ChIMP Vignette

Courtney Schiebout, H. Robert Frost

Load Libraries

Libraries "CAMML" (Schiebout and Frost 2022) and "Seurat" (Satija et al. 2015) need to be loaded to carry out this vignette. Packages will also load additional libraries they depend on.

```
library(CAMML)
library(Seurat)
library(dplyr)
```

Data Processing

The following code outlines how the joint scRNA-seq/CITE-seq data from Lawlor, et al. (2021) (Lawlor et al. 2021), available on the 10X Genomics website, was processed for further analysis.

```
#load data
malt <- Read10X("raw_feature_bc_matrix/")</pre>
## 10X data contains more than one type and is being returned as a list containing matrices of each typ
#isolate the RNA data and make it a Seurat Object
malt.data <- malt$`Gene Expression`</pre>
seurat <- CreateSeuratObject(counts = malt.data, min.cells=10,min.features=100)</pre>
#filter for mitochondrial genes
seurat[["percent.mt"]] <- PercentageFeatureSet(seurat, pattern = "^MT-")</pre>
seurat <- subset(seurat, subset = percent.mt < 10)</pre>
#normalize and scale the RNA data
seurat <- NormalizeData(seurat)</pre>
## Normalizing layer: counts
seurat <- FindVariableFeatures(seurat, selection.method = "vst", nfeatures = 2000)</pre>
## Finding variable features for layer counts
seurat <- ScaleData(seurat)</pre>
## Centering and scaling data matrix
#cluster and visualize
seurat <- RunPCA(seurat)</pre>
## PC 1
## Positive: PCLAF, MKI67, RGS13, TYMS, MYBL2, CDK1, ZWINT, RRM2, UBE2C, AURKB
       TK1, GRN, PKM, TOP2A, BIRC5, ACTB, CCNB2, PHGDH, DHFR, LMO2
##
##
       NUF2, CST3, SPC25, CTSH, SERPINA9, ASPM, GTSE1, CDT1, SHCBP1, MAD2L1
```

