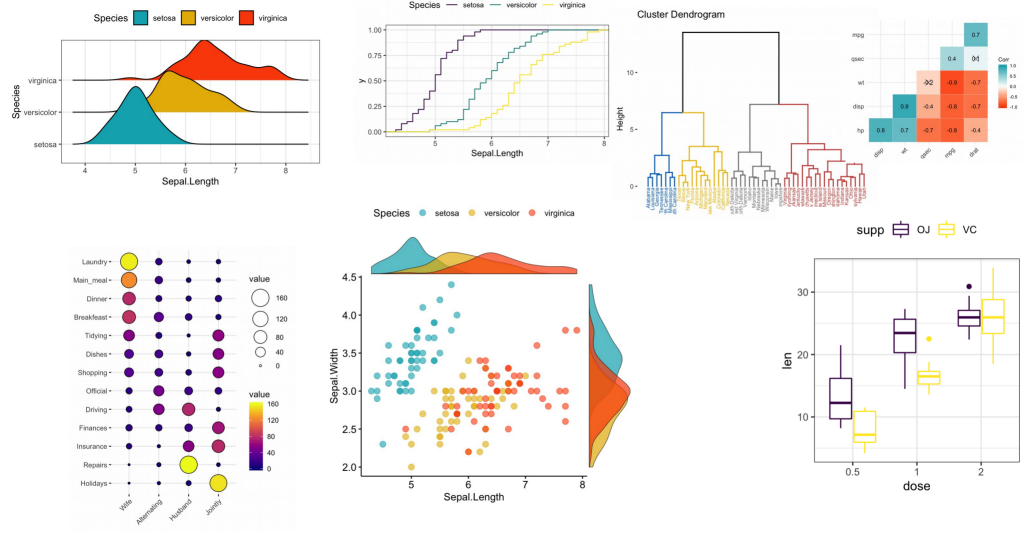
# Illustrating expression data in R

Bioinformatics workshops Jakob Willforss 191023

# What is ggplot2?

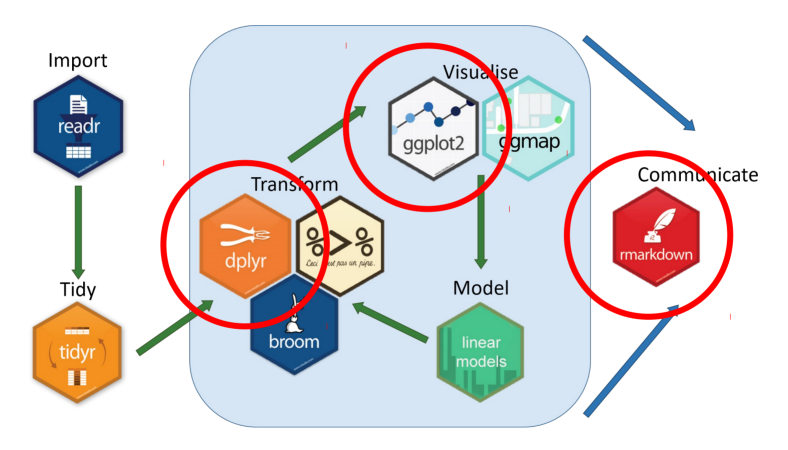
- Data visualization package in R
- Created by Hadley Wickham
- Part of a collection of R packages called **Tidyverse**
- Very flexible and powerful tool for visualizations
- Master a limited set of ideas and you can do a wide array of reproducible visualizations

The difficult part when working with ggplot is not ggplot itself, but getting the input data in the correct format!



Some examples from: https://www.datanovia.com/en/blog/ggplot-examples-best-reference/All on public datasets with runnable code available

#### Tidyverse: A collection of R packages



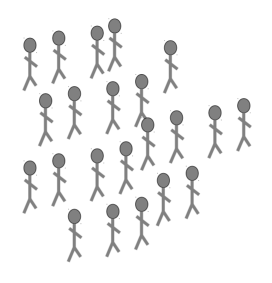
# Demonstration example: Array data

#### Setup:

- 72 array samples
- Survival times
- MIPIcat levels

Preprocessing:

