# A very short, sketchy, introduction to Bioconductor

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**BIOF 339** 

#### Bioconductor

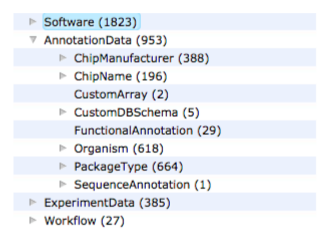
Bioconductor provides tools for the analysis and comprehension of high-throughput genomic and biological data, using R.

- 1823 packages
- Covers the bioinformatic pipeline
- Analysis [GenomicRanges, Biostrings, GenomicAlignments, SummarizedExperiment]
- Annotation (species/platform specific, system) [biomaRt, org.Hs.eg.db, G0.db, KEGG.db]
- Experiments [TENxPBMCData, airway, ALL]
- Workflows [rnaseqGene, TCGAWorkflow]

#### Bioconductor

#### ▼ Software (1823) AssayDomain (732) ▶ BiologicalQuestion (756) ▶ Infrastructure (404) ResearchField (810) StatisticalMethod (652) ► Technology (1160) ▶ WorkflowStep (986) ▶ AnnotationData (953) ExperimentData (385) ▶ Workflow (27) ▶ Software (1823) AnnotationData (953) ExperimentData (385) AssayDomainData (72) ▶ DiseaseModel (88) ▶ OrganismData (132) ▶ PackageTypeData (27) ▶ RepositoryData (91) ReproducibleResearch (20) ▶ SpecimenSource (101) ▶ TechnologyData (254) ▶ Workflow (27)

#### Bioconductor v. 3.10 packages



# Installing Bioconductor packages

Bioconductor is a separate repository and system which uses R. So the process is a bit different than install.packages. The following works for R version 3.5 and higher.

```
install.packages("BiocManager")
BiocManager::install(c('Biobase','limma','hgu95av2.db','Biostrings'))
```

There are several packages that are often installed for each Bioconductor package, and some have functions that have the same name as one's you've used. So

• Use package::function format for calling functions from non-Bioconductor packages

#### Bioconductor basics

```
library(Biostrings)
dna <- DNAStringSet(c("AACAT", "GGCGCCT"))</pre>
reverseComplement(dna)
DNAStringSet object of length 2:
    width seq
[1]
       5 ATGTT
        7 AGGCGCC
library(Biostrings)
data("phiX174Phage")
phiX174Phage
DNAStringSet object of length 6:
    width seq
                                                            names
[1] 5386 GAGTTTTATCGCTTCCATGACGC...ATGATTGGCGTATCCAACCTGCA Genbank
    5386 GAGTTTTATCGCTTCCATGACGC...ATGATTGGCGTATCCAACCTGCA RF70s
    5386 GAGTTTTATCGCTTCCATGACGC...ATGATTGGCGTATCCAACCTGCA SS78
    5386 GAGTTTTATCGCTTCCATGACGC...ATGATTGGCGTATCCAACCTGCA Bull
   5386 GAGTTTTATCGCTTCCATGACGC...ATGATTGGCGTATCCAACCTGCA G97
   5386 GAGTTTTATCGCTTCCATGACGC...ATGATTGGCGTATCCAACCTGCA NEB03
```

## **Bioconductor basics**

```
letterFrequency(phiX174Phage, 'GC', as.prob=TRUE)

G|C

[1,] 0.4476420

[2,] 0.4472707

[3,] 0.4472707

[4,] 0.4470850

[5,] 0.4472707

[6,] 0.4470850
```

