# Tabula Sapiens - MafB

## Domien

### $2025\text{-}09\text{-}19\ 11\text{:}21\text{:}41\ +0200$

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### Introduction

To investigate the conservedness of MafB to imprint Mac identity across species we quantified the correlation between Mafb expression and the expression of macrophage sigature genes and MafB target genes in single cell atlases of mouse (Tabula Muris Senis), human (Tabula Sapiens), lemur (Tabula Microcebus), pig (Pig Cell Atlas), African clawed frog (Xenopus Cell Landscape) and zebrafish (Zebrafish Cell Landscape, ZCL).

### Load packages

```
suppressMessages({
    library(SeuratObject)
    library(Seurat)
    library(reticulate)
    library(sceasy)
    library(ggplot2)
    library(ggrastr)
    library(readxl)
})
```

### Download Tabula Sapiens scanpy object from Figshare server

```
https://figshare.com/articles/dataset/Tabula_Sapiens_release_1_0/14267219?file=40067134
```

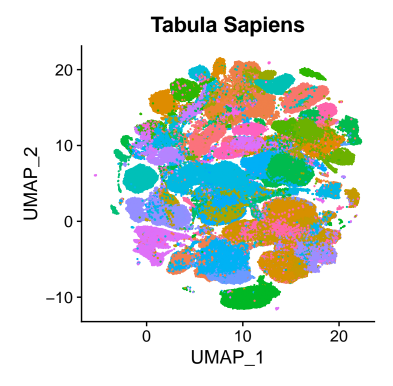
### Convert Scanpy object to Seurat object

```
sceasy::convertFormat("TabulaSapiens.h5ad", from = "anndata",
to = "seurat", outFile = "TBS.rds")
```

### Load Tabula Sapiens Seurat object

```
TBS <- readRDS("/media/uliege/T7/SingleCellAtlas/TBS.rds")
```

### Visualize clusters



# Calculate correlation between MAFB expression and Mac signature score $\,$

```
FeaturePlot(TBS, features = "MAFB", raster = T)
```

<sup>##</sup> Rasterizing points since number of points exceeds 100,000.

<sup>##</sup> To disable this behavior set 'raster=FALSE'

# MAFB 20 20 10 0 10 0 SCVlumap\_1

```
Mac_sign <- read_excel("Mac_sign_human.xlsx")

Mac_sign <- Mac_sign$ortholog_name

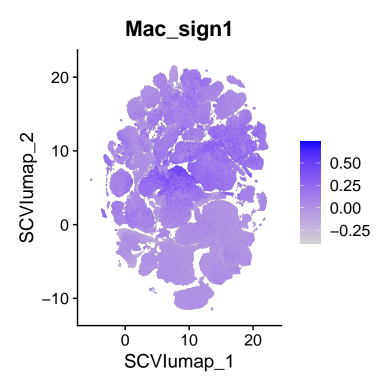
Mac_sign <- intersect(Mac_sign, rownames(TBS))

TBS <- AddModuleScore(TBS, features = list(c(Mac_sign)), name = "Mac_sign")

FeaturePlot(TBS, features = "Mac_sign1", raster = T)</pre>
```

 $\mbox{\tt \#\#}$  Rasterizing points since number of points exceeds 100,000.

## To disable this behavior set 'raster=FALSE'



```
FeatureScatter(TBS, feature1 = "MAFB", feature2 = "Mac_sign1") +
   geom_smooth(method = "lm", colour = "black") + theme(legend.position = "none")
```

```
## Rasterizing points since number of points exceeds 100,000.
## To disable this behavior set 'raster=FALSE'
```

<sup>## &#</sup>x27;geom\_smooth()' using formula = 'y ~ x'

```
0.75

0.50

0.00

0.00

0.00

0.25

0.25

0.25

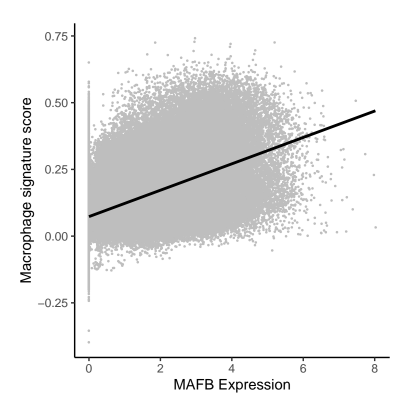
0.25

0.45

0.8
```

