Trajectory Inference

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Overview

##

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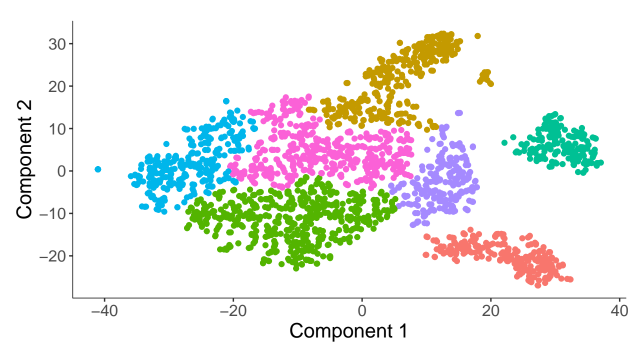
We are going to use 2300 PBMCs single-cells from mouse to construct the trajectory. These cells are selected form 8K PBMCs from 10X website based on the high expression of Cathepsin S (CTSS) gene. This gene is lysosomal cysteine proteinase that may participate in the degradation of antigenic proteins to peptides for presentation on MHC class II molecules. Hence, it is important to track the the regulatory effect of this gene in different cells.

First, load the object.

```
load("2K pbmc subset1.RData", verbose = FALSE)
and packages
suppressMessages(require(monocle))
suppressMessages(require(cellrangerRkit))
suppressMessages(require(dplyr))
We'll select the genes expressed in greater than 5% of the cells and order the cells based on genes.
my_cds_subset <- detectGenes(my_cds_subset, min_expr = 0.1)</pre>
fData(my_cds_subset)$use_for_ordering <- fData(my_cds_subset)$num_cells_expressed >
    0.05 * ncol(my_cds_subset)
# how many genes are used?
table(fData(my_cds_subset)$use_for_ordering)
##
## FALSE
          TRUE
    8641
          6805
We will now perform clustering but without specifying the number of clusters; we will use thresholds on the
cell's local density (rho) and nearest distance (delta) to determine the number of clusters.
my_cds_subset <- reduceDimension(my_cds_subset, max_components = 2, norm_method = "log",
    num_dim = 10, reduction_method = "tSNE", verbose = TRUE)
## Remove noise by PCA ...
## Reduce dimension by tSNE ...
my_cds_subset <- clusterCells(my_cds_subset, verbose = FALSE)</pre>
## Distance cutoff calculated to 3.679052
We'll use rho = 3.6 to cluster the cells again.
my_cds_subset <- clusterCells(my_cds_subset, rho_threshold = 3.6, delta_threshold = 10,
    skip_rho_sigma = T, verbose = FALSE)
table(pData(my_cds_subset)$Cluster)
##
         2
             3
                 4 5
                          6
                               7
```







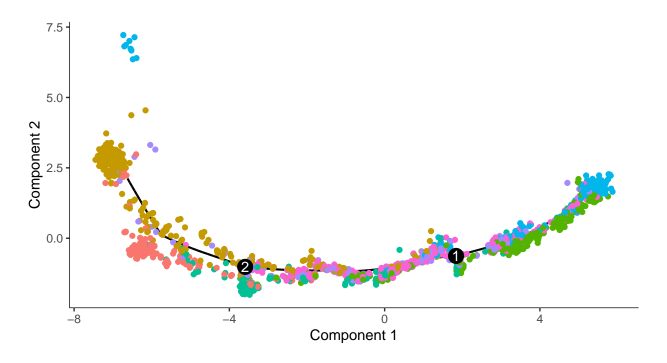
Now we'll perform the differential gene expression analysis as before but across all cell clusters.