#### Bioconductor data structures

- So far we've seen the data.frame or tibble be the unit of data storage
- In Bioconductor, data are stored in containers which can contain many elements of data for an experiment
  - Actual quantitative results of experiments
  - Phenotype data
  - Genotype meta-data
  - Results of analysis
- In Bioconductor workflows, the same container is updated with new elements, which can then be accessed using accessor functions

```
library(Biobase)
data('sample.ExpressionSet')
str(sample.ExpressionSet)
```

```
Formal class 'ExpressionSet' [package "Biobase"] with 7 slots
  ..@ experimentData :Formal class 'MIAME' [package "Biobase"] with 13 slots
 .. .. a name
                             : chr "Pierre Fermat"
 ..... chr "Francis Gatton Eas
..... chr "pfermat@lab.not.exist"
..... chr "Smoking-Cancer Experim
 .... title : chr "Smoking-Cancer Experiment" : chr "An example object of expression set (ExpressionSet) class"
  .. .. ..a url
                             : chr "www.lab.not.exist"
 ..... pubMedIds
                             : chr
 .. .. ..∂ samples
                             : list()
 ..... .. .. .. hybridizations : list()
 .. .. .. a normControls
                             : list()
 .. .. .. a preprocessing
                             : list()
 .. .. ..მ other
                             :List of 1
 ..... s notes: chr "An example object of expression set (exprSet) class"
 .. .. .. വ .__classVersion__:Formal class 'Versions' [package "Biobase"] with 1 slot
 .. .. .. .. .. .. .. .. Data:List of 2
 .. .. .. .. .. .. .. : int [1:3] 1 0 0
 .. .. .. .. ... ... : int [1:3] 1 1 0
 ..@ assayData :<environment: 0x7fcd93cd18d8>
 ..@ phenoData :Formal class 'AnnotatedDataFrame' [package "Biobase"] with 4 slots
 .. .. ..എ varMetadata
                             :'data.frame':
                                              3 obs. of 1 variable:
 ......$ labelDescription: chr [1:3] "Female/Male" "Case/Control" "Testing Score"
                             :'data.frame': 26 obs. of 3 variables:
 .. .. ..@ data
```

varMetadata: labelDescription

These objects are based on a different R structure. Instead of extracting elements using \$, this structure uses **slots** which are accessed using a

We almost never use a. Instead we use accessor functions to extract data

```
pData(sample.ExpressionSet) # Phenotype data
```

```
sex
            type score
A Female Control
                0.75
   Male
           Case
                 0.40
   Male Control
                0.73
   Male
           Case
                 0.42
E Female
           Case 0.93
   Male Control
                0.22
   Male
           Case
                 0.96
   Male
           Case
                 0.79
 Female
           Case
                 0.37
   Male Control
                 0.63
   Male
                0.26
           Case
 Female Control
                0.36
   Male
           Case
                 0.41
   Male
           Case
                 0.80
O Female
           Case 0.10
P Female Control
                0.41
Q Female
           Case 0.16
   Male Control
                0.72
   Male
           Case
                 0.17
 Female
           Case
                 0.74
   Male Control 0.35
 Female Control
                 0.77
   Male Control
                0.27
```

We almost never use a. Instead we use accessor functions to extract data

```
head(exprs(sample.ExpressionSet)) # Expression data
```

```
AFFX-MurIL2 at 192.7420
                        85.75330 176.7570 135.5750 64.49390 76.3569 160.5050
AFFX-MurIL10 at 97.1370 126.19600
                                  77.9216
                                           93.3713 24.39860 85.5088
AFFX-MurIL4 at
                45.8192
                         8.83135
                                  33.0632
                                           28.7072
                                                                     30.9694
AFFX-MurFAS at 22.5445
                        3.60093
                                  14.6883
                                           12.3397 36.86630 11.2568
                                                                    23.0034
AFFX-BioB-5 at 96.7875
                                  46.1271 70.9319 56.17440 42.6756
                        30.43800
AFFX-BioB-M at
                89.0730
                        25.84610
                                  57.2033
                                           69.9766 49.58220 26.1262
                                                                    75.0083
AFFX-MurIL2 at 65.9631 56.9039 135.60800 63.44320 78.2126 83.0943 89.3372
AFFX-MurIL10 at 81.6932 97.8015 90.48380 70.57330 94.5418 75.3455 68.5827
AFFX-MurIL4 at 14.7923 14.2399
                              34.48740 20.35210 14.1554 20.6251 15.9231
AFFX-MurFAS at 16.2134 12.0375
                                         8.51782 27.2852 10.1616 20.2488
AFFX-BioB-5 at 30.7927 19.7183
                                46.35200 39.13260 41.7698 80.2197 36.4903
AFFX-BioB-M at 42.3352 41.1207
                                91.53070 39.91360 49.8397 63.4794 24.7007
AFFX-MurIL2 at 91.0615 95.9377 179.8450 152.4670 180.83400 85.4146 157.98900
AFFX-MurIL10 at 87.4050 84.4581 87.6806 108.0320 134.26300 91.4031
                                                                   -8.68811
AFFX-MurIL4 at 20.1579 27.8139
                              32.7911
                                       33.5292 19.81720 20.4190
                                                                   26.87200
AFFX-MurFAS at 15.7849 14.3276
                               15.9488
                                       14.6753
                                                -7.91911 12.8875
                                                                  11.91860
AFFX-BioB-5 at 36.4021 35.3054 58.6239 114.0620 93.44020 22.5168
                                                                   48.64620
AFFX-BioB-M at 47.4641 47.3578
                                58.1331 104.1220 115.83100 58.1224
                                                                   73.42210
                      V
                              W
                                        Χ
                                                         Ζ
AFFX-MurIL2 at 146.8000 93.8829 103.85500 64.4340 175.61500
AFFX-MurIL10 at 85.0212 79.2998 71.65520 64.2369
```