## 'geom\_smooth()' using formula = 'y ~ x'

```
corr <- FetchData(TBS, vars = "MAFB")</pre>
corr <- cbind(corr, FetchData(TBS, vars = "Mac_sign1"))</pre>
cor.test(corr$MAFB, corr$Mac_sign1, method = "pearson")
##
   Pearson's product-moment correlation
##
##
## data: corr$MAFB and corr$Mac_sign1
## t = 350.48, df = 483150, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.4479783 0.4524746
## sample estimates:
##
         cor
## 0.4502293
p2 <- ggplot(corr, aes(x = MAFB, y = Mac_sign1)) + geom_point(size = 0.1,
    colour = "grey") + geom_smooth(method = "lm", colour = "black") +
    xlab("MAFB Expression") + ylab("Macrophage signature score") +
    theme_classic()
rasterize(p2, layers = "Point", dpi = 600)
```



# Calculate correlation between MAFB expression and MafB target gene signature score

```
Conserved_MafB_target_genes <- read.table("Conserved_MafB_genes.txt")

Conserved_MafB_target_genes <- intersect(Conserved_MafB_target_genes$V1,
    rownames(TBS))

TBS <- AddModuleScore(TBS, features = list(c(Conserved_MafB_target_genes)),
    name = "MafB_target_sign")</pre>
```

```
FeaturePlot(TBS, features = "MafB_target_sign1", raster = T)
```

<sup>##</sup> Rasterizing points since number of points exceeds 100,000.

<sup>##</sup> To disable this behavior set 'raster=FALSE'

# MafB\_target\_sign1 20 20 0.6 0.4 0.2 0.0 -0.2 SCVlumap\_1

```
FeatureScatter(TBS, feature1 = "MAFB", feature2 = "MafB_target_sign1") +
    geom_smooth(method = "lm", colour = "black") + theme(legend.position = "none")
```

```
## Rasterizing points since number of points exceeds 100,000.
## To disable this behavior set 'raster=FALSE'
```

<sup>## &#</sup>x27;geom\_smooth()' using formula = 'y ~ x'

```
0.49

0.6

0.49

0.4

0.0

0.4

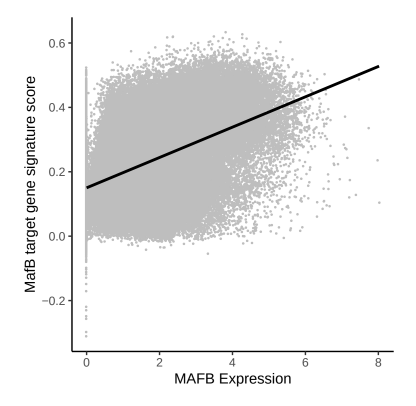
0.2

0 2 4 6 8

MAFB
```

## 'geom\_smooth()' using formula = 'y ~ x'

```
corr <- cbind(corr, FetchData(TBS, vars = "MafB_target_sign1"))</pre>
cor.test(corr$MAFB, corr$MafB_target_sign1, method = "pearson")
##
##
   Pearson's product-moment correlation
##
## data: corr$MAFB and corr$MafB_target_sign1
## t = 392.62, df = 483150, p-value < 2.2e-16
\#\# alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
   0.4896696 0.4939450
## sample estimates:
##
         cor
## 0.4918103
p3 <- ggplot(corr, aes(x = MAFB, y = MafB_target_sign1)) + geom_point(size = 0.1,
    colour = "grey") + geom_smooth(method = "lm", colour = "black") +
    xlab("MAFB Expression") + ylab("MafB target gene signature score") +
    theme_classic()
rasterize(p3, layers = "Point", dpi = 600)
```