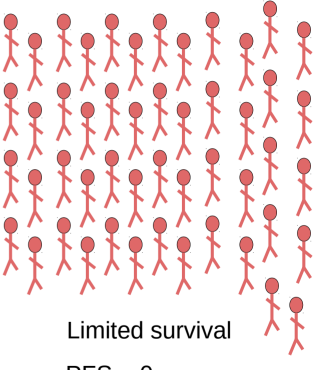
NormalyzerDE



Long survival

**PFS = 1** 

Patients: 22



PFS = 0

Patients: 50

# The data and the design

Design matrix

#### Sample information

sample	PFS	PFS_ye ars	MIPIcat
id1	0	4.3	1
id2	1	9.3	2
id3	1	9.5	1
id4	0	2.3	3

Data matrix

#### Gene information

Gen e	FDR	fold	id1	id2	id3	id4
Α	0.1	1.2	7.5	5.4	4.5	6.5
В	0.9	-2.3	10.2	9.8	11.2	10.3

The sample names is found in one column in the design and should be present as columns in the data

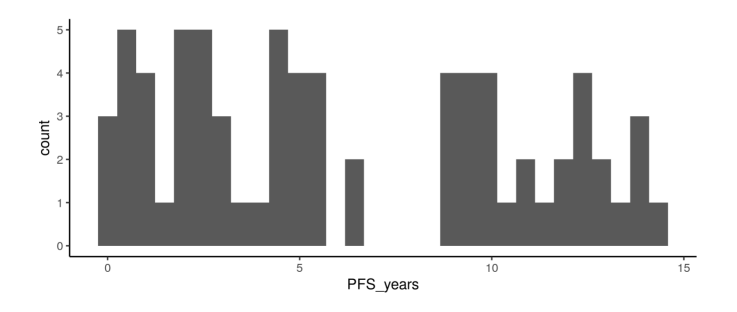
# Loading design matrix

```
> design fp ← "joana data/joana design.tsv"
> design df ← read.table(design_fp, sep="\t", header=TRUE, stringsAsFactors=FALSE)
> dim(design df)
[1] 72 5
> colnames(design df)
[1] "ID"
             "PFS"
                        "PFS years" "MIPIcat" "array_id"
> head(design df)
ID<chr> PFS<int> PFS years<dbl> MIPIcat<int> array id<chr>
    MCL2 006 1 1.547 3 p1615 01 MCL2 006.CEL
                             p1615 02 MCL2 007.CEL
   MCL2 007 1 12.686 1
                             p1615 43 MCL2 008.CEL
    MCL2 008 0 13.994 1
                             p1615 04 MCL2 013.CEL
    MCL2 013 0 12.458 1
   MCL2 031 1 13.100 1
                             p1615 06 MCL2 031.CEL
                              p1615 08 MCL2 032.CEL
    MCL2 032 0 12.175 1
```

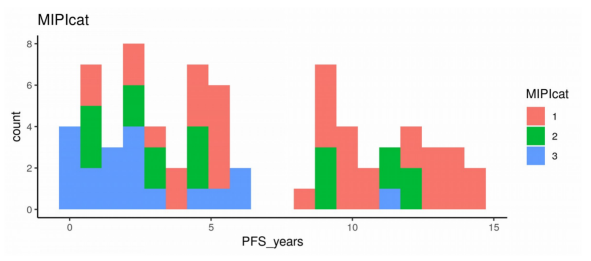
array\_id column matches sample names in data matrix

# Visualizing PFS\_years

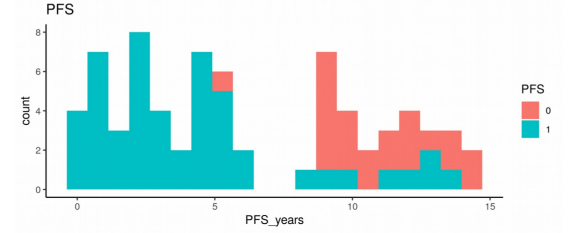
> ggplot(design\_df, aes(x=PFS\_years)) + geom\_histogram()



> ggplot(design\_df, aes(x=PFS\_years)) + geom\_histogram()



> ggplot(design\_df, aes(x=PFS\_years)) + geom\_histogram()



# The ggplot2 command

The target data frame

ggplot(design\_df, aes(x=PFS\_years)) + geom\_histogram()

