## **Bioinformatics Analysis in R**

# **Gene Expression Analysis**

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1 - Give you a overview on the use of R/bioconductor tools for gene expression analysis

2 - Show a real example with all steps necessary for gene expression analysis (based on arrays and RNA-seq)



### **Analysis of Gene Expression**



adapted from: Amit et al. 2016

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**Institute for** 

- 1- Which genes are up/down regulated after treatment?
- differential analysis / clustering genes
- 2 Which cells are more similar?
- clustering samples / PCA
- 3 How to interpret large lists of genes?
- gene ontology enrichment /gene set enrichment analysis (GSEA)

### **Bioinformatics - Gene Expression Analysis**





### **Bioinformatics - Gene Expression Analysis**







### **Affymetrix Arrays - Example**



### **Quantification/Pre-processing**



- Main steps 1 - Quantify gene expression values
- Quality Control<br>• remove had samples 2 - Quality Control • remove bad samples
- 2 Quality check (QC) 3 – Normalization • normalization3 - Correct for Experimental artifacts
	-



### **Why is QC / Normalization important?**

- Systematic errors (array wise)
	- labeling efficiency, scanning parameters, reverse transcriptase, batch effects
- Stochastic errors
	- cross-hybridization, image processing failure, error on probe sequence (manufacturer defect) (gene wise)
	- dust in array, hybridization problems (array wise)

#### **Example of Hybridization Problems**







#### **Normalization Principles**

1 - Most genes don't change expression -> small/same variance 2 - Arrays are hybridized with the same amount of DNA -> same mean



Arrays

**Hybridization intensity distributions across experiments**





#### **Scatter Plots - Comparing 2 arrays**





#### **Scatter Plots - Comparing 2 arrays**





#### **Normalization Results**

Application of BetweenArray normalization from limma package



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Shows systematic dependence between fluorescence intensities between arrays.

- $M = \log R/G$
- A =  $log (R * G)$  (= 1/2  $[log(R) + log(G)])$

For Affymetrix/single channel arrays, R is the intensity of the microarray experiment of interest and the G is the intensity of median values of all the arrays



#### **MA Plots**

"relative expression" (fold change)



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### **Quantification/Pre-processing - Resume**

- Normalization is important to confirm the quality and consistency of data
- Boxplots should also be performed after all steps to assure data standards
- Exclusion of "bad samples" has positive effect on downstream analysis
- **•In doubt, consult a bioinformatician!**



### **Bioinformatics - Gene Expression Analysis**







### **Differential Expression Analysis**

- Identify genes related to a particular condition
	- example van de Laar, et al. 2016, Immunity, 2016.
- We will consider:
	- You Sac Macrophages (YS-Mac)
	- Fetal Liver Monocytes (FL-MO)
	- Bone Marrow Monocytes (BM-MO)
		- 4 replicates per condition



Source: van de Laar, et al. 2016, Immunity, 2016.



### **Differential Expression - Example**



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### **Differential Expression - Example**

Fold change analysis - change  $>$   $|log2(2)|$ 



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- Low expression genes are treated equally as high expression genes
- We lose information about the variance from genes
- No statistical significance
- Is the only alternative when no replicate samples are available (not recommended!)



### **Basic Concepts**

#### **Mean vs. variability**



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#### **T-test**

We can use the t-statistic as an indication of differential expression

$$
t = \frac{\bar{X} - \bar{Y}}{\mathrm{SE}} \cdot \frac{\mathrm{difference\ between\ means}}{\mathrm{variance}}
$$
  

$$
\mathrm{SE} = \sqrt{\frac{s_X^2}{n_X} + \frac{x_Y^2}{n_Y}} \quad \text{and} \quad s_X^2 = \frac{1}{n_X - 1} \sum_{i=1}^{n_X} (X_i - \bar{X})^2 \, .
$$

where X and Y are the mean (log) expression values of a gene in each group sample and  $n_x$  and  $n_y$  are the number of samples on these groups



### **Student T-test**

Test the hypothesis 
$$
H_0 : X - Y = 0
$$
  
 $H_1 : X - Y \neq 0$ 

We can use the t-student distribution to estimate for which t-statistic values the null hypothesis is rejected.

