Creating Dotplot

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Loading package	

```
suppressMessages(library(ggplot2))
suppressMessages(library("forcats"))
suppressMessages(library(formatR))
```

Function adapting the format of tsv file

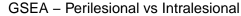
```
modify_GSEA_data <- function(GSEA_results){</pre>
  GSEA_results<- GSEA_results[order(GSEA_results$NES, decreasing = T),]
  GSEA_results$NAME <- gsub("_", " ", GSEA_results$NAME)</pre>
  GSEA_results$NAME <- substr(GSEA_results$NAME, start = 6, stop =
nchar(GSEA_results$NAME))
  GSEA_results
}
```

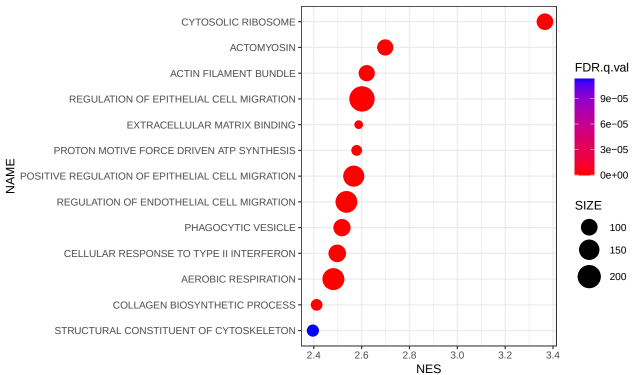
The most relevant ontologies were selected and written in the file "Plot-gsea-Peri vs Intra.docx". The complete list of ontology are described in tsv files.

Dotplot perilesional vs intralesional

```
# loading table
dir <- "../output/perivvsintra/"</pre>
GSEA_BP_res <- read.table(paste0(dir,</pre>
"my_analysis.Gsea.1684311691666_BP2/gsea_report_for_perilesional_1684311691666.tsv"),
header = T, sep = "\t")
GSEA_BP_res <- modify_GSEA_data(GSEA_BP_res)</pre>
```

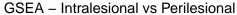
```
GSEA_BP_res \leftarrow GSEA_BP_res[c(6,9,12,16,20,28,53),]
# MF
GSEA_MF_res <- read.table(paste0(dir,</pre>
"my_analysis.Gsea.1684310907505_MF/gsea_report_for_perilesional_1684310907505.tsv"),
header = T, sep = "\t")
GSEA MF res <- modify GSEA data(GSEA MF res)
GSEA_MF_res <- GSEA_MF_res[c(3,11),]</pre>
# CC
GSEA_CC_res <- read.table(paste0(dir,</pre>
"my analysis. Gsea. 1684311503281 CC/gsea report for perilesional 1684311503281.tsv"),
header = T, sep = "\t")
GSEA_CC_res <- modify_GSEA_data(GSEA_CC_res)</pre>
GSEA_CC_res <- GSEA_CC_res[c(1,10,14,18),]</pre>
GSEA_Data <- rbind(GSEA_MF_res, GSEA_CC_res, GSEA_BP_res)</pre>
GSEA Data <- GSEA_Data[order(GSEA_Data$NES, decreasing = F),]</pre>
row.names(GSEA_Data) <- NULL</pre>
# define the column has a factor so that the order of Ontology is maintain in the plot.
GSEA_Data$NAME <- factor(GSEA_Data$NAME, levels = GSEA_Data$NAME)</pre>
ggplot(GSEA_Data, aes(x = NES,y =NAME, size = SIZE, color = FDR.q.val)) + geom_point() +
ggtitle("GSEA - Perilesional vs Intralesional") + scale color gradient(low = "red", high
= "blue") + theme bw()+ scale size(range = c(3,10))
```

