# Microarray Gene Expression Analysis

Acute myeloid leukemia (AML)

Saman Dehestani \* Sajede Fadaei † Aminreza Sefid ‡

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# 1 Microarray

A microarray is a laboratory tool used to detect the expression of genes from a sample at the same time.

Microarrays have thousands of tiny spots in defined positions, with each spot containing a known DNA sequence or gene. The DNA molecules attached to each slide act as probes to detect gene expression.

To perform a microarray, mRNA molecules are typically collected from two different groups. For example, one group is considered as reference, which includes healthy samples and another group is experimental samples collected from a disease individual such as cancer.

The two mRNA samples are then converted into complementary DNA (cDNA) and each sample is labeled with a fluorescent probe of a different color. For instance, the experimental cDNA may be labeled with a red fluorescent dye, whereas the reference cDNA may be labeled with a green fluorescent dye. The two samples are mixed together and allowed to bind to the microarray slide. cDNA molecules bind to the DNA probes on the slides is called hybridization. After hybridization, the microarray is scanned to measure the expression of each gene printed into the slide. The red spots show that the number of experimental samples is higher than the reference sample. If the expression of a particular gene is lower in the experimental sample than in the reference sample, then the corresponding spot on the microarray appears green. Finally, if there is an equal expression in the two samples, then the spot appears yellow.

The microarray output data is in the form of a matrix table, which is finally in three formats SOFT, MINiML and TXT saved. Raw data is provided as a supplementary file.

<sup>\*401208226</sup> 

<sup>†400211513</sup> 

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# 2 Loading libraries

```
[]: library(GEOquery)
  library(limma)
  library(umap)
  library(pheatmap)
  library(ggplots)
  library(ggplot2)
  library(reshape2)
  library(plyr)
  library(repr)
  library(gridExtra)
  library(ggpubr)
  library(Rtsne)
  library(MASS)
  library(FactoMineR)
  library(factoextra)
```

# 3 Loading the GSE48558 series

The getGEO() method from the GEOquery package is used in order to download the GEO SOFT format of the GSE48558 and then parse it to the R structure obtaining the GSE Matrix and the GPL annotations:

```
[2]: gset <- getGEO("GSE48558", GSEMatrix =TRUE, getGPL=T, destdir='../Data/')
    gset <- gset[[1]]

Found 1 file(s)

GSE48558_series_matrix.txt.gz

Using locally cached version: ../Data//GSE48558_series_matrix.txt.gz

Using locally cached version of GPL6244 found here:
    ../Data//GPL6244.soft.gz</pre>
[3]: dim(gset)
```

Features

32321 Samples

170

# 4 Selecting proper samples

The source name column in the GSE48558, stands for the source of the biological material of the sample<sup>1</sup>. There are 11 different sample srources in this data series, including:

- AML cell line
- AML patient
- B ALL cell line
- B ALL patient
- B normal
- T ALL cell line
- T ALL patient
- T normal
- granulocytes normal
- monocytes normal
- CD34+ normal

We will only select the Leukemia AML samples and those with the normal phenotype value. The selection and the grouping were done first in the GEO2R analysis tool provided by *ncbi* website itself. A string will be generated, which can be used to select and group the samples in the R:

Samples are grouped according to the following rules:

- All the AML samples are grouped together.
- Normal samples are grouped based on their source\_name (B, T, granulocytes, monocytes, and CD34+).

Each number in the provided string stands for a group, and the X character means the corresponding sample won't be selected.

Next we will create a factor from these class numbers and add the group column to the gset.

<sup>&</sup>lt;sup>1</sup>https://www.ncbi.nlm.nih.gov/geo/info/qqtutorial.html

# 5 Data Quality Control

#### 5.1 Normalization

#### 5.1.1 Log Transformation

A technique that we can perform to scale the data properly is the *log transformation*. Our data may contain huge feature values that are unevenly distributed. The easiest solution is to scale them down using the log operation.

But before that, we'll check if we need to perform the *log transformation* or not. For this purpose we will simply find the range of feature values to find out if they are already log-scaled or not:

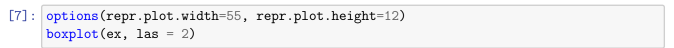
```
[6]: ex <- exprs(gset) # Biobase.exprs() returns a matrix of expression values print(c(min(ex), max(ex)))
```

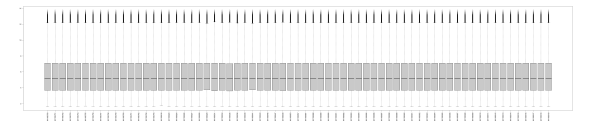
#### [1] 1.611473 13.761536

According to this, min value is 1.6 and max value is 13.76, so they are logarithmic already and data is normalized.