 $P$ -value = Pr(t as extreme or more $H_0$ ),







#### Change: **HIGH Variance: SMALL**

#### **Change: SMALL Variance: HIGH**



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#### **Results - FL-MO vs. BM-MO**

#### **Volcano Plot - combine p-value and fold change**



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- With a p-value of 0.01, we expect to make one mistake every 100 tests
- We have 12.626 genes, therefore 126 mistaken from 1046 DE genes.
- To solve this, a multiple test correction method is necessary (i.e. Benjamini-Hochberg)
	- It is based on the false discovery rate, i.e. the proportion of false DE genes in your list of DE genes



- Higher level analysis are eased by filtering of non-specific genes maximum
	- genes that show no expression changes between arrays
	- i.e. filter genes with low IQR (interquantile range)



- Affymetrix chips has spike-in control probes
	- Should be removed after normalization



### **Differential Analysis - Conclusions**

- Fold-change (alone) -> should be avoided
- For patient samples
	- high number of replicates are necessary (>30)
	- otherwise low DE genes replicability
- For model (mouse) experiments
	- at least 3 samples (and moderated t-test)
	- we can not tell the variance without measuring it!
- All correct for multiple testing! Also, non-specific filtering can help if low number of DE genes is found.



### **Practical Example**

- This data is deposited in the public repository GEO under accession [GSE76999](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE76999)
- This can be found at the materials and methods of papers.
- GEO public database with raw, pre-processed data and experimental details of expression (and other omics) experiments. Source: van de Laar, et al. 2016, Immunity, 2016.







### **GEO - van de Laar, et al. 2016**

**Series GSE76999 Query DataSets for GSE76999** Public on Mar 01, 2016 **Status** Capacity of yolk sac macrophages, fetal liver and adult monocytes to colonize **Title** an empty niche and develop into functional tissue resident macrophages Organism Mus musculus **Experiment type** Expression profiling by array Tissue-resident macrophages can derive from yolk sac macrophages, fetal liver Summary monocytes or adult bone marrow monocytes. Whether these precursors can give rise to transcriptionally identical alveolar macrophages is unknown. Here, we transferred traceable yolk sac macrophages, fetal liver monocytes, adult bone marrow monocytes or adult alveolar macrophages as a control, into the empty alveolar macrophage niche of neonatal Csf2rb-/- mice. All precursors efficiently colonized the alveolar niche and generated alveolar macrophages that were transcriptionally almost identical, with only 22 genes that could be linked to their origin. Underlining the physiological relevance of our findings, all transfer-derived alveolar macrophages self-maintained within the lungs for up to 1 year and durably prevented alveolar proteinosis. Thus, precursor origin does not affect the development of functional self-maintaining tissue-resident macrophages.

Overall design CD45.1+CD45.2+ yolk sac macrophages, fetal liver monocytes, adult bone marrow monocytes or adult alveolar macrophages from the bronchoalveolar lavage were sorted from wild type CD45.1+CD45.2+ mice of indicated ages. From part of these samples RNA was isolated. The other part was transferred intranasally into the lungs of neonate Csf2rb-/- mice. 6 weeks post-transfer, transfer-derived CD45.1+CD45.2+ alveolar macrophages were sorted from the bronchoalveolar lavage. Wild type CD45.1+CD45.2 alveolar macrophages from the bronchoalveolar lavage of 6 week old mice were sorted as control. 36 samples (arrays) in total. RNA was isolated, amplified with Nugene pico kit, converted to cDNA and then hybridised on Affymetrix GeneChip Mouse Gene 1.0 ST Arrays.