# SummarizedExperiment

This is a more common structure related to modern experiments with different technologies

```
# BiocManager::install('SummarizedExperiment')
library(SummarizedExperiment)
data(airway, package="airway")
se <- airway
se</pre>
```

```
class: RangedSummarizedExperiment
dim: 64102 8
metadata(1): ''
assays(1): counts
rownames(64102): ENSG00000000000 ENSG0000000000 ... LRG_98 LRG_99
rowData names(0):
colnames(8): SRR1039508 SRR1039509 ... SRR1039520 SRR1039521
colData names(9): SampleName cell ... Sample BioSample
```

Count data from the scRNA-seq experiment

assay(se)

	SRR1039508	SRR1039509	SRR1039512	SRR1039513	SRR1039516
ENSG00000000003	679	448	873	408	1138
ENSG00000000005	0	0	0	0	0
ENSG00000000419	467	515	621	365	587
ENSG00000000457	260	211	263	164	245
ENSG00000000460	60	55	40	35	78
ENSG00000000938	0	0	2	0	1
ENSG00000000971	3251	3679	6177	4252	6721
ENSG00000001036	1433	1062	1733	881	1424
ENSG00000001084	519	380	595	493	820
ENSG00000001167	394	236	464	175	658
ENSG00000001460	172	168	264	118	241
ENSG00000001461	2112	1867	5137	2657	2735
ENSG00000001497	524	488	638	357	676
ENSG00000001561	71	51	211	156	23
ENSG00000001617	555	394	905	415	727
ENSG00000001626	10	2	9	2	10
ENSG00000001629	1660	1251	2259	1079	2462
ENSG00000001630	59	54	66	23	84
ENSG00000001631	729	692	943	475	1034
ENSG00000002016	201	161	256	99	268
ENSG00000002079	3	0	3	1	4
ENSG00000002330	206	174	184	111	194
ENSG00000002549	1459	1294	1317	998	1451

Genomic ranges for each transcript

```
rowRanges(se)
```

```
GRangesList object of length 64102:
$ENSG00000000003
GRanges object with 17 ranges and 2 metadata columns:
       segnames
                            ranges strand
                                               exon id
                                                             exon name
          <Rle>
                        <IRanges> <Rle>
                                            <integer>
                                                           <character>
   [1]
[2]
[3]
[4]
[5]
              X 99883667-99884983
                                               667145 ENSE00001459322
                                                667146 ENSE00000868868
              X 99885756-99885863
              X 99887482-99887565
                                                667147 ENSE00000401072
              X 99887538-99887565
                                               667148 ENSE00001849132
              X 99888402-99888536
                                                667149 ENSE00003554016
  [13]
              X 99890555-99890743
                                                667156 ENSE00003512331
  [14]
[15]
[16]
              X 99891188-99891686
                                                667158 ENSE00001886883
              X 99891605-99891803
                                                667159 ENSE00001855382
              X 99891790-99892101
                                                667160 ENSE00001863395
  [17]
              X 99894942-99894988
                                                667161 ENSE00001828996
 seqinfo: 722 sequences (1 circular) from an unspecified genome
<64101 more elements>
```

