# Sources of variability in label-free proteomics experiments

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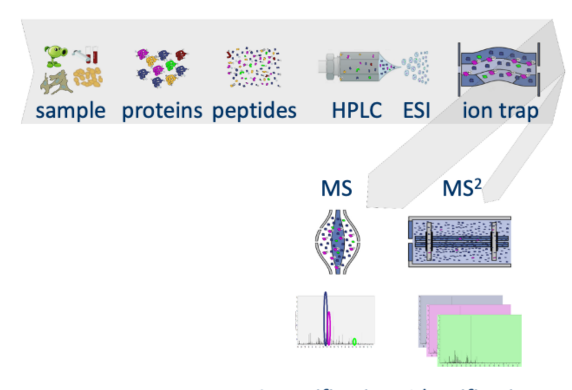
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# 1 Intro

- 1. Background + cptac study
- 2. Sources of variability
- 3. Summarization

The summary  $\cdot$ 

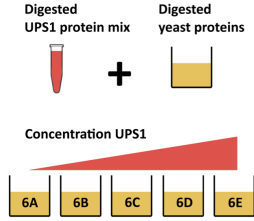
## 1.1 MS-based workflow



# Quantification Identification

- Peptide Characteristics
  - Modifications
  - Ionisation Efficiency: huge variability
  - Identification
    - \* Misidentification  $\rightarrow$  outliers
    - \* MS $^2$  selection on peptide abundance
    - \* Context depending missingness
    - \* Non-random missingness
- $\rightarrow$  Unbalanced pepide identifications across samples and messy data

# 1.2 CPTAC Spike-in Study



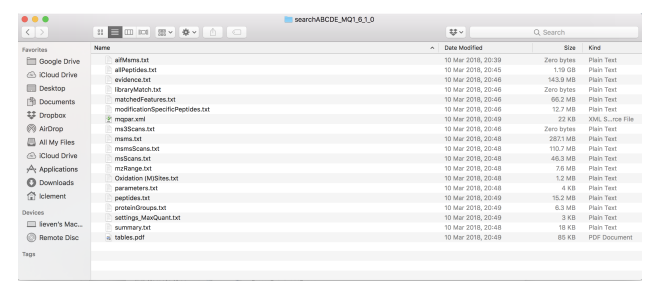
5 spike-in concentrations: 6A to 6E



- Same trypsin-digested yeast proteome background in each sample
- Trypsin-digested Sigma UPS1 standard: 48 different human proteins spiked in at 5 different concentrations (treatment A-E)
- Samples repeatedly run on different instruments in different labs
- After MaxQuant search with match between runs option
  - -41% of all proteins are quantified in all samples
  - 6.6% of all peptides are quantified in all samples

 $<sup>\</sup>rightarrow$  vast amount of missingness

#### 1.2.1 Maxquant output



#### 1.2.2 Read data

Click to see background and code

1. We use a peptides.txt file from MS-data quantified with maxquant that contains MS1 intensities summarized at the peptide level.

peptidesFile <- "https://raw.githubusercontent.com/statOmics/PDA21/data/quantification/fullCptacDatasSe</pre>

2. Maxquant stores the intensity data for the different samples in columns that start with Intensity. We can retreive the column names with the intensity data with the code below:

```
ecols <- grep("Intensity\\.", names(read.delim(peptidesFile)))</pre>
```

3. Read the data and store it in QFeatures object

```
pe <- readQFeatures(
  table = peptidesFile,
  fnames = 1,
  ecol = ecols,
  name = "peptideRaw", sep="\t")</pre>
```

#### 1.2.3 Design

Click to see background and code

```
## CharacterList of length 1
## [["peptideRaw"]] Intensity.6A_1 Intensity.6A_2 ... Intensity.6E_9
```

- Note, that the sample names include the spike-in condition.
- They also end on a number.

```
- 1-3 is from lab 1,
```