Negative: ANXA1, GPR171, GZMK, CCL5, CD8A, SPRY1, GZMA, GTSCR1, RTKN2, TRGC2

```
BIRC5, TK1, SPC25, CDCA5, MYBL2, GTSE1, DHFR, NUF2, CDT1, CCNB2
       CDCA7, MAD2L1, SERPINA9, RMI2, ASPM, CD79A, GINS2, CHEK1, ASF1B, SHCBP1
##
## Negative: CEBPD, CST3, TNFAIP2, LYZ, TYROBP, CSF2RA, NDRG2, LGALS2, NECTIN2, FCER1G
       SERPINA1, RAB32, GOS2, ETS2, PLAUR, ALDH2, IFITM3, C15orf48, CXCL8, VEGFA
##
##
       ACO20656.1, CLEC7A, IL1B, CXCL2, PKP2, DST, AIF1, TIMP1, PLXDC2, CFP
## PC_ 3
## Positive: HLA-DRA, HLA-DQA1, HLA-DPA1, HLA-DQB1, HLA-DRB1, HLA-DPB1, CD74, IGHM, MS4A1, CD79A
       HLA-DMA, IGKC, LY9, TCF4, BASP1, FCRL5, BCL2A1, TNFRSF13B, CTSZ, CD22
##
##
       FTL, SWAP70, ID3, ITGAX, ARID3A, IGHA1, LDLRAD4, CTSH, SYNGR2, H3F3A
## Negative: ITM2A, IL32, ANXA1, MAF, GZMK, CTLA4, BATF, LDHB, TIGIT, HMGB2
       ICA1, MT2A, KLRB1, CORO1B, H2AFZ, CCL5, LDHA, NCOA7, GZMA, S100A4
##
       TNFRSF4, NKG7, GPR171, MAGEH1, PCLAF, GAPDH, CH25H, S100A10, PTPN13, ID2
## PC_ 4
## Positive: LYZ, TYROBP, LGALS2, FCER1G, CST3, ACO20656.1, AIF1, IL1B, MS4A6A, CSF2RA
       JAML, SERPINA1, CPVL, CFP, CD1E, AXL, GOS2, ITGAX, ID2, LST1
##
       NLRP3, CLEC7A, EREG, C1QA, DUSP4, C15orf48, CLEC10A, CD4, PLAUR, C1QB
##
## Negative: CD79A, MS4A1, IGHM, S100A16, DSP, TM4SF1, RBP1, HLA-DRA, FOXC1, S100A14
       CDC42EP1, EDN1, SOX9, NFIB, KRT7, ELF3, CD74, PITX1, ADIRF, TRIM29
       S100A2, KRT8, TJP1, CALD1, GABRP, PALMD, TACSTD2, RND3, MEIS2, MIA
##
## PC 5
## Positive: PCLAF, MKI67, RRM2, BIRC5, CDK1, AURKB, LYZ, ZWINT, UBE2C, TCL1A
       TOP2A, TK1, TYMS, LGALS2, SPC25, CDCA5, CST3, GTSE1, NUF2, CCNB2
       AIF1, ASPM, IGLC2, ESCO2, ACO20656.1, DHFR, Clorf162, MAD2L1, IL1B, PBK
##
## Negative: DUSP4, ARID3A, TNFRSF13B, SLAMF7, CPEB4, IGHA1, TRPV3, NEAT1, PDGFA, CLNK
       S100A4, LITAF, CD27, RAMP1, GUSB, RGS1, BCAR3, SPN, SQSTM1, UGCG
       CYTOR, FCRL4, ACY3, CD63, YWHAH, VOPP1, RAB11FIP1, BSG, ACP5, CLECL1
##
seurat <- FindNeighbors(seurat, dims = 1:10)</pre>
## Computing nearest neighbor graph
## Computing SNN
seurat <- FindClusters(seurat, resolution = 0.5)</pre>
\#\# Modularity Optimizer version 1.3.0 by Ludo Waltman and Nees Jan van Eck
##
## Number of nodes: 6438
## Number of edges: 216819
## Running Louvain algorithm...
## Maximum modularity in 10 random starts: 0.8886
## Number of communities: 11
## Elapsed time: 0 seconds
seurat <- RunUMAP(seurat, dims = 1:10)</pre>
## Warning: The default method for RunUMAP has changed from calling Python UMAP via reticulate to the R
## To use Python UMAP via reticulate, set umap.method to 'umap-learn' and metric to 'correlation'
## This message will be shown once per session
## 15:17:02 UMAP embedding parameters a = 0.9922 b = 1.112
## Found more than one class "dist" in cache; using the first, from namespace 'spam'
```

NKG7, CD40LG, KRT1, KLRD1, IFNG, CRTAM, CD8B, LINC02446, ITM2A, LYPD3

ITGA6, ID1, CDKN1C, KLRB1, TRDC, TRGC1, ALKAL2, KLRC2, LINCO1871, ENC1

Positive: PCLAF, MKI67, RRM2, ZWINT, CDK1, AURKB, UBE2C, TYMS, TOP2A, RGS13

##

PC 2

```
## Also defined by 'BiocGenerics'
## 15:17:02 Read 6438 rows and found 10 numeric columns
## 15:17:02 Using Annoy for neighbor search, n_neighbors = 30
## Found more than one class "dist" in cache; using the first, from namespace 'spam'
## Also defined by 'BiocGenerics'
## 15:17:02 Building Annoy index with metric = cosine, n_trees = 50
                      40
                           50
                                60
                                     70
                                          80
                                                    100%
## **************
## 15:17:02 Writing NN index file to temp file /var/folders/wv/91qlnj1571q8w6tn77wg10pr0000gp/T//Rtmpic
## 15:17:02 Searching Annoy index using 1 thread, search_k = 3000
## 15:17:04 Annoy recall = 100%
## 15:17:04 Commencing smooth kNN distance calibration using 1 thread with target n_neighbors = 30
## 15:17:06 Initializing from normalized Laplacian + noise (using RSpectra)
## 15:17:06 Commencing optimization for 500 epochs, with 266746 positive edges
## 15:17:15 Optimization finished
UMAPPlot(seurat, label = T)
     5
                                                                                0
     0
umap_2
                                                                                5
    -5
                                                                                6
                                                                                7
                                                                                8
                                                                                9
   -10
                                                                                10
                          -5
          -10
                                           0
                                                          5
                                    umap_1
```

scRNA-seq and CITE-seq Integration

Following the RNA data processing, the CITE-seq data needs to be added back into the data as an additional assay in the Seurat Object (Satija et al. 2015; Stoeckius et al. 2017). Since we filtered the data, the CITE-seq data needs aligned with the remaining cells.