#### 5.1.2 Plotting the *expression* matrix

We can also plot the data to check if any normalization process is needed. If samples vary in range and scale, it's imperative to perform the normalization:





As the plot represents, samples are well-scaled with approximately identical  $5 number summary^2$  values.

<sup>&</sup>lt;sup>2</sup>minimum, lower-hinge, median, upper-hinge, maximum

# 5.2 Dimensionality Reduction

First, let's check our data dimensions again:

[8]:	<pre>dim(gset)</pre>					
]	Features	32321 Samples	67			

As you see, we've got a 32321-dimensional data, which is enormous! What we're about to do, is reduce the dimensions of our data while keeping as much variation in the original data as possible.

#### but How exactly is this dimensionality reduction going to help us?

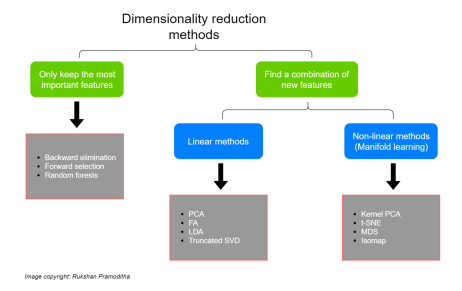
- 1. Lower dimensionality means less computational resources, less training time, and more performance (because in a high-dimensional space, most data points are likely far from each other, and it's hard for the model to train effectively on such data curse of dimensionality).
- 2. When there are many features, the model tends to become more complex, resulting in over-fitting. So the *dimensionality reduction* avoids the problem of **overfitting**.
- 3. Imagine you want to **visualize** this 323321-dimentional data! It's not that practical, is it? But when you reduce the dimensionality of higher dimension data to 2 or 3 components, it can easily be plotted on a 2D or 3D plot.

#### 4. multicollinearity

It occurs when features are highly correlated with one or more of the other features in the dataset and affect the performance of regression and classification models. *Dimensionality reduction* takes advantage of multicollinearity and combines the highly correlated variables into a set of uncorrelated variables.

5. When you keep the essential features and remove the others, the data **noise** is more likely to be removed, leading to higher model accuracy.

You can see different methods of dimensionality reduction in the diagram below:



## 5.2.1 Principal Component Analysis (PCA)

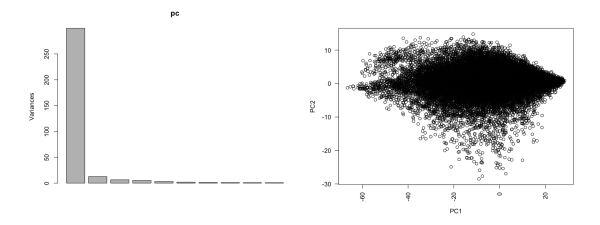
The first dimension reduction method that we're going to use is the Principal Component Analysis (PCA). Principal component analysis is used to extract the important information from a multivariate data table and to express this information as a set of few new variables called principal components. PCA assumes that the directions with the largest variances are the most **important** and the amount of variance retained by each principal component is measured by the **eigenvalue**. We will use the **prcomp()** method to obtain the PCA and then plot it:

```
[9]: pc <- prcomp(ex)
get_eigenvalue(pc)[1:5, ]</pre>
```

		eigenvalue <dbl></dbl>	variance.percent <dbl></dbl>	cumulative.variance.percent <dbl></dbl>
•	Dim.1	299.598423	87.2509394	87.25094
A data.frame: $5 \times 3$	Dim.2	12.643203	3.6820333	90.93297
	Dim.3	6.430822	1.8728244	92.80580
	Dim.4	5.144998	1.4983587	94.30416
	Dim.5	3.140644	0.9146382	95.21879

As you can see, about 90.93 of the variance is retained by the first two *principal components*. Before going further, let's first plot the data according to PC1 and PC2:

```
[10]: options(repr.plot.width=15, repr.plot.height=6)
    par(mfrow = c(1, 2))
    p1 <- plot(pc)
    p2 <- plot(pc$x[, 1:2], las = 2)</pre>
```



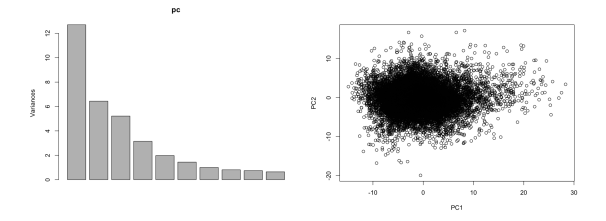
The problem is, PC1 seems to present a misleading variance in the data points, acquired probably by the uninteresting expression differences between genes which often expressed (e.g. housekeeping genes) and those expressed rarely.

