3.6 Tutorial-6: Differential Gene analysis in longitudinal data (CNP0001102)

This tutorial allows users to identify differential expressed genes in direction of time-points. As an example single cell data from Zhu et al. 2020 downloaded from here. Metadata is downloaded from table and can be found in the data. The dataset consists of 5 Covid-19 donors, 2 Flu donors with longitudinal data and 3 controls. To explore differential expressed gened in each celltype of each donor we used hurdle model based modeling on input data to retrieve the DEGs. To infer DEGs in each celltype towards time progression (timepoints considered as continuous if more than 2), please follow following steps.

3.6.1 Load data and clinical metadata (Time ~ 30sec)

Single cell object CNP0001102

```
#Single cell object CNP0001102
pbmc <- readRDS("data/CNP0001102_Final_nCoV_0716_upload.RDS")
#Add column Sample
pbmc@meta.data$Sample <- pbmc@meta.data$batch</pre>
```

3.6.2 Clinical annotations Table S1. Clinical data of the enrolled subjects (Time ~ 5sec)

```
metadata <- read.csv("data/CNP0001102-annotation.csv", stringsAsFactors = F)
row.names(metadata) <- metadata$Sample</pre>
```

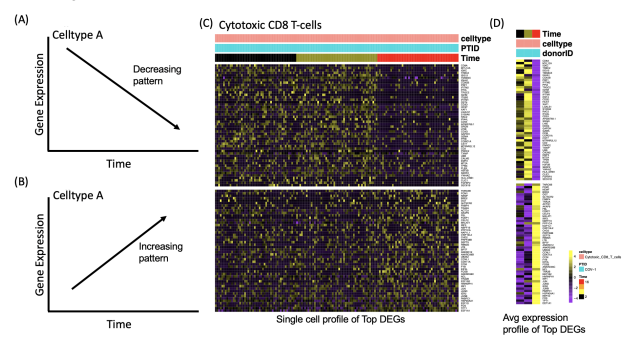
3.6.3 Load library and run (Time ~ 15 min)

```
#Load Library
library("PALMO")
library("tidyverse")
#Create PALMO object
palmo_obj <- createPALMOobject(anndata=metadata, data=pbmc)</pre>
#Assign Sample, PTID and Time parameters
palmo_obj <- annotateMetadata(data_object=palmo_obj,</pre>
                               sample column= "Sample",
                               donor_column= "Participant",
                               time_column= "Day")
#Sample overlap and final matrix
palmo_obj <- mergePALMOdata(data_object=palmo_obj, datatype="singlecell")</pre>
#Perform longitudinal differential analysis
palmo_obj <- sclongitudinalDEG(data_object=palmo_obj, scassay="RNA",</pre>
                                group_column="cell_type")
>Fitting a ZLM model for donorID: COV-1 ...
>Fitting a ZLM model for donorID: COV-2 ...
>Fitting a ZLM model for donorID: COV-3 ...
>Fitting a ZLM model for donorID: COV-4 ...
>Fitting a ZLM model for donorID: COV-5 ...
>Fitting a ZLM model for donorID: IAV-1 ...
>Fitting a ZLM model for donorID: IAV-2 ...
```

Check output folder for tabular results and visualization.

```
#Plots can be seen in output directory output
DEGres <- palmo_obj@result$degs</pre>
head(DEGres[order(DEGres$coef, decreasing = T),])
#primerid contrast
                                     coef
                                                   adjP donorID celltype dir
#IGHG4
         TimeD9 1.701579e-26 3.056092 1.453999e-23
                                                               Plasma upregulated at D9
                                                       IAV-2
#JCHAIN
          TimeD9 8.759407e-32 2.647757 2.245474e-28
                                                        IAV-2
                                                                Plasma upregulated at D9
         TimeD9 8.470810e-22 2.485250 2.412769e-19
                                                               Plasma upregulated at D9
#IGHG3
                                                       IAV-2
#IGLC2
         TimeD9 1.352490e-16 2.289544 8.890022e-15
                                                       IAV-2
                                                               Plasma upregulated at D9
#IGHG1
         TimeD9 1.292885e-16 2.219146 8.608601e-15
                                                       IAV-2
                                                               Plasma upregulated at D9
#SYNE2
         TimeD4 7.249010e-13 2.209541 1.215659e-09
                                                       COV-4 XCL+_NKs upregulated at D4
#Or interested celltypes
celltype oi <- c("Activated CD4 T cells", "Naive B cells")</pre>
palmo_obj <- sclongitudinalDEG(data_object=palmo_obj, scassay="RNA",</pre>
                            group_column="cell_type",
                            group_oi = celltype_oi)
```

General analysis schema and differential results in each donor over timepoints in celltype Cytotoxic CD8 T-cells using PALM shown below.