Get Gene Sets and Run CAMML

Centering and scaling data matrix

In order to run CAMML and ChIMP, a gene set of cell types needs to be accessed. In the following code, "GetGeneSets" is used to load a pre-built gene set of 5 immune cell types. This can then be used to run CAMML. For this example HSCs will be removed.

```
#get gene sets
gene.set.df <- GetGeneSets(data = "immune.cells")

#filter out HSC
gene.set.df <- gene.set.df[-which(gene.set.df$cell.type == "HSC_CD34+"),]
gene.set.df</pre>
```

```
##
      gene.symbol cell.type gene.weight
                                              ensembl.id
## 1
            ABCB4
                     B_cell
                               5.929227 ENSG00000005471
## 3
             AIM2
                     B cell
                               5.750254 ENSG00000163568
## 5
            BANK1
                     B_cell
                               7.942142 ENSG00000153064
## 8
              BLK
                     B_cell
                               7.548835 ENSG00000136573
                     B cell
## 9
            BTNL9
                               5.071020 ENSG00000165810
## 12
             CD19
                     B cell
                               8.853964 ENSG00000177455
                     B_cell
                               6.364610 ENSG00000012124
## 13
             CD22
## 19
            CD79A
                     B_cell
                               8.582138 ENSG00000105369
## 24
            CPNE5
                     B_cell
                               6.580999 ENSG00000124772
## 30
             E2F5
                     B_cell
                               7.159631 ENSG00000133740
                     B_cell
## 32
            FCRL1
                               8.492002 ENSG00000163534
## 33
            FCRL2
                     B_cell
                               6.669676 ENSG00000132704
## 34
            FCRLA
                     B_cell
                               8.898029 ENSG00000132185
## 38
          HLA-DOB
                     B_cell
                               7.261022 ENSG00000241106
## 55
        LINC00926
                     B_cell
                               5.223783 ENSG00000247982
                               7.927719 ENSG00000156738
## 60
            MS4A1
                     B_cell
```

```
## 66
            P2RX5
                      B cell
                                5.862599 ENSG00000083454
## 69
          PKHD1L1
                      B cell
                                6.031826 ENSG00000205038
                      B cell
##
  70
          PLEKHG1
                                5.831675 ENSG00000120278
             PNOC
                      B_cell
                                7.489833 ENSG00000168081
##
  71
##
   74
          RALGPS2
                      B cell
                                5.228555 ENSG00000116191
  85
                      B cell
##
             SPIB
                                6.160680 ENSG00000269404
                      B cell
## 86
            STAP1
                                5.787053 ENSG00000035720
## 92
            TLR10
                      B cell
                                7.109604 ENSG00000174123
##
  2
           ADGRE2
                   Monocyte
                                7.729049 ENSG00000127507
## 11
            C5AR1
                    Monocyte
                                8.813094 ENSG00000197405
## 14
          CD300LF
                    Monocyte
                                6.470491 ENSG00000186074
## 18
             CD68
                   Monocyte
                                5.983464 ENSG00000129226
##
  22
           CDKN1C
                   Monocyte
                                5.755567 ENSG00000129757
## 25
           CPPED1
                                5.344299 ENSG00000103381
                    Monocyte
## 27
            CSF1R
                                6.984429 ENSG00000182578
                   Monocyte
## 28
           CXCL16
                    Monocyte
                                6.330266 ENSG00000161921
## 36
              HCK
                   Monocyte
                                7.334923 ENSG00000101336
                    Monocyte
##
   37
              HK3
                                5.814265 ENSG00000160883
## 51
                                6.479104 ENSG00000104974
           LILRA1
                    Monocyte
## 52
           LILRA2
                    Monocyte
                                6.321271 ENSG00000239998
## 53
           LILRA5
                   Monocyte
                                5.485005 ENSG00000187116
## 54
           LILRB2
                    Monocyte
                                8.092027 ENSG00000131042
## 57
             LST1
                    Monocyte
                                5.607879 ENSG00000204482
            MS4A7
                                6.592873 ENSG00000166927
## 61
                    Monocyte
## 62
             MSR1
                    Monocyte
                                6.644375 ENSG00000038945
  67
           PAPSS2
                   Monocyte
                                5.102467 ENSG00000198682
  68
            PILRA
                                7.658938 ENSG00000085514
##
                   Monocyte
##
   77
         SERPINA1
                    Monocyte
                                6.364474 ENSG00000197249
## 80
          SLC31A2
                   Monocyte