Contributor(s) van de Laar L, Saelens W, De Prijck S, Martens L, Scott CL, Van Isterdael G, Hoffmann E, Beyaert R, Saeys Y, Lambrecht BN, Guilliams M van de Laar L, Saelens W, De Prijck S, Martens L et al. Yolk Sac Macrophages, Citation(s) Fetal Liver, and Adult Monocytes Can Colonize an Empty Niche and Develop into Functional Tissue-Resident Macrophages. Immunity 2016 Apr 19;44(4):755-68. PMID: 26992565

#### GEO ID

Information about the study

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### **GEO - van de Laar, et al. 2016**



### **GEO - van de Laar, et al. 2016**



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# **Hands on!**

Handout Steps 1,2 and 3



### **Principal Component Analysis**

- **• method for dimension reduction** 
	- **• find combination of genes explaining cells with distinct expression**
- **• finding directions with highest variance**



Recommended reading: Ringner M., *Nature Biotechnology* 26, 303 - 304 (2008)

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#### **Gene Expression - PCA Example 1**

#### **Can be interpreted as a computational FACs sorting (without knowing the markers)**



#### **First 2 PCs on the analysis of 5000 samples from Array Express/EBI**



#### **Gene Expression - PCA Example 2**

#### **PCA Analysis of van de Leer, 2016 data**



#### **First 2 PCs van de Leer, 2016 data**





#### **PCA Analysis - Conclusions**

- **• PCA allows an "blind" cell sorting**
	- **• only works if variant directions split the groups**
	- **• is complementary to clustering**
- **• Weights allow interpretation of relevant variables**
- **• Can also be used for quality check** 
	- **• samples not fitting to groups**
- **• Alternatives to PCA:**
	- **• tSNE very commonly used in single cell RNA-seq**



#### **Clustering / Heatmaps**





#### **Clustering / Heatmaps**



**clustering methods: k-means, hierarchical clustering, …**



#### **Distance**

• For a expression matrix *X* (genes vs. arrays), measure the distance between expression values of two genes (*x<sub>i</sub>* e *x<sub>j</sub>*)

• Euclidean distance (sensitive to scale)

$$
d(x_i, x_j) = \sqrt{\sum_{l=1}^{L} (x_{il} - x_{jl})^2}
$$

• Pearson correlation (not sensitive to scale / similarity measure)

$$
d(x_i, x_j) = \frac{\sum_{i=1}^{L} (x_{il} - \overline{x}_i)(x_{jl} - \overline{x}_j)}{\sigma_i^2 \sigma_j^2}
$$



#### **Distance**

• Which distance for gene expression?

• - example of two genes for 15 cancer patients



• Euclidean - not similar • Correlation - similar

• z-score normalised expression



• Euclidean - similar • Correlation - similar



## **Hierarchical Clustering**



• Botton up method

Starting with a distance (similarity) matrix and each object as a group

Repeat:

- Joint two most similar groups
- Until the dendrogram has only one group





Gene 1

Single-Linkage

- Join two groups where two examples are close
- Find groups with linear shapes



#### Distance Matrix



5  $\overline{4}$  $\overline{3}$  $\mathcal{L}$ 





#### Distance Matrix







#### Distance Matrix



$$
d_{(1,2),3} = \min\{d_{1,3}, d_{2,3}\} = \min\{6,3\} = 3
$$
  

$$
d_{(1,2),4} = \min\{d_{1,4}, d_{2,4}\} = \min\{10,9\} = 9
$$
  

$$
d_{(1,2),5} = \min\{d_{1,5}, d_{2,5}\} = \min\{9,8\} = 8
$$





#### **Hierarchical Clustering**



$$
d_{(1,2,3),4} = \min\{d_{(1,2),4}, d_{3,4}\} = \min\{9,7\} = 7
$$

$$
d_{(1,2,3),5} = \min\{d_{(1,2),5}, d_{3,5}\} = \min\{8,5\} = 5
$$



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#### **Hierarchical Clustering**



$$
d_{(1,2,3),(4,5)} = \min\{d_{(1,2,3),4}, d_{(1,2,3),5}\} = 5
$$







Single-Linkage

- Groups with closest genes
- linear shapes

Complete-Linkage

- Closest groups with more far genes
- Compact clusters

Average Linkage

- Groups with closest centroids (middle)
- Gene 1 **Constant Construction Construction** Construction Construction Construction Construction Construction Const



# Which linkage?

# Which distance?



## **Hierarchical Clustering - Complete Linkage**





## **Hierarchical Clustering - Average Linkage**





#### **Hierarchical Clustering - Single Linkage**





#### **Hierarchic Clustering - Final Results**

**Color Key**





## **Clustering - Resume**

- Clustering allow detection of unknown groups in the data
- Classical methods (hierarchical or k-mean) work well in general
- How to choose distance and linkage?
	- Pearson or Euclidean (followed by z-transform)
	- Heatmaps usually only like nice with ztransform
- How to find number of groups?
	- No simple solution!