6 Session info

```
sessionInfo()
#> R version 4.0.3 (2020-10-10)
#> Platform: x86_64-apple-darwin17.0 (64-bit)
#> Running under: macOS Catalina 10.15.7
#> Matrix products: default
#> BLAS: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRblas.dylib
#> LAPACK: /Library/Frameworks/R.framework/Versions/4.0/Resources/lib/libRlapack.dylib
#> locale:
#> [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
#> attached base packages:
#> [1] grid
                stats graphics grDevices utils
                                                       datasets methods
#> [8] base
#> other attached packages:
#> [1] PALMO_0.1.0
#> loaded via a namespace (and not attached):
   [1] readxl_1.3.1
                                   backports 1.2.0
#>
    [3] circlize_0.4.11
                                  plyr_1.8.6
    [5] igraph_1.2.8
                                  lazyeval_0.2.2
                                 listenv_0.8.0
GenomeInfoDb_1.24.2
digest_0.6.28
#>
    [7] splines_4.0.3
    [9] scattermore_0.7
#> [11] ggplot2_3.3.5
                                 fansi\_0.5.0 tensor\_1.5
#> [13] htmltools 0.5.2
#> [15] magrittr_2.0.1
#> [17] cluster_2.1.0
                                  ROCR_1.0-11
                                 globals_0.14.0
#> [19] ComplexHeatmap_2.4.3
#> [21] readr_1.4.0
                                  modelr_0.1.8
#> [23] matrixStats 0.61.0
                                   colorspace 2.0-2
#> [25] rvest_0.3.6
                                   blob_1.2.1
#> [27] qqrepel_0.9.1
                                  haven_2.3.1
#> [29] xfun_0.25
                                  dplyr_1.0.7
                                 crayon_1.4.2
#> [31] RCurl_1.98-1.2
#> [33] jsonlite_1.7.2
                                  lme4_1.1-25
                                  spatstat.data\_2.1-0
#> [35] spatstat_1.64-1
#> [37] survival_3.2-7
                                  zoo_1.8-9
#> [39] qlue_1.5.0
                                  polyclip_1.10-0
#> [41] gtable_0.3.0
                                   zlibbioc_1.34.0
#> [43] XVector_0.28.0
                                   leiden 0.3.9
#> [45] DelayedArray_0.14.1 GetoptLong_1.0.4
#> [47] SingleCellExperiment_1.10.1 future.apply_1.8.1
#> [49] shape_1.4.5
                               BiocGenerics\_0.34.0
#> [51] abind_1.4-5
                                   scales_1.1.1
#> [53] pheatmap_1.0.12
                                   DBI_1.1.0
#> [55] miniUI_0.1.1.1
                                  Rcpp_1.0.7
                                 xtable\_1.8-4
#> [57] viridisLite 0.4.0
#> [59] clue_0.3-57
                                   reticulate_1.22
                               htmlwidgets\_1.5.4
#> [61] stats4_4.0.3
```

```
#> [63] httr_1.4.2
                                     RColorBrewer_1.1-2
#> [65] ellipsis_0.3.2
                                     Seurat_4.0.0
#> [67] factoextra_1.0.7.999
                                     ica_1.0-2
                                     pkgconfig_2.0.3
#> [69] farver_2.1.0
#> [71] uwot_0.1.10
                                     dbplyr_1.4.4
#> [73] deldir_1.0-6
                                     utf8_1.2.2
#> [75] tidyselect_1.1.1
                                     rlang_0.4.12
#> [77] reshape2_1.4.4
                                     later 1.3.0
#> [79] munsell_0.5.0
                                     cellranger_1.1.0
#> [81] tools_4.0.3
                                     generics_0.1.1
#> [83] broom_0.7.2
                                     ggridges_0.5.3
#> [85] evaluate_0.14
                                    stringr_1.4.0
#> [87] fastmap_1.1.0
                                     yaml_2.2.1
#> [89] goftest_1.2-3
                                    knitr_1.30
#> [91] fs_1.5.0
                                    fitdistrplus_1.1-6
#> [93] purrr_0.3.4
                                    RANN_2.6.1
#> [95] pbapply_1.5-0
                                     future_1.23.0
#> [97] nlme_3.1-149
                                     mime_0.12
#> [99] xml2_1.3.2
                                     compiler_4.0.3
#> [101] plotly_4.10.0
                                     png_0.1-7
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                                    reprex_0.3.0
#> [105] tweenr_1.0.1
                                     tibble_3.1.6
#> [107] statmod_1.4.35
                                     stringi_1.7.5
#> [109] forcats_0.5.0
                                     lattice_0.20-41
#> [111] Matrix 1.3-4
                                     nloptr 1.2.2.2
#> [113] vctrs 0.3.8
                                    pillar 1.6.4
#> [115] lifecycle_1.0.1
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#> [117] GlobalOptions_0.1.2
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#> [119] bitops_1.0-7
                                     data.table_1.14.2
#> [121] cowplot_1.1.1
                                     irlba_2.3.3
#> [123] GenomicRanges_1.40.0
                                     httpuv_1.6.3
#> [125] patchwork_1.1.1
                                     R6_2.5.1
#> [127] promises_1.2.0.1
                                     KernSmooth\_2.23-17
#> [129] qridExtra_2.3
                                     IRanges_2.22.2
#> [131] parallelly_1.28.1
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#> [133] boot_1.3-25
                                     MASS_7.3-53
#> [135] assertthat_0.2.1
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#> [137] MAST_1.14.0
                                     rjson_0.2.20
#> [139] SeuratObject_4.0.2
                                     sctransform_0.3.2
#> [141] GenomeInfoDbData_1.2.3
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#> [143] mgcv_1.8-33
                                     parallel_4.0.3
#> [145] hms_0.5.3
                                     rpart_4.1-15
#> [147] tidyverse_1.3.0
                                     tidyr_1.1.4
#> [149] minga_1.2.4
                                     rmarkdown_2.5
#> [151] Rtsne_0.15
                                     ggforce_0.3.2
#> [153] Biobase_2.48.0
                                     shiny_1.7.1
#> [155] lubridate_1.7.9
```