We'll use the top 500 most significantly differentially expressed genes as the set of ordering genes and perform the dimension reduction and the trajectory analysis (using the orderCells() function).

```
my_ordering_genes <- row.names(clustering_DEG_genes)[order(clustering_DEG_genes$qval)][1:500]
my_cds_subset <- setOrderingFilter(my_cds_subset, ordering_genes = my_ordering_genes)
my_cds_subset <- reduceDimension(my_cds_subset, method = "DDRTree")
my_cds_subset <- orderCells(my_cds_subset)

plot_cell_trajectory(my_cds_subset, color_by = "Cluster")</pre>
```

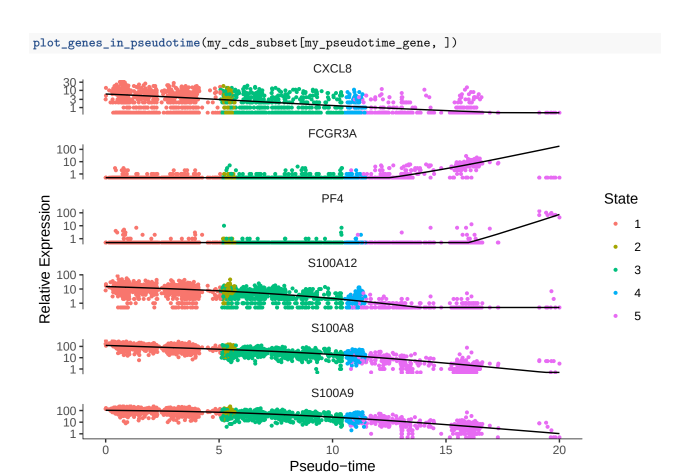




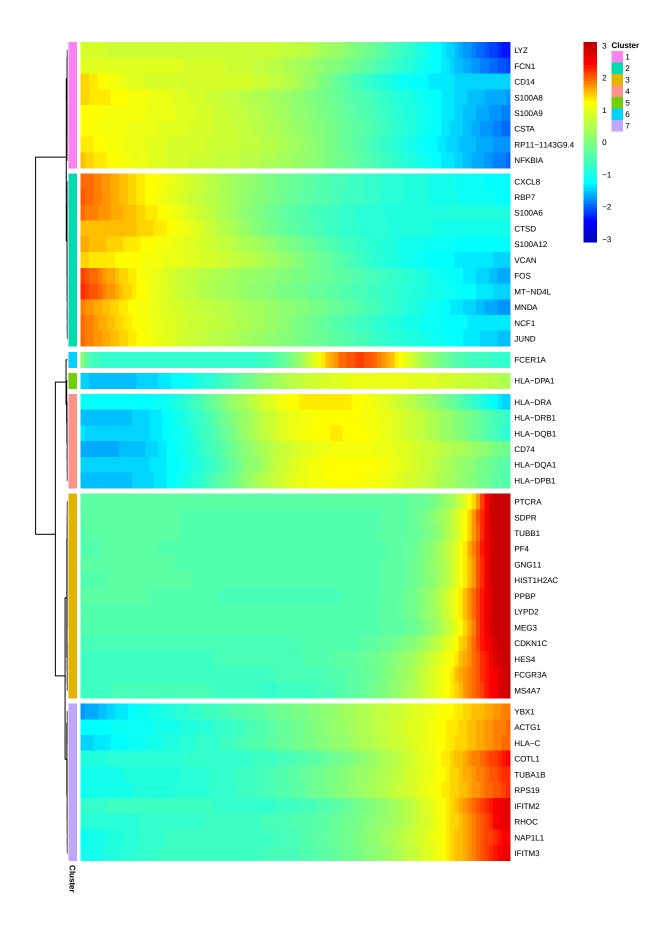
Finding Genes that Change as a Function of Pseudotime

Once we have a trajectory, we can use differential GeneTest() to find genes that have an expression pattern that varies according to pseudotime.

```
my_pseudotime_de <- differentialGeneTest(my_cds_subset, fullModelFormulaStr = "~sm.ns(Pseudotime)",
    cores = 8)
my_pseudotime_de %>% arrange(qval) %>% head()
##
                       family pval qval
                                                      id gene_short_name
     status
                                      0 ENSG00000163220
## 1
         OK negbinomial.size
                                                                  S100A9
## 2
         OK negbinomial.size
                                      0 ENSG00000163221
                                                                 S100A12
## 3
         OK negbinomial.size
                                 0
                                      0 ENSG00000143546
                                                                  S100A8
## 4
         OK negbinomial.size
                                 0
                                      0 ENSG00000203747
                                                                  FCGR3A
## 5
         OK negbinomial.size
                                 0
                                      0 ENSG00000169429
                                                                   CXCL8
## 6
                                      0 ENSG00000163737
         OK negbinomial.size
                                                                     PF4
     num_cells_expressed use_for_ordering
##
## 1
                    2285
                                      TRUE
                                      TRUE
## 2
                    1887
## 3
                    2270
                                      TRUE
## 4
                     377
                                      TRUE
## 5
                    1278
                                      TRUE
                      76
                                      TRUE
## 6
my_pseudotime_gene <- my_pseudotime_de %>% arrange(qval) %>% head() %>% select(id)
my_pseudotime_gene <- my_pseudotime_gene$id
```