- 4-6 from lab 2 and
- 7-9 from lab 3.
- We update the colData with information on the design

```
colData(pe)$lab <- rep(rep(paste0("lab",1:3),each=3),5) %>% as.factor
colData(pe)$condition <- pe[["peptideRaw"]] %>% colnames %>% substr(12,12) %>% as.factor
colData(pe)$spikeConcentration <- rep(c(A = 0.25, B = 0.74, C = 2.22, D = 6.67, E = 20),each = 9)</pre>
```

• We explore the colData

```
colData(pe)
```

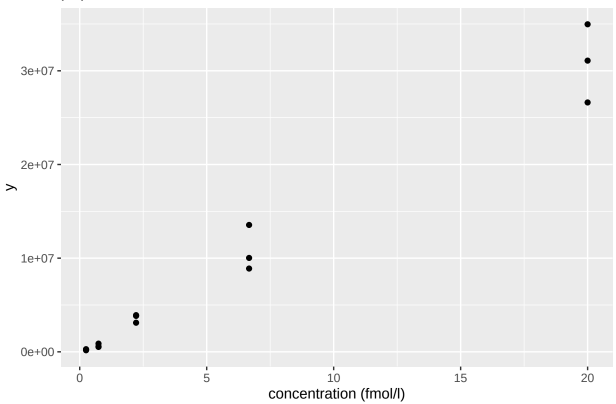
```
## DataFrame with 45 rows and 3 columns
##
                        lab condition spikeConcentration
##
                  <factor> <factor>
                                                <numeric>
## Intensity.6A_1
                      lab1
                                    Α
                                                     0.25
## Intensity.6A_2
                      lab1
                                    Α
                                                     0.25
## Intensity.6A_3
                      lab1
                                    Α
                                                     0.25
## Intensity.6A_4
                      lab2
                                    Α
                                                     0.25
## Intensity.6A_5
                                    Α
                                                     0.25
                      lab2
## ...
                        . . .
## Intensity.6E_5
                                                       20
                      lab2
                                    Ε
## Intensity.6E_6
                      lab2
                                    Ε
                                                       20
                                    Ε
## Intensity.6E_7
                                                       20
                      lab3
## Intensity.6E_8
                                    Ε
                                                       20
                      lab3
                                    Ε
                                                       20
## Intensity.6E_9
                       lab3
```

# 2 Sources of variation

## 2.1 Intensities of one peptide

Peptide AALEELVK from spiked-in UPS protein P12081. We only show data from lab1.

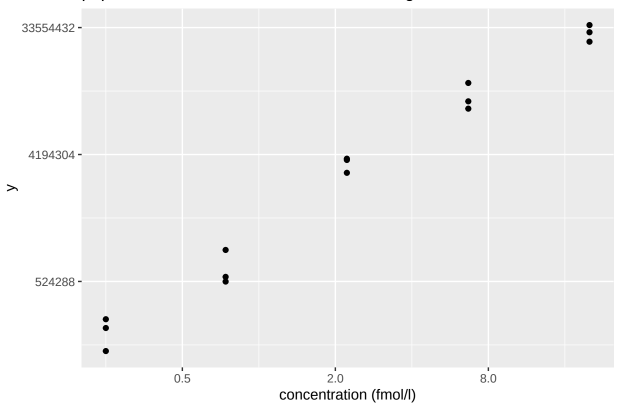
# peptide AALEELVK in lab1



• Variance increases with the mean  $\rightarrow$  Multiplicative error structure

## plotLog

# peptide AALEELVK in lab1 with axes on log scale



- Data seems to be homoscedastic on log-scale  $\rightarrow$  log transformation of the intensity data
- In quantitative proteomics analysis on log<sub>2</sub>
- $\rightarrow$  Differences on a  $\log_2$  scale:  $\log_2$  fold changes

$$\log_2 B - \log_2 A = \log_2 \frac{B}{A} = \log F C_{\text{B - A}}$$
 
$$log_2 F C = 1 \rightarrow F C = 2^1 = 2$$
 
$$log_2 F C = 2 \rightarrow F C = 2^2 = 4$$

# 2.2 Log-transform

Click to see code to log-transfrom the data

• We calculate how many non zero intensities we have for each peptide and this can be useful for filtering.