Phenotype data

```
colData(se)
```

```
DataFrame with 8 rows and 9 columns
           SampleName
                          cell
                                    dex
                                           albut
                                                         Run avgLength
             <factor> <factor> <factor> <factor>
                                                    <factor> <integer>
SRR1039508 GSM1275862
                       N61311
                                           untrt SRR1039508
                                                                   126
                                  untrt
                                                                   126
SRR1039509 GSM1275863
                       N61311
                                  trt
                                           untrt SRR1039509
SRR1039512 GSM1275866
                                                                   126
                       N052611
                                           untrt SRR1039512
                                  untrt
SRR1039513 GSM1275867
                       N052611
                                  trt
                                           untrt SRR1039513
                                                                    87
SRR1039516 GSM1275870
                       N080611
                                           untrt SRR1039516
                                                                   120
                                  untrt
SRR1039517 GSM1275871
                       N080611
                                           untrt SRR1039517
                                                                   126
                                  trt
SRR1039520 GSM1275874
                                           untrt SRR1039520
                       N061011
                                                                   101
                                  untrt
                                           untrt SRR1039521
SRR1039521 GSM1275875
                       N061011
                                  trt
                                                                    98
                                   BioSample
           Experiment
                         Sample
             <factor> <factor>
                                    <factor>
SRR1039508 SRX384345 SRS508568 SAMN02422669
SRR1039509 SRX384346 SRS508567 SAMN02422675
SRR1039512 SRX384349 SRS508571 SAMN02422678
SRR1039513 SRX384350 SRS508572 SAMN02422670
SRR1039516 SRX384353 SRS508575 SAMN02422682
SRR1039517 SRX384354 SRS508576 SAMN02422673
SRR1039520 SRX384357 SRS508579 SAMN02422683
SRR1039521
           SRX384358 SRS508580 SAMN02422677
```

Experimental meta-data

metadata(se)

```
[[1]]
Experiment data
    Experimenter name: Himes BE
    Laboratory: NA
    Contact information:
    Title: RNA-Seq transcriptome profiling identifies CRISPLD2 as a glucocorticoid responsive gene that modulates
    URL: http://www.ncbi.nlm.nih.gov/pubmed/24926665
    PMIDs: 24926665

Abstract: A 226 word abstract is available. Use 'abstract' method.
```

Data subsetting

```
se[1:5, 1:3]
```

```
class: RangedSummarizedExperiment
dim: 5 3
metadata(1): ''
assays(1): counts
rownames(5): ENSG00000000000 ENSG0000000000 ENSG000
    ENSG0000000457 ENSG00000000460
rowData names(0):
colnames(3): SRR1039508 SRR1039509 SRR1039512
colData names(9): SampleName cell ... Sample BioSamp
```

```
se[,se$cell=='N61311']
```

```
class: RangedSummarizedExperiment
dim: 64102 2
metadata(1): ''
assays(1): counts
rownames(64102): ENSG00000000003 ENSG00000000005 ...
rowData names(0):
colnames(2): SRR1039508 SRR1039509
colData names(9): SampleName cell ... Sample BioSamp
```

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# Annotation

#### biomaRt

The biomaRt package allows access to many public annotation databases

# Identifying attributes

```
searchAttributes(mart=ensemblHuman, pattern='affy')
```

```
description
                     name
                                                               page
107
             affv hc g110
                                   AFFY HC G110 probe feature page
            affy_hg_focus
108
                                  AFFY HG Focus probe feature page
109
            affy hg u133a
                                  AFFY HG U133A probe feature page
110
          affy hg u133a 2
                                AFFY HG U133A 2 probe feature page
111
            affy hg u133b
                                  AFFY HG U133B probe feature page
112
      affy hg u133 plus 2
                            AFFY HG U133 Plus 2 probe feature page
113
             affy hg u95a
                                   AFFY HG U95A probe feature page
114
           affy_hg_u95av2
                                 AFFY HG U95Av2 probe feature page
115
             affy hg u95b
                                   AFFY HG U95B probe feature page
116
             affy hg u95c
                                   AFFY HG U95C probe feature page
117
             affy hg u95d
                                   AFFY HG U95D probe feature page
118
             affy hg u95e
                                   AFFY HG U95E probe feature page
119
             affy hta 2 0
                                   AFFY HTA 2 0 probe feature page
120
      affy huex 1 0 st v2
                            AFFY HuEx 1 0 st v2 probe feature page
121
            affy hugenefl
                                  AFFY HuGeneFL probe feature page
   affy hugene 1 0 st v1 AFFY HuGene 1 0 st v1 probe feature page
123 affy_hugene_2_0_st_v1 AFFY HuGene 2 0 st v1 probe feature_page
           affy primeview
                                 AFFY PrimeView probe feature page
124
125
            affy u133 x3p
                                  AFFY U133 X3P probe feature page
```