Specify aesthetics columns from the target data

Common aesthetics are: x, y, color, fill, label, shape

geometric - how should the data with aesthetics be illustrated

Common geoms are: geom\_histogram() geom\_point() geom\_line() geom\_col() ... and many many more

# The data and the design

Design matrix

#### Sample information

sample	PFS	PFS_ye ars	MIPIcat
id1	0	4.3	1
id2	1	9.3	2
id3	1	9.5	1
id4	0	2.3	3

Data matrix

#### Gene information

Gen e	FDR	fold	id1	id2	id3	id4
Α	0.1	1.2	7.5	5.4	4.5	6.5
В	0.9	-2.3	10.2	9.8	11.2	10.3

The sample names is found in one column in the design and should be present as columns in the data

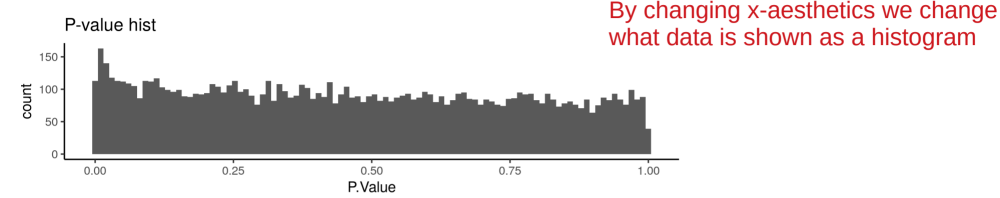
#### Let's demonstrate on the data

```
> data fp ← "joana data/normalyzerde stats.tsv"
> data df ← read.table(data fp, sep="\t", header=TRUE, stringsAsFactors=FALSE)
> dim(data df)
[1] 9190 80
> colnames(data df)
[1] "PROBEID"
                             "SYMBOL"
                                                      "GENENAME"
                                                                               "array.iD"
                                                      "log2Fold"
 [5] "P.Value"
                              "adi.P.Val"
                                                                                "AvgExpr"
 [9] "p1615_01_MCL2_006.CEL" "p1615_02_MCL2_007.CEL" "p1615_43_MCL2_008.CEL" "p1615_04_MCL2_013.CEL"
[13] "p1615_06_MCL2_031.CEL" "p1615_08_MCL2_032.CEL" "p1615_61_MCL2_038.CEL" "p1615_52_MCL2_039.CEL"
[13] "p1615 06 MCL2 031.CEL"
[17] "p1615 10 MCL2 048.CEL"
                              "p1615 11 MCL2 054.CEL"
                                                       "p1615 12 MCL2 058.CEL" "p1615 13 MCL2 059.CEL"
> head(data df)
SYMBOL <chr>
                    P.Value <dbl> adj.P.Val <dbl> log2Fold <dbl> AvgExpr <dbl> (then sample cols)
LOC100287497
                    0.8679910
                                                  0.01752942
                                                                 7.002756 6.483045
                                  0.9812370
                                  0.9715940
                                                  0.01497248
                                                                 4.296178 4.449927
LINC01128
                    0.8088450
SAMD11
                   0.8325184
                                   0.9743369
                                                  0.01496062
                                                                 6.343670 6.143292
                   0.2982318
                                   0.8626470
                                                  -0.04156739
                                                                 5.837616 5.759693
KLHL17
                                                                 6.342759 6.108941
                   0.1575169
                                                  -0.08993811
PLEKHN1
                                   0.7926791
ISG15
                    0.1390773
                                   0.7756221
                                                  0.14539873
                                                                 6.195273 6.515469
```

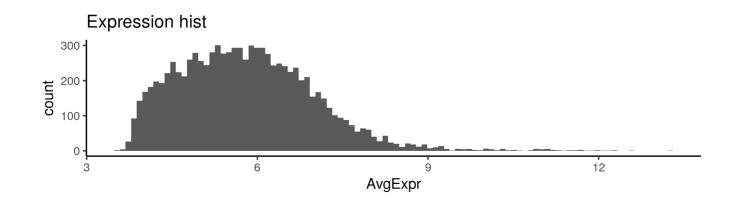
# Specifying what features are 'significant'

```
> fdr thres \leftarrow 0.25
> p thres ← 0.05
> data df$IsPSig ← data df$P.Value 
> data df$IsFDRSig ← data df$adj.P.Val < fdr thres</pre>
> # head(data df) to double check
> table(data df$IsSig)
FALSE TRUE
 8487 702
> table(data df$IsFDRSig)
FALSE TRUE
 9178
```

