr3:197066754-197081051
                                                                             DLG1
                                                                                         cryptic
                                                                                                     2.32e
```

```
[7]: write.table(clusters, file="cluster_ds_results_annotated.txt", sep="\t", quote=FALSE, row.names=FALSE)
```

1.3 Generate plots

1.3.1 Define functions

```
[8]: filter_intron_table <- function(introns, clu){
    d < - introns %>% filter(clusterID == clu) %>%
        select(chr, start, end, verdict, deltapsi) %>%
        arrange(desc(abs(deltapsi))) %>%
        rename("APSI" = deltapsi)
    row.names(d) <- letters[1:nrow(d)] # letters is just a:z
    return(d)</pre>
```

```
}
getGeneLength <- function(gene_name, exons_table){</pre>
              <- exons_table[ exons_table$gene_name == gene_name, ]</pre>
    geneStart <- min(exons$start)</pre>
             <- max(exons$end)
    geneEnd
    geneLength <- geneEnd - geneStart</pre>
    if( geneLength >1e6){
        pixels <- 5000
    } else if ( geneLength > 5e5 & geneLength < 1e6){
        pixels <- 3000
    } else if ( geneLength > 1.5e5 & geneLength <= 5e5){</pre>
        pixels <- 2000
    } else {
        stopifnot(geneLength <= 1.5e5)</pre>
        pixels <- "auto"</pre>
    return(pixels)
}
select_data <- function(sel, clusters, exons_table){</pre>
    gene <- clusters[ sel, ]$gene</pre>
    width <- getGeneLength(gene, exons_table)</pre>
    clusterID <- clusters[ sel, ]$clusterID</pre>
    coord <- clusters[ sel, ]$coord</pre>
    return(list(gene = gene, width = width, cluster = clusterID, coord = coord))
}
```

1.3.2 Plot top 6 clusters

```
dev.off()
    if (is.numeric(mydata$width)) {
        new_width = mydata$width / 100
    } else {
        new_width = mydata$width
    }
    pdf(file=pasteO(dir, gene_name, '_allClusters_top_',num,'.pdf'),
        width=new_width, height=6)
    try(print(make_gene_plot(mydata$gene,
                         counts = counts,
                         introns = introns,
                         exons_table = exons_table,
                         cluster_list = clusters,
                         clusterID = mydata$cluster,
                         introns_to_plot = introns_to_plot, debug=F)))
    dev.off()
}
```

1.4 Plot splicing

```
[10]: dir.create("top10")
      for(num in 1:10){
          ii = plot_cluster(num, clusters, "top10/")
      }
     Warning message:
     "`guides(<scale> = FALSE)` is deprecated. Please use `guides(<scale> = "none")`
     instead."
     TableGrob (2 x 1) "arrange": 2 grobs
             cells
                      name
                                      grob
     1 1 (1-1,1-1) arrange gtable[layout]
     2 2 (2-2,1-1) arrange gtable[layout]
     Warning message:
     "'mode(width)' differs between new and previous
              ==> NOT changing 'width'"
     Warning message:
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     instead."
     TableGrob (2 x 1) "arrange": 2 grobs
       z
             cells
                      name
                                      grob
     1 1 (1-1,1-1) arrange gtable[layout]
     2 2 (2-2,1-1) arrange gtable[layout]
     Warning message:
     "'mode(width)' differs between new and previous
```