# Identifying attributes

```
name description page
64 hgnc_id HGNC ID feature_page
65 hgnc_symbol HGNC symbol feature_page
96 hgnc_trans_name Transcript name ID feature_page
```

# **Annotating probsets**

We first grab some probesets from the sample. ExpressionSet Affy experiment

```
affyIDs <- rownames(featureData(sample.ExpressionSet))</pre>
```

Now let's find annotation

```
getBM(attributes = c('affy_hg_u95av2','entrezgene_id'),
    filters = 'affy_hg_u95av2',
    values = affyIDs[200:203],
    mart = ensemblHuman)
```

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# A RNA-seq pipeline

#### The workflow

- Exploratory data analysis
- Differential expression analysis with DESeq2
- Visualization
- We will start after reads have been aligned to a reference genome and reads overlapping known genes have been counted

#### The experiment

- In the experiment, four primary human airway smooth muscle cell lines were treated with 1 micromolar dexamethasone for 18 hours.
- For each of the four cell lines, we have a treated and an untreated sample.

#### Get data

```
# BiocManager::install('airway')
library(airway)
data(airway)
se <- airway
head(assay(airway))</pre>
```

```
SRR1039508 SRR1039509 SRR1039512 SRR1039513 SRR1039516
ENSG00000000003
                        679
                                   448
                                               873
                                                          408
                                                                     1138
ENSG00000000005
                          0
                                     0
                                                 0
                                                            0
                                                                        0
                                                                      587
ENSG00000000419
                        467
                                   515
                                               621
                                                           365
ENSG00000000457
                        260
                                   211
                                               263
                                                           164
                                                                      245
ENSG00000000460
                         60
                                    55
                                                40
                                                           35
                                                                       78
ENSG00000000938
                          0
                                                            0
                SRR1039517 SRR1039520 SRR1039521
ENSG00000000003
                       1047
                                   770
                                               572
ENSG00000000005
                          0
                                     0
                                                 0
ENSG00000000419
                        799
                                   417
                                               508
ENSG00000000457
                        331
                                   233
                                               229
ENSG00000000460
                         63
                                    76
                                                60
ENSG00000000938
                          0
                                     0
                                                 0
```

# Create a DESeqDataSet

```
# BiocManager::install('DESeq2')
library("DESeq2")
dds <- DESeqDataSet(se, design = ~ cell + dex)
dds</pre>
```

```
class: DESeqDataSet
dim: 64102 8
metadata(2): '' version
assays(1): counts
rownames(64102): ENSG00000000003 ENSG00000000005 ... LRG_98 LRG_99
rowData names(0):
colnames(8): SRR1039508 SRR1039509 ... SRR1039520 SRR1039521
colData names(9): SampleName cell ... Sample BioSample
```

# Run differential expression pipeline

dds <- DESeq(dds) estimating size factors estimating dispersions gene-wise dispersion estimates mean-dispersion relationship final dispersion estimates fitting model and testing