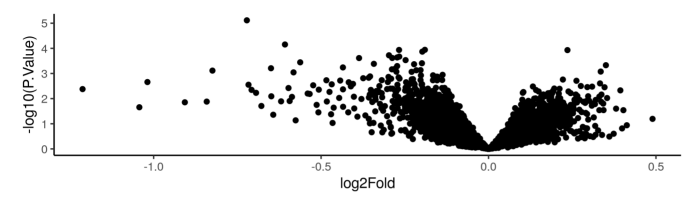
> ggplot(data\_df, aes(x=P.Value)) + geom\_histogram(bins=100) +
ggtitle("P-value hist")



> ggplot(data\_df, aes(x=AveExpr)) + geom\_histogram(bins=100) +
ggtitle("Expression hist")

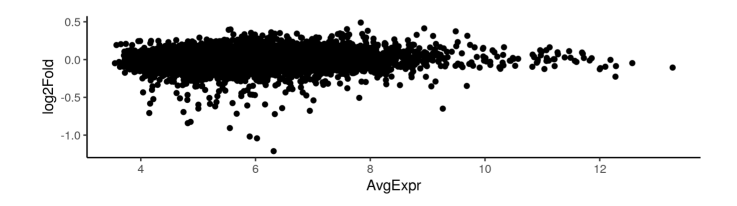


> ggplot(data\_df, aes(x=log2Fold, y=-log10(P.Value))) + geom\_point()

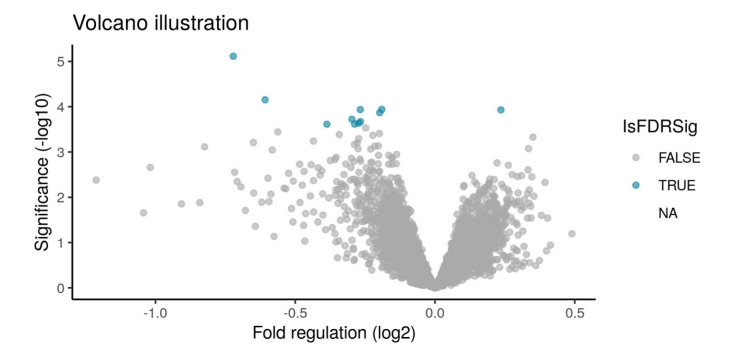


Again, we simply change the aesthetics to use different columns in data\_df

> ggplot(data\_df, aes(x=AvgExpr, y=log2Fold)) + geom\_point()



```
gray_blue_colors ← c("#AAAAAA", "#0C81A2")
ggplot(full_data_df, aes(x=AvgExpr, y=log2Fold, color=IsFDRSig)) +
    geom_point(alpha=0.6) +
    ggtitle("MA") +
    xlab("Expression level") +
    ylab("Fold change (log2)") +
    scale_color_manual(values=gray_blue_colors)
```



## Hands-on time!

### Single feature illustration

We want to: Illustrate the feature with lowest FDR

# Single feature illustration

```
> target cols ← c("SYMBOL", "GENENAME", "adj.P.Val", "log2Fold")
> top hits ← head(arrange(data df, P.Value), 5)
> top hits[, target cols]
SYMBOL <chr>
              GENENAME <chr>
                                               adj.P.Val <dbl> log2Fold <dbl>
                                              0.1445887
SH3BGRL2
              SH3 domain binding glutamate ...
                                                             -0.7216677
              keratin 19
KRT19
                                               0.2184205
                                                             -0.6078616
              solute carrier family 2 member 10 0.2184205
SLC2A10
                                                             -0.1900941
              pleckstrin homology domain ... 0.2184205
PLEKHA5
                                                             -0.2671958
              toll like receptor 9
TLR9
                                               0.2184205
                                                             0.2356864
```

**arrange** is a Tidyverse command organizing the data frame in rising order based on given column - note that the P.Value column is specified without quotes (""), same as within ggplot's aes