                                5.431567 ENSG00000136867
## 81
           SLC7A7
                    Monocyte
                                5.888768 ENSG00000155465
          SMPDL3A
## 83
                    Monocyte
                                5.007270 ENSG00000172594
## 84
             SPI1
                    Monocyte
                                5.337993 ENSG00000066336
##
  87
           TBC1D8
                    Monocyte
                                5.634734 ENSG00000204634
                                5.145335 ENSG00000059377
##
  88
           TBXAS1
                    Monocyte
##
  97
             VMO1
                    Monocyte
                                7.072633 ENSG00000182853
                                7.119036 ENSG00000169583
##
  23
                     NK_cell
            CLIC3
## 31
            FASLG
                     NK cell
                                5.610501 ENSG00000117560
## 40
          IL18RAP
                     NK_cell
                                5.080865 ENSG00000115607
## 43
          KIR2DL4
                     NK_cell
                                5.716879 ENSG00000189013
                     NK_cell
## 44
          KIR3DL1
                                6.315612 ENSG00000167633
          KIR3DL2
                     NK cell
                                5.439694 ENSG00000240403
  45
## 46
            KLRF1
                     NK cell
                                5.481712 ENSG00000150045
            KRT86
                     NK_cell
##
  48
                                5.458481 ENSG00000170442
##
  73
            PRR5L
                     NK_cell
                                5.241528 ENSG00000135362
## 76
            S1PR5
                     NK_cell
                                5.062385 ENSG00000180739
## 78
           SH2D1B
                     NK_cell
                                7.356306 ENSG00000198574
## 82
           SLFN13
                     NK_cell
                                5.210182 ENSG00000154760
## 98
             XCL1
                     NK_cell
                                6.650246 ENSG00000143184
## 99
             YES1
                     NK_cell
                                5.168530 ENSG00000176105
## 6
           BCL11B
                     T_cells
                                5.564075 ENSG00000127152
## 15
             CD3D
                     T_cells
                                6.559457 ENSG00000167286
             CD3E
## 16
                     T_cells
                                5.457156 ENSG00000198851
## 17
             CD3G
                     T_cells
                                7.904570 ENSG00000160654
## 20
             CD8A
                     T cells
                                5.715826 ENSG00000153563
```

```
## 29
             DPP4
                    T_cells
                                5.142898 ENSG00000197635
## 39
                                6.000820 ENSG00000163600
             ICOS
                    T cells
           INPP4B
## 41
                    T_cells
                                5.664718 ENSG00000109452
## 42
              ITK
                    T cells
                                5.013501 ENSG00000113263
## 47
            KLRG1
                    T cells
                                5.244996 ENSG00000139187
## 50
             LEF1
                    T cells
                                5.500641 ENSG00000138795
## 56
            LRRN3
                    T cells
                                7.577637 ENSG00000173114
## 58
              MAL
                    T_cells
                                7.694320 ENSG00000172005
## 63
            NELL2
                    T_cells
                                6.411428 ENSG00000184613
## 65
           OXNAD1
                    T_cells
                                5.250065 ENSG00000154814
## 75
           RNF157
                    T_cells
                                5.376098 ENSG00000141576
## 79
            SIRPG
                    T_cells
                                5.538280 ENSG00000089012
                                5.002518 ENSG00000165929
## 89
             TC2N
                    T_cells
## 90
             TCF7
                    T_cells
                                5.961207 ENSG00000081059
## 91
           THEMIS
                    T_cells
                                6.889632 ENSG00000172673
## 93
          TRABD2A
                    T_cells
                                5.447930 ENSG00000186854
## 94
             TRAC
                    T cells
                                5.025658 ENSG00000277734
## 95
            TRAT1
                    T_cells
                                7.189648 ENSG00000163519
## 96
          UBASH3A
                    T cells
                                5.305995 ENSG00000160185
#run CAMML
seurat <- CAMML(seurat, gene.set.df)</pre>
## Computing VAM distances for 4 gene sets, 6438 cells and 15518 genes.
## Min set size: 11, median size: 23.5
## Warning: Feature names cannot have underscores ('_'), replacing with dashes
## ('-')
## Warning: Feature names cannot have underscores ('_'), replacing with dashes
## ('-')
## Warning: Feature names cannot have underscores ('_'), replacing with dashes
## Warning: Feature names cannot have underscores ('_'), replacing with dashes
## ('-')
## Warning: Key 'vamcdf ' taken, using 'camml ' instead
```