 $\bullet$  Peptides with zero intensities are missing peptides and should be represent with a NA value rather than 0.

```
pe <- zeroIsNA(pe, "peptideRaw") # convert 0 to NA
```

• Logtransform data with base 2

```
pe <- logTransform(pe, base = 2, i = "peptideRaw", name = "peptideLog")</pre>
```

## 2.3 Filtering

Click to see code to filter the data

1. Handling overlapping protein groups

In our approach a peptide can map to multiple proteins, as long as there is none of these proteins present in a smaller subgroup.

```
pe <- filterFeatures(pe, ~ Proteins %in% smallestUniqueGroups(rowData(pe[["peptideLog"]])$Proteins))</pre>
```

2. Remove reverse sequences (decoys) and contaminants

We now remove the contaminants, peptides that map to decoy sequences, and proteins which were only identified by peptides with modifications.

```
pe <- filterFeatures(pe,~Reverse != "+")
pe <- filterFeatures(pe,~ Potential.contaminant != "+")</pre>
```

3. Drop peptides that were only identified in one sample

We keep peptides that were observed at last twice.

```
pe <- filterFeatures(pe,~ nNonZero >=2)
nrow(pe[["peptideLog"]])
```

```
## [1] 10478
```

We keep 10478 peptides upon filtering.

## 2.4 Technical Variability

Click to see code for plot

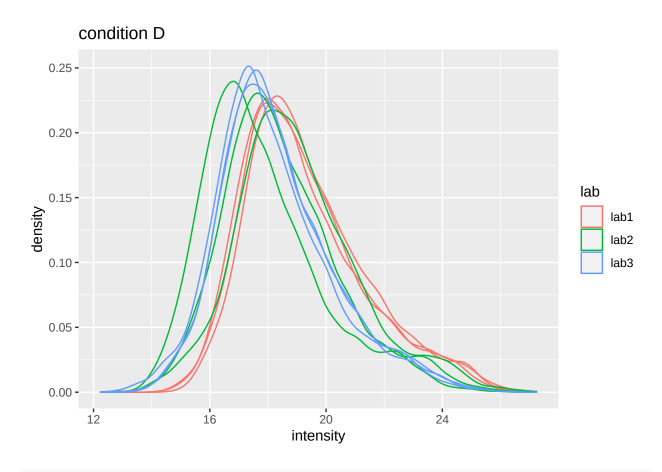
```
densityConditionD <- pe[["peptideLog"]][,colData(pe)$condition=="D"] %>%
  assay %>%
  as.data.frame() %>%
  gather(sample, intensity) %>%
  mutate(lab = colData(pe)[sample,"lab"]) %>%
  ggplot(aes(x=intensity,group=sample,color=lab)) +
    geom_density() +
```

```
ggtitle("condition D")

densityLab2 <- pe[["peptideLog"]][,colData(pe)$lab=="lab2"] %>%
   assay %>%
   as.data.frame() %>%
   gather(sample, intensity) %>%
   mutate(condition = colData(pe)[sample,"condition"]) %>%
   ggplot(aes(x=intensity,group=sample,color=condition)) +
   geom_density() +
   ggtitle("lab2")
```