# Run differential expression pipeline

```
(res <- results(dds))</pre>
```

```
log2 fold change (MLE): dex untrt vs trt
Wald test p-value: dex untrt vs trt
DataFrame with 64102 rows and 6 columns
                 baseMean log2FoldChange
                                                                  pvalue
                                          lfcSE
                                                        stat
                <numeric>
                               <numeric> <numeric> <numeric>
                                                                <numeric>
ENSG00000000003 708.6022
                               0.3812540 0.100654 3.787752 0.000152016
ENSG00000000005
                   0.0000
                                                NA
                                                          NA
ENSG00000000419 520.2979
                              -0.2068126 0.112219 -1.842943 0.065337292
ENSG00000000457 237.1630
                              -0.0379204 0.143445 -0.264356 0.791505742
ENSG00000000460 57.9326
                               0.0881682 0.287142 0.307054 0.758801924
. . .
                                     . . .
LRG 94
                                      NA
                                                NA
                                                          NA
                                                                      NA
LRG 96
                                      NA
                                                NA
                                                          NA
                                                                      NA
LRG_97
                                      NA
                                                NA
                                                          NA
                                                                      NA
LRG 98
                                      NA
                                                NA
                                                          NA
                                                                      NA
LRG 99
                                      NA
                                                NA
                                                          NA
                      padj
                 <numeric>
ENSG00000000000 0.00128363
ENSG00000000005
ENSG00000000419 0.19654609
ENSG00000000457 0.91145948
ENSG00000000460 0.89503278
. . .
LRG 94
LRG_96
```

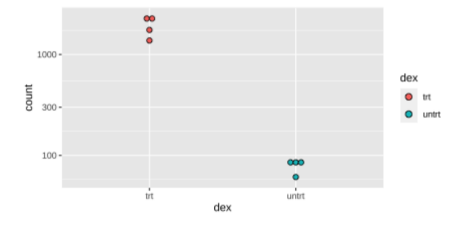
## Summarizing results

```
library(tidyverse)
as.data.frame(res) %>%
  rownames_to_column(var = 'ID') %>%
  filter(padj < 0.1) %>%
  arrange(desc(abs(log2FoldChange))) %>% head()
```

```
baseMean log2FoldChange
                                               lfcSE
                                                           stat
                                                                       pvalue
                                  -9.505975 1.0545032 -9.014647 1.975049e-19
1 ENSG00000179593
                  67.243048
2 ENSG00000109906 385.071029
                                  -7.352626 0.5363887 -13.707645 9.137621e-43
3 ENSG00000250978 56.318194
                                  -6.327383 0.6777973 -9.335214 1.007876e-20
4 ENSG00000132518
                  5.654654
                                  -5.885114 1.3240439 -4.444803 8.797262e-06
5 ENSG00000128285
                  6.624741
                                 5.325904 1.2578147 4.234251 2.293144e-05
6 ENSG00000127954 286.384119
                                  -5.207158 0.4930818 -10.560435 4.545484e-26
          padi
1 1.253739e-17
2 2.256617e-40
3 7.210311e-19
4 1.000612e-04
5 2.380012e-04
6 5.058395e-24
```

#### A visualization

```
topGene <- rownames(res)[which.min(res$padj)]
dat <- plotCounts(dds, gene=topGene, intgroup=c("dex"), returnData=TRUE)
ggplot(dat, aes(x = dex, y = count, fill=dex))+
   geom_dotplot(binaxis='y', stackdir='center')+
   scale_y_log10()</pre>
```



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# A Seurat pipeline

#### Grab the data and convert for Seurat

```
An object of class Seurat

13714 features across 2700 samples within 1 assay
Active assay: RNA (13714 features, 0 variable features)
```

#### A bit of QC

```
# The [[ operator can add columns to object metadata. This is a great place to stash QC stats rownames(pbmc@assays$RNA@counts) <- r2 rownames(pbmc[['RNA']]@meta.features) <- r2 rownames(pbmc@assays$RNA@data) <- r2 # Change to gene names from Ensembl pbmc[["percent.mt"]] <- PercentageFeatureSet(pbmc, pattern = "^MT-") # percentage in mitochondria genome head(pbmc@meta.data)
```

```
orig.ident nCount_RNA nFeature_RNA percent.mt
Cell1
          pbmc3k
                      2419
                                    779 3.0177759
Cell2
          pbmc3k
                      4903
                                   1352 3.7935958
Cell3
                      3147
          pbmc3k
                                   1129 0.8897363
Cell4
          pbmc3k
                      2639
                                    960 1.7430845
Cell5
         pbmc3k
                       980
                                    521 1.2244898
Cell6
          pbmc3k
                      2163
                                    781 1.6643551
```