7.828488 ENSG00000172116

T_cells

CD8B

21

Integrate CITE-seq via ChIMP into CAMML with k-means Disretization

Following the running of CAMML, the cell type scores can be altered by the inclusion of CITE-seq data using ChIMP. In order to use this, a list of cell types and their corresponding CITE-seq markers needs to be built. This list, the Seurat Object, and a vector of booleans will then be fed into ChIMP. The vector serves to designate whether, in cases of multiple marker proteins in a cell type, any marker protein can be present to maintain the CAMML score or if ChIMP should require all marker proteins to be present to maintin the CAMML score.

In other words, if a user designates both CD4 and CD8 for T cells, anyMP=TRUE would require that just one of the two markers be present in a cell for the cell to have a nonzero cell type score. However, if anyMP=FALSE, both markers would have to be present in a cell for the cell type score to be nonzero.

In this example, we use anyMP=FALSE for monocytes to single out cells that are positive for both CD14 and CD16. We use anyMP=TRUE to select for T cells that are either CD4 or CD8 positive.

```
#compare ADT markers and cell types
rownames(seurat@assays$ADT)
  [1] "CD3-TotalSeqB"
                                  "CD4-TotalSeqB"
##
   [3] "CD8a-TotalSeqB"
                                  "CD14-TotalSeqB"
  [5] "CD15-TotalSeqB"
                                  "CD16-TotalSeqB"
##
  [7] "CD56-TotalSeqB"
                                  "CD19-TotalSeqB"
## [9] "CD25-TotalSeqB"
                                  "CD45RA-TotalSeqB"
## [11] "CD45RO-TotalSeqB"
                                  "PD-1-TotalSeqB"
## [13] "TIGIT-TotalSeqB"
                                  "CD127-TotalSeqB"
## [15] "IgG2a-control-TotalSeqB" "IgG1-control-TotalSeqB"
## [17] "IgG2b-control-TotalSeqB"
rownames(seurat@assays$CAMML)
## [1] "B-cell"
                  "Monocyte" "NK-cell" "T-cells"
#create CITE list
markers <- cbind(c(rownames(seurat),rownames(seurat)[2],rownames(seurat)[4]),</pre>
      (rownames(seurat@assays$ADT)[c(8,4,7,2,6,3)]))
citelist <- list()</pre>
for (i in 1:length(rownames(seurat))){
  citelist[[i]] <- markers[which(markers[,1]==rownames(seurat)[i]),2]</pre>
}
names(citelist) <- rownames(seurat)</pre>
citelist
## $`B-cell`
## [1] "CD19-TotalSeqB"
## $Monocyte
## [1] "CD14-TotalSeqB" "CD16-TotalSeqB"
##
## $`NK-cell`
## [1] "CD56-TotalSeqB"
##
## $`T-cells`
## [1] "CD4-TotalSeqB" "CD8a-TotalSeqB"
#run ChIMP
seuratk <- ChIMP(seurat, citelist, anyMP = c(T,F,T,T),</pre>
                 greater = rep(T, length(unlist(citelist))))
#visualize the cell type scores
seurat.markers = FindAllMarkers(seuratk, assay="ChIMP", only.pos = TRUE)
## Calculating cluster 0
## For a (much!) faster implementation of the Wilcoxon Rank Sum Test,
## (default method for FindMarkers) please install the presto package
## install.packages('devtools')
## devtools::install_github('immunogenomics/presto')
## -----
## After installation of presto, Seurat will automatically use the more
```

```
## efficient implementation (no further action necessary).
## This message will be shown once per session
## Calculating cluster 1
## Calculating cluster 2
## Calculating cluster 3
## Calculating cluster 4
## Calculating cluster 5
## Warning in FindMarkers.default(object = data.use, slot = data.slot, counts =
## counts, : No features pass logfc.threshold threshold; returning empty
## data.frame
## Calculating cluster 6
## Calculating cluster 7
## Calculating cluster 8
## Calculating cluster 9
## Calculating cluster 10
DefaultAssay(object = seuratk) = "ChIMP"
top.pathways <- seurat.markers %>% group_by(cluster) %>% top_n(n = 3, wt = avg_log2FC)
DoHeatmap(seuratk, slot="data", features = top.pathways$gene,
          size=2, label=T, raster = F) + NoLegend()
                                                                                 1 890
                                                                               6
  B-cell
```