### sessionInfo()

```
## R version 4.4.1 (2024-06-14)
## Platform: x86_64-pc-linux-gnu
## Running under: Ubuntu 20.04.6 LTS
##
## Matrix products: default
           /usr/lib/x86_64-linux-gnu/blas/libblas.so.3.9.0
## LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.9.0
##
## 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] readxl_1.4.3
                          ggrastr_1.0.2
                                             ggplot2_3.4.4
                                                                sceasy_0.0.7
## [5] reticulate_1.39.0 Seurat_4.3.0
                                             SeuratObject_4.1.3 sp_2.1-4
## loaded via a namespace (and not attached):
```

```
##
     [1] deldir 2.0-4
                                 pbapply_1.4-3
                                                         gridExtra 2.3
##
     [4] formatR 1.14
                                                         magrittr_2.0.3
                                 rlang_1.1.4
     [7] RcppAnnoy 0.0.18
##
                                 spatstat.geom_3.3-3
                                                         matrixStats 1.4.1
                                 compiler_4.4.1
##
    [10] ggridges_0.5.3
                                                         mgcv_1.9-1
##
    [13] png_0.1-8
                                 vctrs_0.6.5
                                                         reshape2_1.4.4
##
    [16] stringr 1.5.1
                                 pkgconfig 2.0.3
                                                         fastmap 1.2.0
    [19] labeling 0.4.3
                                 utf8 1.2.4
                                                         promises 1.2.0.1
                                 ggbeeswarm_0.7.2
    [22] rmarkdown 2.28
                                                         purrr_1.0.2
##
##
    [25] xfun 0.47
                                 jsonlite 1.8.9
                                                         goftest_1.2-2
    [28] highr_0.11
                                 later_1.2.0
##
                                                         spatstat.utils_3.1-0
   [31] irlba_2.3.5.1
                                                        cluster_2.1.6
                                 parallel_4.4.1
                                                         stringi_1.6.2
##
    [34] R6_2.5.1
                                 ica_1.0-2
##
    [37] RColorBrewer_1.1-3
                                 spatstat.data_3.1-2
                                                         parallelly_1.26.0
   [40] spatstat.univar_3.0-1
                                 cellranger_1.1.0
                                                         lmtest_0.9-38
##
##
   [43] scattermore_0.7
                                 Rcpp_1.0.13
                                                         knitr_1.48
##
    [46] tensor_1.5
                                 future.apply_1.7.0
                                                         zoo_1.8-9
##
    [49] sctransform_0.4.1
                                 httpuv_1.6.1
                                                         Matrix_1.6-1.1
    [52] splines 4.4.1
                                 igraph 1.2.6
                                                         tidyselect_1.2.1
   [55] abind_1.4-5
                                 rstudioapi_0.16.0
                                                         yaml_2.2.1
##
##
    [58] spatstat.random 3.3-2
                                 codetools 0.2-19
                                                         miniUI 0.1.1.1
##
    [61] spatstat.explore_3.3-2 listenv_0.8.0
                                                         lattice_0.22-5
    [64] tibble 3.2.1
                                 plyr_1.8.6
                                                         withr 3.0.1
   [67] shiny_1.9.1
##
                                 ROCR_1.0-11
                                                         evaluate_1.0.0
    [70] Rtsne 0.15
                                 future 1.21.0
                                                         survival 3.7-0
##
                                 fitdistrplus_1.1-5
  [73] polyclip_1.10-0
                                                         pillar 1.9.0
##
   [76] KernSmooth 2.23-24
                                 plotly_4.10.4
                                                         generics 0.1.0
##
   [79] munsell_0.5.1
                                 scales_1.3.0
                                                         globals_0.14.0
                                                         lazyeval_0.2.2
##
    [82] xtable_1.8-4
                                 glue_1.7.0
##
   [85] tools_4.4.1
                                 data.table_1.14.0
                                                         RANN_2.6.1
##
   [88] leiden_0.3.8
                                 Cairo_1.6-2
                                                         cowplot_1.1.1
##
    [91] grid_4.4.1
                                 tidyr_1.3.1
                                                         colorspace_2.1-1
##
   [94] nlme_3.1-165
                                 patchwork_1.1.1
                                                         beeswarm_0.4.0
   [97] vipor_0.4.7
                                 cli_3.6.3
                                                         spatstat.sparse_3.1-0
## [100] fansi_1.0.6
                                 viridisLite_0.4.2
                                                         dplyr_1.1.4
## [103] uwot 0.2.2
                                 gtable 0.3.5
                                                         digest 0.6.37
## [106] progressr_0.14.0
                                 ggrepel_0.9.6
                                                         farver_2.1.2
## [109] htmlwidgets 1.5.3
                                 htmltools 0.5.8.1
                                                         lifecycle 1.0.4
## [112] httr_1.4.7
                                 mime_0.11
                                                         MASS_7.3-61
```