See how analyses results are added to the same container. The idea is to keep all the experimental information together. This was a philosophic choice maide by the Bioconductor developers, inspired by the MIAME requirements and how data are stored on genomics databases like GEO

## Visualization

```
# Visualize QC metrics as a violin plot
(plt <- VlnPlot(pbmc, features = c("nFeature_RNA", "nCount_RNA", "percent.mt"), ncol = 3))</pre>
```

# Normalization

pbmc <- NormalizeData(pbmc)</pre>

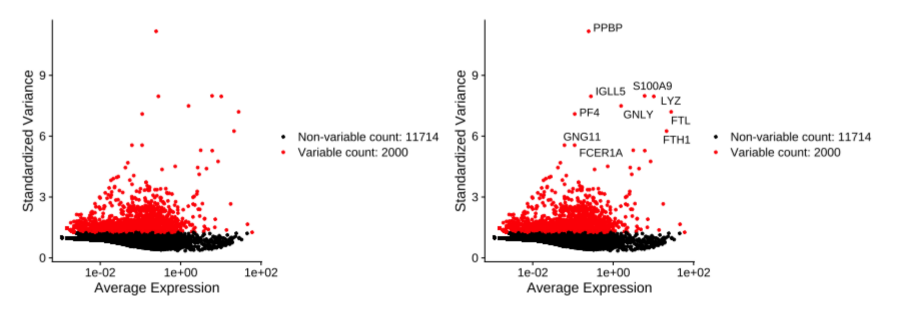
Note, we're saving in the same object

#### Feature selection

```
rownames(pbmc[['RNA']]@meta.features) <- r2
pbmc <- FindVariableFeatures(pbmc, selection.method = "vst", nfeatures = 2000)

# Identify the 10 most highly variable genes
top10 <- head(VariableFeatures(pbmc), 10)

# plot variable features with and without labels
plot1 <- VariableFeaturePlot(pbmc)
plot2 <- LabelPoints(plot = plot1, points = top10, repel = TRUE)
CombinePlots(plots = list(plot1, plot2))</pre>
```



#### **PCA**

```
pbmc <- ScaleData(pbmc)
pbmc <- RunPCA(pbmc, features = VariableFeatures(object = pbmc))
print(pbmc[["pca"]], dims = 1:5, nfeatures = 5)</pre>
```

```
PC_ 1
Positive: MALAT1, LTB, IL32, CD2, ACAP1
Negative: CST3, TYROBP, LST1, AIF1, FTL
PC_ 2
Positive: CD79A, MS4A1, TCL1A, HLA-DQA1, HLA-DRA
Negative: NKG7, PRF1, CST7, GZMA, GZMB
PC_ 3
Positive: HLA-DQA1, CD79A, CD79B, HLA-DQB1, HLA-DPB1
Negative: PPBP, PF4, SDPR, SPARC, GNG11
PC_ 4
Positive: HLA-DQA1, CD79A, CD79B, HIST1H2AC, HLA-DQB1
Negative: VIM, S100A8, S100A6, S100A4, S100A9
PC_ 5
Positive: GZMB, FGFBP2, NKG7, GNLY, PRF1
Negative: LTB, VIM, AQP3, PPA1, MAL
```

Important to note that each step just adds elements to the Seurat object

### **PCA**

DimPlot(pbmc, reduction = "pca")

### t-SNE

```
pbmc <- RunTSNE(pbmc, dims=1:10)
DimPlot(pbmc, reduction='tsne')</pre>
```

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# Heatmaps

#### Heatmaps

There are several ways of doing heatmaps in R:

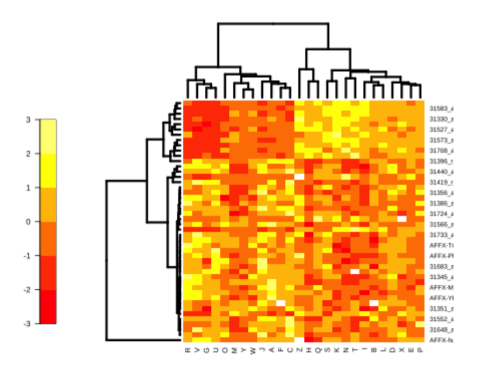
- http://sebastianraschka.com/Articles/heatmaps\_in\_r.html
- https://plot.ly/r/heatmaps/