Clustering Genes by Pseudotemporal Expression Pattern

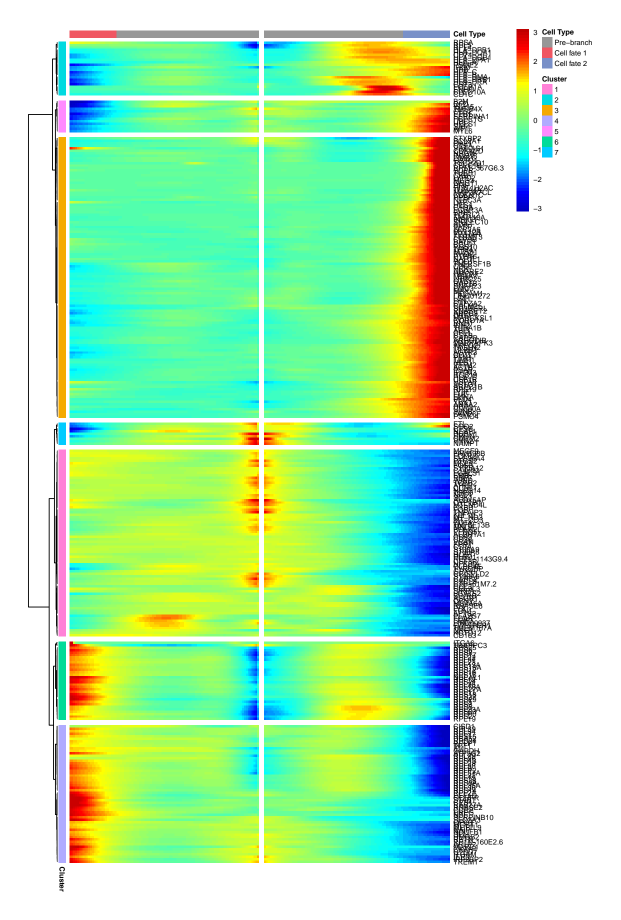


Analyzing Branches in Single-Cell Trajectories

Our trajectory has four branches, which represents cells that have alternative gene expression patterns. These represent cells that have supposedly gone through different developmental paths. We will now identify the genes that differ at a particular branch point. Here is the trajectory again.

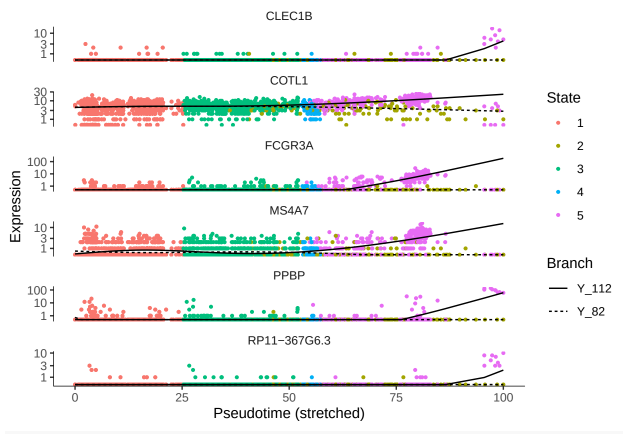
```
BEAM_res <- BEAM(my_cds_subset, branch_point = 1, cores = 8)
BEAM_res <- BEAM_res[order(BEAM_res$qval), ]
BEAM_res <- BEAM_res[, c("gene_short_name", "pval", "qval")]

# The heatmap shows how some genes are over-expressed or under-expressed
# depending on the trajectory path.
my_branched_heatmap <- plot_genes_branched_heatmap(my_cds_subset[row.names(subset(BEAM_res, qval < 1e-04)), ], branch_point = 1, num_clusters = 7, cores = 8, use_gene_short_name = TRUE, show_rownames = TRUE, return_heatmap = TRUE)</pre>
```



