Introduction to CytoDx

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Introduction

CytoDx is a method that predicts clinical outcomes using single cell data without the need of cell gating. It first predicts the association between each cell and the outcome using a linear statistical model (Figure 1). The cell level predictions are then averaged within each sample to represent the sample level predictor. A second model is used to make prediction at the sample level (Figure 1). Compare to traditional gating based methods, CytoDX have multiple advantages.

- 1. Robustness. CytoDx is able to robustly predict clinical outcomes using data acquired by different cytometry platforms in different research institutes.
- 2. Interpretability. CytoDx identifies cell markers and cell subsets that are most associated with the clinical outcome, allowing researchers to interpret the result easily.
- 3. Simplicity. CytoDx associates cytometry data with clinical features without any cell gating steps, therefore simplifying the diagnostic process in clinical settings.

In section 2, we demonstrate how to install CytoDx.

In section 3, we show an example where CytoDx is used to diagnose acute myeloid leukemia (AML). In addition to perform diagnosis, we also show that CytoDx can be used to identify the cell subsets that are associated with AML.





Installation

You can install the stable version of CytoDx from Bioconductor: BiocManager::install("CytoDx")

You can also install the latest develop version of CytoDx from github: devtools::install_github("hzc363/CytoDx")

Example: diagnosing AML using flow cytometry

In this example, we build a CytoDx model to diagnose acute myeloid leukemia (AML) using flow cytometry data. We train the model using data from 5 AML patients and 5 controls and test the performance in a test dataset.

Step 1: Prepare data

The CytoDx R package contains the fcs files and the ground truth (AML or normal) that are needed for our example. We first load the ground truth.

library(CytoDx)

```
# Find data in CytoDx package
path <- system.file("extdata",package="CytoDx")
# read the ground truth
fcs_info <- read.csv(file.path(path,"fcs_info.csv"))
# print out the ground truth
knitr::kable(fcs_info)</pre>
```

| fcsName | Label | dataset |
|---------------|----------------------|------------------------|
| sample1.fcs | aml | test |
| sample2.fcs | aml | test |
| sample3.fcs | aml | test |
| sample4.fcs | aml | test |
| sample5.fcs | aml | test |
| sample6.fcs | normal | test |
| sample 7. fcs | normal | test |
| sample 8. fcs | normal | test |
| sample9.fcs | normal | test |
| sample10.fcs | normal | test |
| sample11.fcs | aml | train |
| sample12.fcs | aml | train |
| sample13.fcs | aml | train |
| sample14.fcs | aml | train |
| sample15.fcs | aml | train |
| sample16.fcs | normal | train |
| sample17.fcs | normal | train |
| sample18.fcs | normal | train |
| sample19.fcs | normal | train |
| sample20.fcs | normal | train |

We then read the cytometry data for training samples using the fcs2DF function.

```
# Find the training data
train_info <- subset(fcs_info,fcs_info$dataset=="train")
# Specify the path to the cytometry files
fn <- file.path(path,train_info$fcsName)
# Read cytometry files using fcs2DF function
train_data <- fcs2DF(fcsFiles=fn,</pre>
```

```
y=train_info$Label,
assay="FCM",
b=1/150,
excludeTransformParameters=
c("FSC-A","FSC-W","FSC-H","Time"))
```

The CytoDx is flexible to data transformations. It can be applied to rank transformed data to reduce batch effects. Here, we transform the original data to rank data.

```
# Perfroms rank transformation
x_train <- pRank(x=train_data[,1:7],xSample=train_data$xSample)
# Convert data frame into matrix. Here we included the 2-way interactions.</pre>
```

Step 2: Build CytoDx model

We use training data to build a predictive model.

x_train <- model.matrix(~.*.,x_train)</pre>

Step 3: Predict AML using testing data

We first load and rank transform the test data.

We use the built CytoDx model to predict AML.

```
# Predict AML using CytoDx.ped function
pred <- CytoDx.pred(fit,xNew=x_test,xSampleNew=test_data$xSample)</pre>
```

We plot the prediction. In this example, CytoDx classifies the sample into AML and normal perfectly.





We use a decision tree to find cell subsets that are associated the AML. In this step, the original cytometry data should be used, rather than the ranked data.



Session Infomation

sessionInfo()

R version 4.3.0 RC (2023-04-13 r
84269) Platform: x86_64-pc-linux-gnu (64-bit) Running under: Ubuntu 22.04.2 LTS

 $Matrix \ products: \ default \ BLAS: \ /home/biocbuild/bbs-3.17-bioc/R/lib/libRblas.so \ LAPACK: \ /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0$

locale: [1] LC_CTYPE=en_US.UTF-8 LC_NUMERIC=C

- [3] LC_TIME=en_GB LC_COLLATE=C
- 5 LC_MONETARY=en_US.UTF-8 LC_MESSAGES=en_US.UTF-8
- [7] LC_PAPER=en_US.UTF-8 LC_NAME=C
- [9] LC_ADDRESS=C LC_TELEPHONE=C

[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

time zone: America/New_York tzcode source: system (glibc)

attached base packages: [1] stats graphics grDevices utils datasets methods base

other attached packages: [1] CytoDx_1.20.0

loaded via a namespace (and not attached): [1] Matrix_1.5-4 glmnet_4.1-7 highr_0.10

- [4] dplyr_1.1.2 compiler_4.3.0 rpart_4.1.19
- [7] tidyselect_1.2.0 Rcpp_1.0.10 Biobase_2.60.0
- [10] cytolib_2.12.0 parallel_4.3.0 splines_4.3.0
- [13] yaml_2.3.7 fastmap_1.1.1 lattice_0.21-8
- [16] R6_2.5.1 RProtoBufLib_2.12.0 generics_0.1.3
- [19] rpart.plot_3.1.1 shape_1.4.6 knitr_1.42
- [22] BiocGenerics_0.46.0 iterators_1.0.14 tibble_3.2.1
- [25] pillar_1.9.0 rlang_1.1.0 utf8_1.2.3
- [28] flowCore_2.12.0 xfun_0.39 doParallel_1.0.17
- [31] cli_3.6.1 withr_2.5.0 magrittr_2.0.3
- [34] digest 0.6.31 foreach 1.5.2 grid 4.3.0
- [37] lifecycle_1.0.3 S4Vectors_0.38.0 vctrs_0.6.2
- [40] evaluate_0.20 glue_1.6.2 codetools_0.2-19
- [43] survival_3.5-5 stats4_4.3.0 fansi_1.0.4

- [46] rmarkdown_2.21 matrixStats_0.63.0 tools_4.3.0 [49] pkgconfig_2.0.3 htmltools_0.5.5