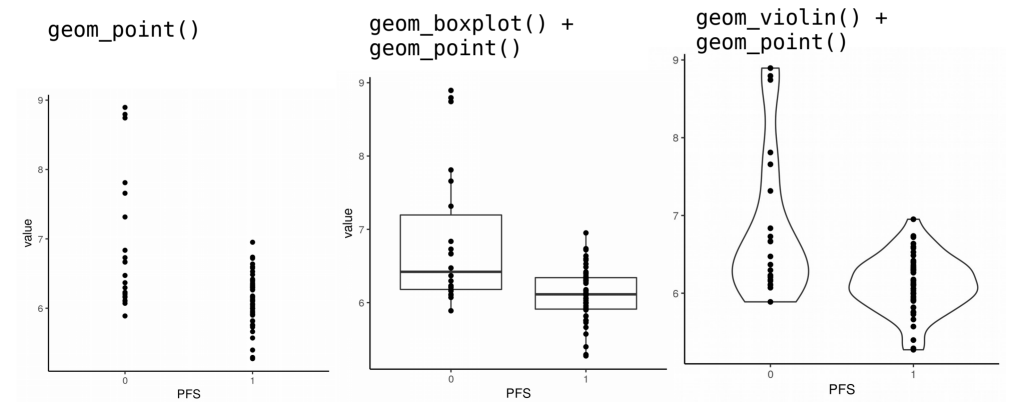
top\_hits[, target\_cols] slices out only the given columns

# Getting the best hits

```
> best hit ← data df[data df$SYMBOL == "SH3BGRL2", ]
> best hit vals <- best hit[, design df$array id]</pre>
> best hit df <- cbind(value=unlist(best_hit_vals), design_df)</pre>
                      We bind the values from SH3BGRL2 to the design matrix,
                      calling the new column 'value'
> head(best hit df, 4)
value <dbl> ID <chr> PFS <fctr> PFS years <dbl> MIPIcat <fctr> array_id <chr>
5.290603 MCL2_006 1 1.547 3 p1615_01_MCL2_006.CEL
6.329583 MCL2_007 1 12.686 1 p1615_02_MCL2_007.CEL
7.810174 MCL2 008 0 13.994 1 p1615 43 MCL2 008.CEL
          MCL2 013
                      0 12.458 1 p1615 04 MCL2 013.CEL
6.158796
```

```
ggplot(best_hit_df, aes(x=PFS, y=value)) + geom_point()
ggplot(best_hit_df, aes(x=PFS, y=value)) + geom_boxplot() + geom_point()
ggplot(best_hit_df, aes(x=PFS, y=value)) + geom_violin() + geom_point()
```

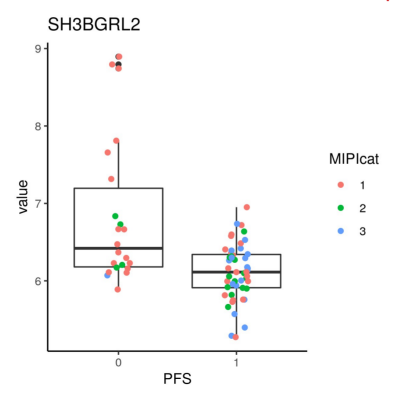
Here, we illustrate the data differently simply by changing the geom-layers



```
ggplot(best_hit_df, aes(x=PFS, y=value)) +
    geom_boxplot() +
    geom_point(position=position jitter(0.1), aes(color=MIPIcat)) +
    gqtitle("SH3BGRL2")
```

Jitter 'shakes up' the positions of the dots

If we only want to color one layer, we specify the aesthetic within it



## Hands-on time!

## Wide versus long format

- Expression data is commonly found in "wide" format
- ggplot expects the input to be in "long" format Long format

#### Wide format

Gene	ID1	ID2	ID3
gene_A	9.4	7.5	8.4
gene_B	6.5	6.7	7.3

Gene	ID	value
gene_A	ID1	9.4
gene_A	ID2	7.5
gene_A	ID3	8.4
gene_B	ID1	6.5
gene_B	ID2	6.7
gene_B	ID3	7.3