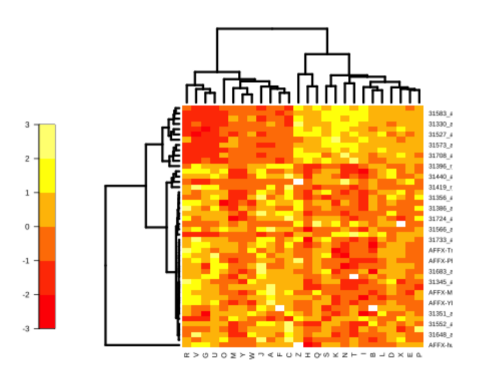
- http://moderndata.plot.ly/interactive-heat-maps-for-r/
  http://www.siliconcreek.net/r/simple-heatmap-in-r-with-ggplot2
  https://rud.is/b/2016/02/14/making-faceted-heatmaps-with-ggplot2/

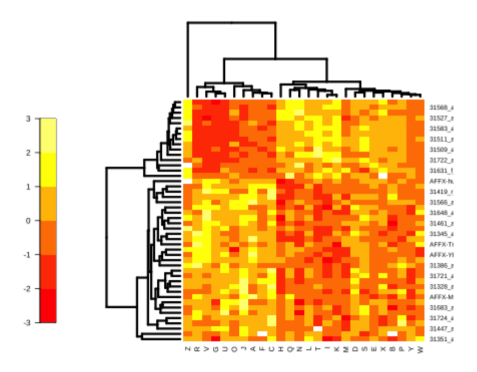
## Some example data

```
library(Biobase)
data(sample.ExpressionSet)
exdat <- sample.ExpressionSet
library(limma)
design1 <- model.matrix(~type, data=pData(exdat))
lm1 <- lmFit(exprs(exdat), design1)
lm1 <- eBayes(lm1) # compute linear model for each probeset
geneID <- rownames(topTable(lm1, coef=2, num=100, adjust='none',p.value=0.05))
exdat2 <- exdat[geneID,] # Keep features with p-values < 0.05
exdat2</pre>
```

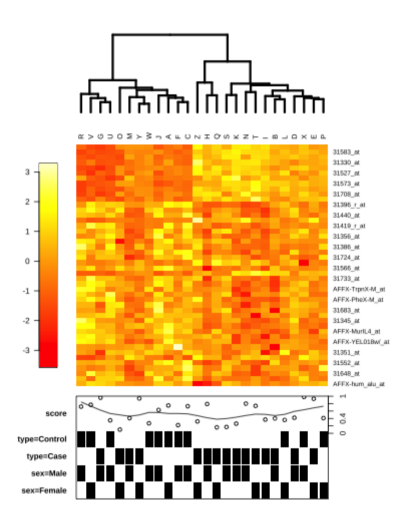
```
ExpressionSet (storageMode: lockedEnvironment)
assayData: 46 features, 26 samples
   element names: exprs, se.exprs
protocolData: none
phenoData
   sampleNames: A B ... Z (26 total)
   varLabels: sex type score
   varMetadata: labelDescription
featureData: none
experimentData: use 'experimentData(object)'
Annotation: hgu95av2
```

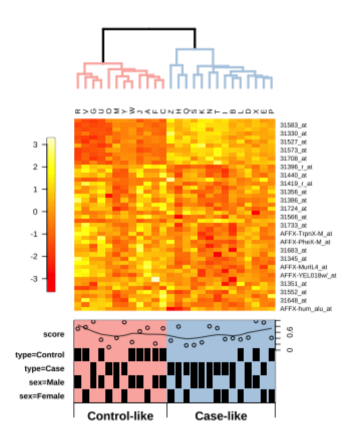






ann1 <- annHeatmap(exprs(exdat2), ann=pData(exdat2), col = heat.colors)
plot(ann1)</pre>





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#### Link

Put your mouse over each point :)

# Resources

- Organizing Single Cell Analysis with BioconductorBioconductor Courses
- 2019 Bioconductor workshops