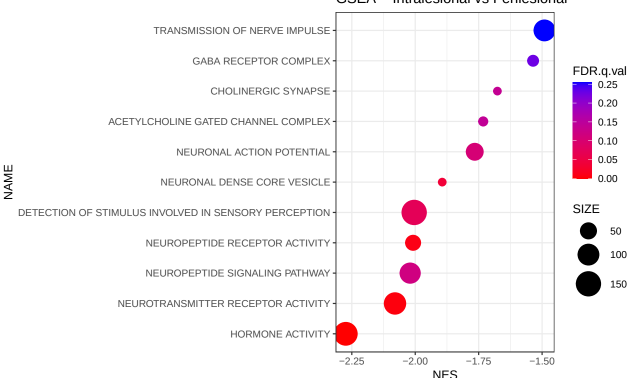




Dotplot intralesional vs perilesional

```
# BP
GSEA_BP_res <- read.table(paste0(dir,
"my_analysis.Gsea.1684311691666_BP2/gsea_report_for_intralesional_1684311691666.tsv"),
header = T, sep = "\t")
GSEA_BP_res <- modify_GSEA_data(GSEA_BP_res)</pre>
GSEA BP res <- GSEA BP res[rev(rownames(GSEA BP res)),]
GSEA_BP_res \leftarrow GSEA_BP_res[c(1,2,21,78),]
# MF
GSEA_MF_res <- read.table(paste0(dir,</pre>
"my_analysis.Gsea.1684310907505_MF/gsea_report_for_intralesional_1684310907505.tsv"),
header = T, sep = "\t")
GSEA_MF_res <- modify_GSEA_data(GSEA_MF_res)</pre>
GSEA_MF_res <- GSEA_MF_res[rev(rownames(GSEA_MF_res)),]</pre>
GSEA_MF_res <- GSEA_MF_res[c(2,4,6),]</pre>
# CC
GSEA_CC_res <- read.table(paste0(dir,</pre>
"my_analysis.Gsea.1684311503281_CC/gsea_report_for_intralesional_1684311503281.tsv"),
header = T, sep = "\t")
GSEA_CC_res <- modify_GSEA_data(GSEA_CC_res)</pre>
GSEA_CC_res <- GSEA_CC_res[rev(rownames(GSEA_CC_res)),]</pre>
GSEA CC res \leftarrow GSEA CC res[c(2,3,5,9),]
GSEA Data <- rbind(GSEA MF res, GSEA CC res, GSEA BP res)
GSEA_Data <- GSEA_Data[order(GSEA_Data$NES, decreasing = F),]</pre>
row.names(GSEA_Data) <- NULL</pre>
# define the column has a factor so that the order of Ontology is maintain in the plot.
GSEA_Data$NAME <- factor(GSEA_Data$NAME, levels = GSEA_Data$NAME)
ggplot(GSEA_Data, aes(x = NES,y =NAME, size = SIZE, color = FDR.q.val)) + geom_point() +
ggtitle("GSEA - Intralesional vs Perilesional") + scale_color_gradient(low = "red", high
= "blue") + theme_bw()+ scale_size(range = c(3,10))
```





#qqsave("GSEA IntravsPeri.pdf", width = 12, height = 8)

sessionInfo()

```
## R version 4.3.1 (2023-06-16)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 22.04.3 LTS
##
## Matrix products: default
          /usr/lib/x86 64-linux-gnu/openblas-pthread/libblas.so.3
## LAPACK: /usr/lib/x86_64-linux-gnu/openblas-pthread/libopenblasp-r0.3.20.so; LAPACK version 3.10.0
##
## locale:
## [1] LC_CTYPE=en_US.UTF-8
                                  LC NUMERIC=C
## [3] LC_TIME=fr_BE.UTF-8
                                  LC_COLLATE=en_US.UTF-8
## [5] LC MONETARY=fr BE.UTF-8
                                  LC MESSAGES=en US.UTF-8
## [7] LC_PAPER=fr_BE.UTF-8
                                  LC_NAME=C
## [9] LC_ADDRESS=C
                                  LC_TELEPHONE=C
## [11] LC_MEASUREMENT=fr_BE.UTF-8 LC_IDENTIFICATION=C
## time zone: Europe/Brussels
## tzcode source: system (glibc)
##
## attached base packages:
## [1] stats
                graphics grDevices utils
                                              datasets methods
                                                                   base
## other attached packages:
## [1] formatR_1.14 forcats_1.0.0 ggplot2_3.4.2
##
```

```
## loaded via a namespace (and not attached):
## [1] vctrs_0.6.3
                        cli_3.6.1
                                         knitr_1.43
                                                          rlang_1.1.1
## [5] xfun_0.39
                        highr_0.10
                                                          labeling_0.4.2
                                         generics_0.1.3
## [9] glue_1.6.2
                        colorspace_2.1-0 htmltools_0.5.5 scales_1.2.1
                                         grid_4.3.1
                                                          evaluate_0.21
## [13] fansi_1.0.4
                        rmarkdown_2.23
## [17] munsell_0.5.0
                        tibble_3.2.1
                                         fastmap_1.1.1
                                                          yaml_2.3.7
## [21] lifecycle_1.0.3 compiler_4.3.1
                                         dplyr_1.1.2
                                                          pkgconfig_2.0.3
## [25] rstudioapi_0.14 farver_2.1.1
                                         digest_0.6.33
                                                          R6_2.5.1
## [29] tidyselect_1.2.0 utf8_1.2.3
                                         pillar_1.9.0
                                                          magrittr_2.0.3
## [33] withr_2.5.0
                                         gtable_0.3.3
                        tools_4.3.1
```