# Converting to long format

```
> dim(data df)
9190 82
> long df ← pivot longer(data df, design df$sample,
names to="array id", values to="value")
> dim(long df)
661680 12
> head(long df)[, c("SYMBOL", "sample", "value")]
SYMBOL <chr>
               array id <chr>
                                       value <dbl>
               p1615 01 MCL2 006.CEL
                                       6.483045
L0C100287497
               p1615 02 MCL2 007.CEL
L0C100287497
                                       6.878746
               p1615 43 MCL2 008.CEL
L0C100287497
                                       8.497729
               p1615 04 MCL2 013.CEL
L0C100287497
                                       7.477194
               p1615 06 MCL2 031.CEL
                                       7.058754
L0C100287497
               p1615 08 MCL2 032.CEL
L0C100287497
                                       6.819554
```

Now we have only one column with values, with the "array\_id" column specifying from where each value comes

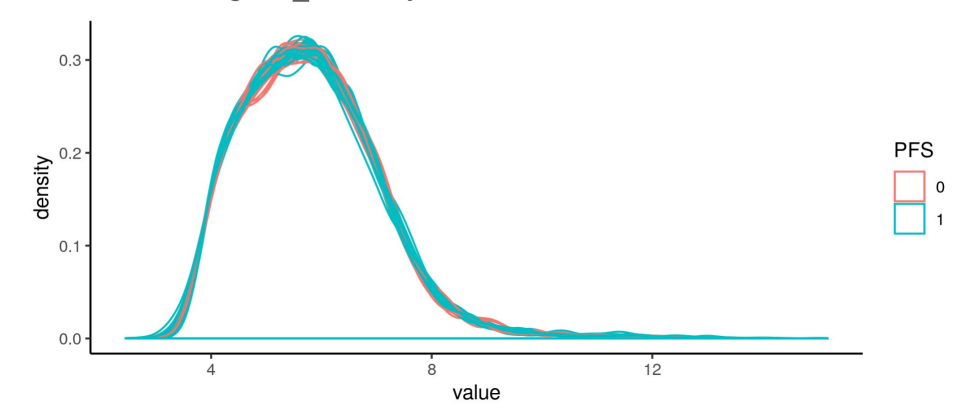
# Adding sample annotations

```
> annot long df ← left join(long df, design df, by=
"array id")
> head(annot long df)
ID <chr> PFS <fctr> PFS years <dbl> MIPIcat <fctr>
p1615 01 MCL2 006.CEL
                       6.483045 MCL2 006 1
                                            1.547 3
p1615 02 MCL2 007.CEL
                       6.878746 MCL2 007 1 12.686
p1615 43 MCL2 008.CEL
                       8.497729 MCL2 008 0 13.994
p1615 04 MCL2 013.CEL
                      7.477194 MCL2 013 0 12.458
p1615 06 MCL2 031.CEL
                      7.058754 MCL2 031 1
                                           13.100
p1615 08 MCL2 032.CEL
                       6.819554 MCL2 032 0
                                            12.175
```

"left\_join" from Tidyverse lets us merge information from another matrix by specifying a common column. Here, the design-matrix information is added based on the array IDs. Now we have everything we need for sample-wide illustrations.

## The density plot

```
> ggplot(annot_long_data, aes(x=value, group=array_id,
color=PFS)) + geom density()
```



## Sample boxplots

```
> ggplot(annot long data, aes(x=array id, y=value,
color=PFS)) + geom boxplot() + theme(axis.text.x =
element text(angle=90, hjust=1)
gqtitle("Sample intensity levels")
                                                 Rotate axis-x labels
 Sample intensity levels
                             array id
```

# Saving your plots

```
> plt_obj ← ggplot(annot_long_data, aes(x=value,
group=array_id, color=PFS)) + geom_density()
> ggsave(plt_obj, filename="plots/density.png", width=7,
height=7, dpi=300)
```

- plt\_obj An object containing the plot
- filename Where to write the new plot. Note that the file ending is important here we generate a figure in PNG format
- width / height Size specified in inches
- dpi The resolution of the written figure

# Investigating further plots

Chord diagrams

https://www.r-graph-gallery.com/chord-diagram.html

Survival curves

https://rpkgs.datanovia.com/survminer/index.html

**Enrichment illustrations** 

https://yulab-smu.github.io/clusterProfiler-book/

# The End