We can return genes that belong to specific clusters that were identified by BEAM().

```
head(my_branched_heatmap$annotation_row)
##
                 Cluster
## S100A9
                       1
## S100A8
                       1
## S100A12
                       1
## LYZ
                       1
## RP11-1143G9.4
                       1
## VCAN
dim(my_branched_heatmap$annotation_row)
## [1] 322
table(my_branched_heatmap$annotation_row$Cluster)
##
##
     1
         2
             3
                 4
                     5
                         6
                             7
## 76 22 114 56 13 32
                             9
my_row <- my_branched_heatmap$annotation_row</pre>
my_row <- data.frame(cluster = my_row$Cluster, gene = row.names(my_row), stringsAsFactors = FALSE)</pre>
head(my_row[my_row$cluster == 3, "gene"])
## [1] "FCGR3A"
                      "CLEC1B"
                                      "RP11-367G6.3" "COTL1"
## [5] "PPBP"
                      "MS4A7"
my_gene <- row.names(subset(fData(my_cds_subset), gene_short_name %in% head(my_row[my_row$cluster ==
    3, "gene"])))
# plot genes that are expressed in a branch dependent manner
plot_genes_branched_pseudotime(my_cds_subset[my_gene, ], branch_point = 1, ncol = 1)
```



sessionInfo()

```
## R version 3.5.0 (2018-04-23)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: macOS 10.14.2
##
## Matrix products: default
## BLAS: /Library/Frameworks/R.framework/Versions/3.5/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/3.5/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] splines
                  stats4
                            parallel stats
                                                graphics grDevices utils
    [8] datasets methods
##
                            base
##
## other attached packages:
##
   [1] dplyr_0.8.3
                             cellrangerRkit_2.0.0 Rmisc_1.5
##
   [4] plyr_1.8.4
                             lattice_0.20-38
                                                  bit64_0.9-7
   [7] bit_1.1-14
                             RColorBrewer_1.1-2
                                                  monocle_2.10.1
## [10] DDRTree_0.1.5
                             irlba_2.3.3
                                                  VGAM_1.1-1
  [13] ggplot2_3.2.1
                             Biobase_2.42.0
                                                  BiocGenerics_0.28.0
## [16] Matrix_1.2-17
##
## loaded via a namespace (and not attached):
   [1] viridis_0.5.1
                             viridisLite_0.3.0
                                                  assertthat_0.2.1
   [4] yaml_2.2.0
                             slam_0.1-45
                                                  ggrepel_0.8.1
```

## ## ## ## ## ## ## ## ##	[10] [13] [16] [19] [22] [25] [28] [31] [34] [40] [43] [46] [49] [52]	pillar_1.4.2 densityClust_0.3 fastICA_1.2-2 pheatmap_1.0.12 purrr_0.3.2 Rtsne_0.15 combinat_0.0-8 sparsesvd_0.2 crayon_1.3.4 tools_3.5.0 matrixStats_0.55.0 munsell_0.5.0 rlang_0.4.0 rstudioapi_0.10 rmarkdown_1.15 R6_2.4.0	glue_1.3.1 digest_0.6.21 htmltools_0.3.6 HSMMSingleCell_1.2.0 scales_1.0.0 proxy_0.4-23 docopt_0.6.1 lazyeval_0.2.2 evaluate_0.14 data.table_1.12.2 stringr_1.4.0 cluster_2.1.0 rhdf5_2.26.2 igraph_1.2.4.1 gtable_0.3.0 gridExtra_2.3	RANN_2.6.1 tibble_2.1.3 withr_2.1.2 magrittr_1.5 FNN_1.1.3 formatR_1.7 Rhdf5lib_1.4.3 compiler_3.5.0 grid_3.5.0 labeling_0.3 reshape2_1.4.3 knitr_1.25
##	[55]	R6_2.4.0 stringi_1.4.3 xfun_0.9	gridExtra_2.3 Rcpp_1.0.2	knitr_1.25 tidyselect_0.2.5