Integrate CITE-seq via ChIMP into CAMML with Quantile Disretization

This example follows the same pipeline as the above example but uses a median discretization for CITE-seq instead of k-means for comparison.

```
#compare ADT markers and cell types
rownames(seurat@assays$ADT)
```

```
## [1] "CD3-TotalSeqB"
                                   "CD4-TotalSeqB"
## [3] "CD8a-TotalSeqB"
                                   "CD14-TotalSeqB"
## [5] "CD15-TotalSeqB"
                                   "CD16-TotalSeqB"
## [7] "CD56-TotalSeqB"
                                   "CD19-TotalSeqB"
## [9] "CD25-TotalSeqB"
                                   "CD45RA-TotalSeqB"
## [11] "CD45RO-TotalSeqB"
                                   "PD-1-TotalSeqB"
## [13] "TIGIT-TotalSeqB"
                                   "CD127-TotalSeqB"
## [15] "IgG2a-control-TotalSeqB" "IgG1-control-TotalSeqB"
## [17] "IgG2b-control-TotalSeqB"
rownames(seurat@assays$CAMML)
## [1] "B-cell"
                  "Monocyte" "NK-cell" "T-cells"
#create CITE list
markers <- cbind(c(rownames(seurat),rownames(seurat)[2],rownames(seurat)[4]),</pre>
      (rownames(seurat@assays$ADT)[c(8,4,7,2,6,3)]))
citelist <- list()</pre>
for (i in 1:length(rownames(seurat))){
  citelist[[i]] <- markers[which(markers[,1]==rownames(seurat)[i]),2]</pre>
names(citelist) <- rownames(seurat)</pre>
citelist
## $`B-cell`
## [1] "CD19-TotalSeqB"
##
## $Monocyte
## [1] "CD14-TotalSeqB" "CD16-TotalSeqB"
##
## $`NK-cell`
## [1] "CD56-TotalSeqB"
##
## $`T-cells`
## [1] "CD4-TotalSeqB" "CD8a-TotalSeqB"
#run ChIMP
seuratq <- ChIMP(seurat, citelist, method = "q", anyMP = c(T,F,T,T),</pre>
                 greater = rep(T, length(unlist(citelist))))
#visualize the cell type scores
seurat.markers = FindAllMarkers(seuratq, assay="ChIMP", only.pos = TRUE)
## Calculating cluster 0
## Calculating cluster 1
## Calculating cluster 2
## Calculating cluster 3
## Calculating cluster 4
## Calculating cluster 5
## Warning in FindMarkers.default(object = data.use, slot = data.slot, counts =
## counts, : No features pass logfc.threshold threshold; returning empty
## data.frame
## Calculating cluster 6
```



References

Calculating cluster 7

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