To overcome this problem, we are going to center<sup>3</sup> our data columns using the scale() method:

```
[11]: ex.scale <- t(scale(t(ex), scale = F))
   pc <- prcomp(ex.scale)
   get_eigenvalue(pc)[1:5, ]</pre>
```

```
eigenvalue
                                         variance.percent
                                                           cumulative.variance.percent
                             <dbl>
                                         <dbl>
                                                           <dbl>
                     Dim.1
                             12.698222
                                         28.876999
                                                           28.87700
A data.frame: 5 \times 3 Dim.2
                             6.434864
                                         14.633510
                                                           43.51051
                     Dim.3
                             5.211775
                                         11.852086
                                                           55.36259
                     Dim.4
                             3.142359
                                         7.146032
                                                           62.50863
                     Dim.5
                             1.972313
                                         4.485232
                                                           66.99386
```

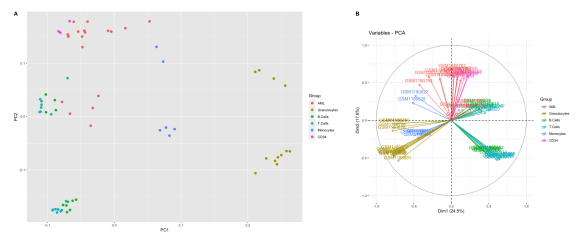
```
[12]: options(repr.plot.width=15, repr.plot.height=6)
    par(mfrow = c(1, 2))
    plot(pc)
    plot(pc$x[, 1:2])
```



As you can see, now other components are contributing as well.

 $<sup>^3 {\</sup>rm Subtracting}$  columns mean from their corresponding columns.

We can also plot the PCA results according to the variables (which are the samples in this case) instead of the individuals (which are the genes in this case) to demonstrate different sample groups and their correlation:



## About the figure B:

- Positively correlated variables are grouped together.
- Negatively correlated variables are positioned on opposite sides of the plot origin (opposed quadrants).

## 5.2.2 t-distributed stochastic neighbor embedding (tSNE)

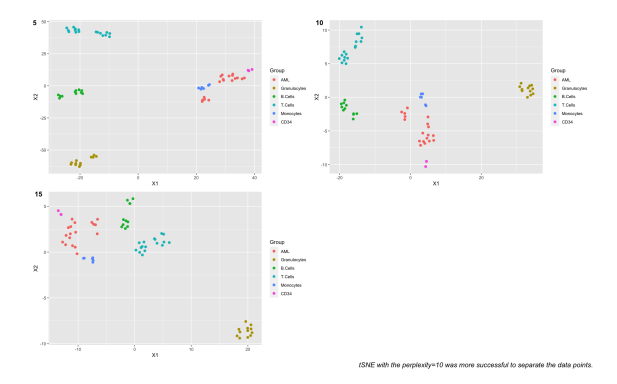
The next dimension reduction algorithm that we're going to try is the tSNE which as mentioned in figure 1, is a non-linear algorithm. Non-linearity means that it's capable of separating data which can't be separated by a straight line, unlike the linear algorithms (e.g. PCA).

We're going to use the Rtsne() method from the Rtsne package, to calculate the tSNE of our expression matrix. One parameter that we need to pass to the Rtsne() method is the perplexity. A perplexity is more or less a target number of neighbors for our central point. Basically, the higher the perplexity is the higher the variance value.

So we will calculate the tSNE for 3 perplexity values of 5, 10 and 15. After plotting the result, we can choose the best value based on how well it separated the group clusters:

```
[14]: tsne_results <- list(Rtsne(t(ex), perplexity=5, check_duplicates = FALSE),
                            Rtsne(t(ex), perplexity=10, check_duplicates = FALSE),
                            Rtsne(t(ex), perplexity=15, check_duplicates = FALSE))
      options(repr.plot.width=16, repr.plot.height=10)
      tsne.plots.list <- list()</pre>
      for(i in seq_along(tsne_results)) {
        tsne <- data.frame(tsne_results[[i]]$Y[, 1:2], Group=gs)</pre>
        tsne.plots.list[[i]] <- ggplot(tsne, aes(X1, X2, color = Group)) +
       →geom_point(size=2) + theme_gray()
      ggarr <- ggarrange(plotlist = tsne.plots.list,</pre>
                          ncol = 2,
                          nrow = 2,
                          labels = c(5, 10, 15))
      annotate_figure(ggarr,
                      bottom = text_grob("tSNE with the perplexity=10 was more_
       ⇒successful to separate the data points.",
                                         hjust = 1,
                                         x = 1,
                                         face = "italic",
                                         size = 14)
                      )
```

<sup>&</sup>lt;sup>4</sup>https://towardsdatascience.com/t-sne-clearly-explained-d84c537f53a



According to the plots, tSNE with the <code>perplexity=10</code> was more successful to separate the data points.