```
==> NOT changing 'width'"
Warning message:
"`guides(<scale> = FALSE)` is deprecated. Please use `guides(<scale> = "none")`
instead."
TableGrob (2 x 1) "arrange": 2 grobs
       cells
                 name
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       cells
                name
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```

```
TableGrob (2 x 1) "arrange": 2 grobs
             cells
       z
                      name
                                     grob
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     2 2 (2-2,1-1) arrange gtable[layout]
     Warning message:
     "'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> = "none")'
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     TableGrob (2 x 1) "arrange": 2 grobs
             cells
                      name
     1 1 (1-1,1-1) arrange gtable[layout]
     2 2 (2-2,1-1) arrange gtable[layout]
     Warning message:
     "'mode(width)' differs between new and previous
              ==> NOT changing 'width'"
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     instead."
     TableGrob (2 x 1) "arrange": 2 grobs
             cells
                      name
     1 1 (1-1,1-1) arrange gtable[layout]
     2 2 (2-2,1-1) arrange gtable[layout]
[11]: dir.create("x_chromosome")
      x_clu = clusters %>% filter(str_detect(coord, "chrX"), FDR < 0.05)</pre>
      x_clu
```

```
<dbl>
                         < chr >
                                                <chr>
                                                                          <chr>
                                                                                    <chr>
                         clu 755 -
                                               chrX:53217966-53220839
                                                                         KDM5C
                                                                                    annotated
                         clu 744 -
                                       10
                                                chrX:53176622-53193437
                                                                         KDM5C
                                                                                    cryptic
                         clu 61504 +
                                                                          UBA1
                                       3
                                                chrX:47199106-47199480
                                                                                    cryptic
                         clu 62902 +
                                       10
                                                chrX:153765580-153769162
                                                                         PLXNB3
                                                                                    cryptic
                         clu 1810 -
                                       3
                                                                         PNMA5
                                                                                    cryptic
                                                chrX:152989331-152991916
                         clu 752 -
                                                                         KDM5C
                                       3
                                                chrX:53210576-53211497
                                                                                    cryptic
                         clu 62904 +
                                       7
                                                                         PLXNB3
                                                chrX:153771081-153771864
                                                                                    cryptic
                         clu 62916 +
                                       5
                                                                         PLXNB3
                                                chrX:153777688-153778396
                                                                                    cryptic
                         clu 62431 +
                                       3
                                                chrX:115636978-115640408
                                                                         PLS3
                                                                                    annotated
                         clu_261_-
                                       11
                                                                         AP1S2
                                                                                    cryptic
                                               chrX:15825984-15845379
     A data.frame: 22 \times 6
                         clu_206_-
                                       5
                                                chrX:13785808-13938507
                                                                         GPM6B
                                                                                    annotated
                         clu 1841 -
                                       9
                                               chrX:154032557-154097604
                                                                         MECP2
                                                                                    cryptic
                         clu 1482 -
                                        3
                                               chrX:120428626-120441730
                                                                         LAMP2
                                                                                    annotated
                         clu 60959 +
                                       9
                                                chrX:2861722-2910874
                                                                         GYG2
                                                                                    cryptic
                         clu 62906 +
                                       5
                                                                         PLXNB3
                                                chrX:153772287-153773230
                                                                                    cryptic
                         clu 62231 +
                                       4
                                                chrX:103310981-103330984
                                                                         TCEAL7
                                                                                    cryptic
                         clu 480 -
                                       3
                                                chrX:41610025-41626604
                                                                         CASK
                                                                                    annotated
                         clu 12601 ?
                                                                         LAMP2
                                       4
                                                chrX:120428626-120441730
                                                                                    cryptic
                         clu 62948 +
                                       5
                                                chrX:154399941-154400702
                                                                         RPL10
                                                                                    cryptic
                         clu 61084 +
                                       6
                                                chrX:13735220-13736656
                                                                         OFD1
                                                                                    cryptic
                         clu_62377_+
                                       3
                                                chrX:108584525-108586615
                                                                         COL4A5
                                                                                    cryptic
                         clu 61903 +
                                                chr X:71167800-71171654
                                                                         NLGN3
                                                                                    cryptic
[12]: for(num in 1:dim(x_clu)[1]){
          plot_cluster(num, x_clu, "x_chromosome/")
      }
     Warning message:
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                                       grob
                       name
     1 1 (1-1,1-1) arrange gtable[layout]
     2 2 (2-2,1-1) arrange gtable[layout]
```

clusterID

Ν

coord

FDR

< dbl >

2.27e-43

3.64e-40

2.45e-16

 $1.06\mathrm{e}\text{-}09$

4.00e-09

8.61e-09

1.38e-07

1.14e-06

7.28e-06

1.37e-04

1.53e-04

9.14e-04

1.94e-03

8.68e-03

8.68e-03

1.14e-02

1.23e-02

1.70e-02

2.73e-02

3.09e-02

4.60e-02

4.93e-02

annotation

gene

```
Warning message:
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```

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     Warning message:
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     Warning message:
     "Removed 1 row(s) containing missing values (geom_path)."
     TableGrob (2 x 1) "arrange": 2 grobs
       7.
             cells
                      name
                                      grob
     1 1 (1-1,1-1) arrange gtable[layout]
     2 2 (2-2,1-1) arrange gtable[layout]
     Warning message:
     "'mode(width)' differs between new and previous
              ==> NOT changing 'width'"
     Error: Aesthetics must be either length 1 or the same as the data (1): x, xend,
     group and colour
[13]: dir.create("ank3")
      ank3 = clusters %>% filter(gene == 'ANK3')
      ank3
                                                                                       FDR.
                       clusterID
                                    Ν
                                            coord
                                                                    gene
                                                                            annotation
                        <chr>
                                     <dbl>
                                             <chr>
                                                                    <chr>
                                                                            <chr>
                                                                                       <dbl>
     A data.frame: 2 \times 6
                       clu 22299 -
                                    3
                                            chr10:60059430-60063111
                                                                    ANK3
                                                                            annotated
                                                                                       0.00159
                       clu 22304 -
                                            chr10:60139087-60166824
                                                                    ANK3
                                                                            annotated
                                                                                       0.00895
                                    3
[14]: for(num in 1:dim(ank3)[1]){
          plot_cluster(num, ank3, "ank3/")
      }
     Warning message:
     "`guides(<scale> = FALSE)` is deprecated. Please use `guides(<scale> = "none")`
```