# **Functional Analysis**

Clustering/Differential Expression (DE) returns lists of hundreds of genes How to functionally characterize these?

**Solution 1** - Look at each gene individually

**Solution 2** - Relate these genes to annotations from databases

- Gene Ontology, pathways, gene sets, disease ontology, …



## **Databases**

Manually or automatic curated annotation of genes



# **Gene Ontology**

Controlled vocabulary to describe gene and gene product attributes in any organism

Formed by three ontologies

- 1. Biological Process (BP)
- 2. Molecular Function (MF)
- 3. Celular Component (CC)

Annotation (Organism depend) - genes are associated to terms manually (literature) or automatically (sequence homology)



# **Gene Ontology**





# **Gene Ontology**



#### inheritance property



# **GO Enrichment Analysis**



#### How probable is that 3 up regulated genes are annotated to the GO term?



# **GO Enrichment Analysis**





For a given gene list:

- 1. evaluate the the overlap of the list vs. all gene sets i.e. GO terms, pathways, …
- 2. Estimate p-value (corrected by multiple testing)
- 3. Rank gene sets by lowest p-value



We interface for enrichment analysis with: Gene Ontology, KEGG Pathway and TF binding

http://biit.cs.ut.ee/gprofiler/index.cgi

Check the results for my favorite genes:

Irf8 Id2 Spi1 Klf4 Runx2 Egr1



Perform a functional evaluation of ranking of genes

- i.e all genes ranked by fold change cond. A vs. B

Advantages over "Normal" enrichment analysis:

- do not require previous DE analysis
- works when effects of the experiment are low

GSEA Gene Sets

- GO Terms, KEGG Pathways
- experimentally derived Gene Sets
	- DE genes from microarray studies from GEO
	- Can be obtained at mysigdb

[\(software.broadinstitute.org/gsea/msigdb/\)](http://software.broadinstitute.org/gsea/msigdb/)

http://www.broadinstitute.org/gsea









For a given gene ranking:

- 1. evaluate ES score for all gene sets
- 2. estimate p-value(corrected)
- 3. rank gene sets by lowest pvalue



http://www.broadinstitute.org/gsea


# **Hands on!**

Handout Step 4 to 7



### **Integrative Analysis - ImmGen**

• ImmGen - expression data of immune cells under standardized conditions



- How do cells from **van de Leer, 2016** compares to monocyte/macrophages from ImmGenn?
- we obtained/pre-processed ImmGen data (v1) from GEO (GSE15907)



#### **Integrative Analysis - Problem**

• Batch Effects - Arrays from distinct lab tends to cluster together



See: Leek JT,.... (2016). sva: Surrogate Variable Analysis. R package version 3.22.0.

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# **Integrative Analysis - Problem**

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#### **Integrative Analysis - PCA After Combat**

- Solution Batch effect removal with COMBAT
	- annotation of your data: tissue of origin, cell type, experimental batches

# Hands on! Handout Step 7

See: Leek JT,…. (2016). sva: Surrogate Variable Analysis. R package version 3.22.0.

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## **Bioinformatics - Gene Expression Analysis**





## **Afternoon Exercise**

- Analyse gene expression data (steps 1-7 of handout) of the following paper:
	- Spence JR, Mayhew CN, Rankin SA, Kuhar MF et al. Directed differentiation of human pluripotent stem cells into intestinal tissue in vitro. Nature 2011 Feb 3;470(7332):105-9.

Try to get answers to the following questions with your analysis:

- Are the stem cells and induced pluripotent cells the same?
- If not, what are the reasons?
	- Are there changes in gene expression between these cells?
	- Does GO analysis indicate any functional differences between these cells?





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