#### 5.2.3 Multidimensional scaling (MDS)

MDS returns an optimal solution to represent the data in a lower-dimensional space, where the number of dimensions k is pre-specified by the analyst. For example, choosing k = 2 optimizes the object locations for a two-dimensional scatter plot.

The input data for MDS is a dissimilarity matrix representing the distances between pairs of objects. So first we will compute the distance matrix using the dist() method:

```
[15]: ex_dist <- dist(t(ex.scale))</pre>
```

Now we can pass the ex\_dist to the cmdscale from the stats package to compute the mds result:

```
[16]: options(repr.plot.width=6, repr.plot.height=4)

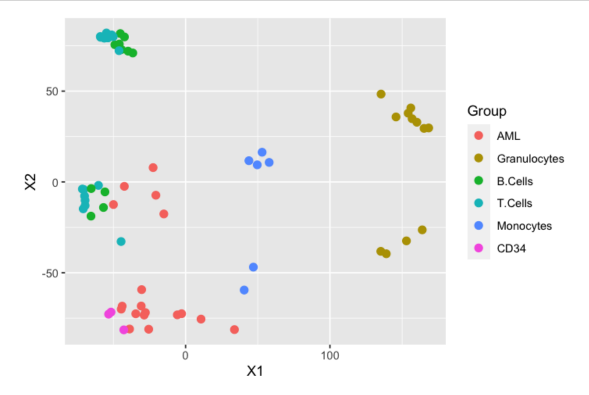
mds_result <- cmdscale(ex_dist, eig=TRUE, k=2)

mds_df <- data.frame(mds_result$points[,1:2], Group=gs)

mds.plot <- ggplot(mds_df, aes(X1,X2, color = Group)) + geom_point(size=2.5) +

theme_gray()

mds.plot</pre>
```



# 5.2.4 PCA vs. tSNE vs. MDS

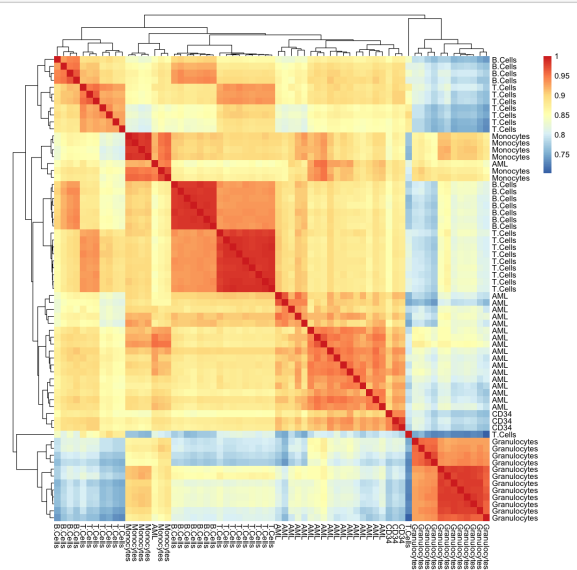
To compare the results of these 3 methods, we will plot their results one more time to make it easier for comparing:

According to plots, we believe that the tSNE was able to separate groups of samples more precisely.

# 5.3 Correlation between samples

Heat map shows correlations between different samples. For example each group has high corealation with itself that is determined by red color or granulocytes have a low correlation with B-cells and T-cells that is determined by blue color.

According to the heatmap, AML has a high correlation with CD34 and Monocytes which makes them proper candidates to check whether there are any significant expression differences between their genes and the AML ones.



# 6 References

- https://www.digitalocean.com/community/tutorials/normalize-data-in-r
- $\bullet \ https://www.v7labs.com/blog/data-preprocessing-guide$
- https://machinelearningmastery.com/dimensionality-reduction-for-machine-learning/
- $\bullet\ https://towards datascience.com/11-dimensionality-reduction-techniques-you-should-know-in-2021-dcb9500d388b$
- $\bullet \ http://www.sthda.com/english/articles/31-principal-component-methods-in-r-practical-guide/118-principal-component-analysis-in-r-prcomp-vs-princomp/ \\$
- $\bullet \ http://www.sthda.com/english/articles/31-principal-component-methods-in-r-practical-guide/112-pca-principal-component-analysis-essentials/ \\$
- https://ajitjohnson.com/tsne-for-biologist-tutorial/
- https://towardsdatascience.com/t-sne-clearly-explained-d84c537f53a