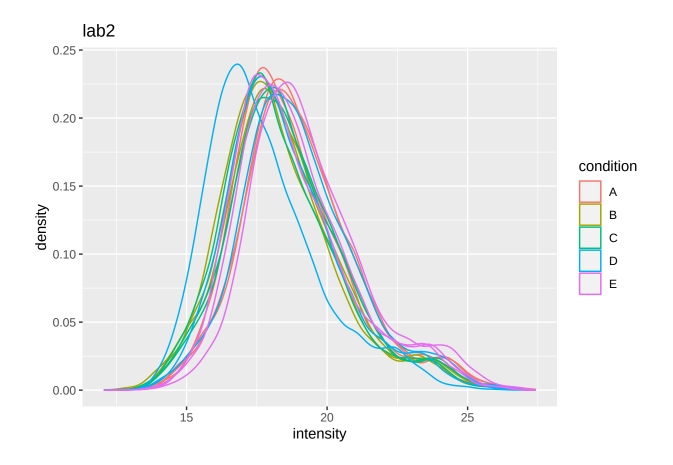
densityConditionD

## Warning: Removed 39179 rows containing non-finite values (stat\_density).



densityLab2

## Warning: Removed 44480 rows containing non-finite values (stat\_density).



- Even in very clean synthetic dataset (same background, only 48 UPS proteins can be different) the marginal peptide intensity distribution across samples can be quite distinct
- Considerable effects between and within labs for replicate samples
- Considerable effects between samples with different spike-in concentration

# $\rightarrow$ Normalization is needed

## 2.5 Normalization

Normalization of the data by median centering

$$y_{ip}^{\text{norm}} = y_{ip} - \hat{\mu}_i$$

with  $\hat{\mu}_i$  the median intensity over all observed peptides in sample i.

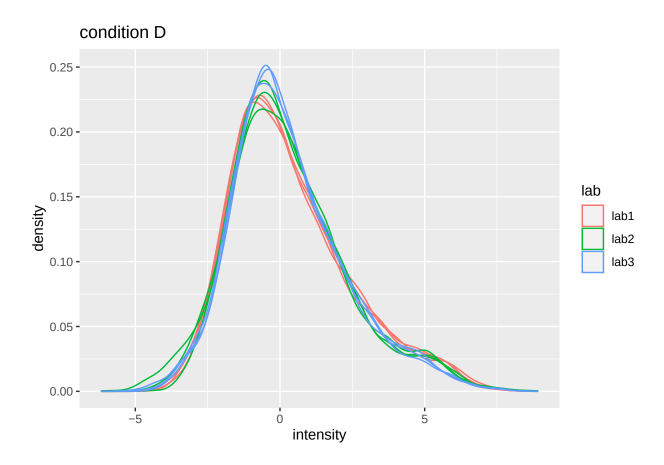
Click to see R-code to normalize the data

Click to see code to make plot

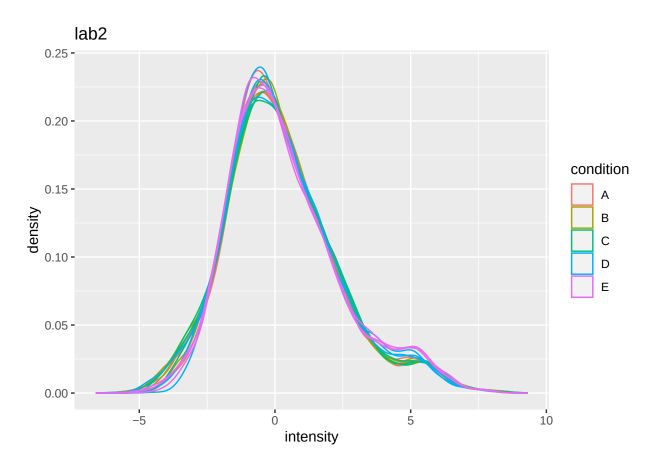
```
densityConditionDNorm <- pe[["peptideNorm"]][,colData(pe)$condition=="D"] %>%
  assay %>%
  as.data.frame() %>%
  gather(sample, intensity) %>%
  mutate(lab = colData(pe)[sample, "lab"]) %>%
  ggplot(aes(x=intensity,group=sample,color=lab)) +
    geom_density() +
    ggtitle("condition D")
densityLab2Norm <- pe[["peptideNorm"]][,colData(pe)$lab=="lab2"] %>%
  assay %>%
  as.data.frame() %>%
  gather(sample, intensity) %>%
  mutate(condition = colData(pe)[sample,"condition"]) %>%
  ggplot(aes(x=intensity,group=sample,color=condition)) +
    geom_density() +
    ggtitle("lab2")
```

 ${\tt densityConditionDNorm}$ 

## Warning: Removed 39179 rows containing non-finite values (stat\_density).



## Warning: Removed 44480 rows containing non-finite values (stat\_density).

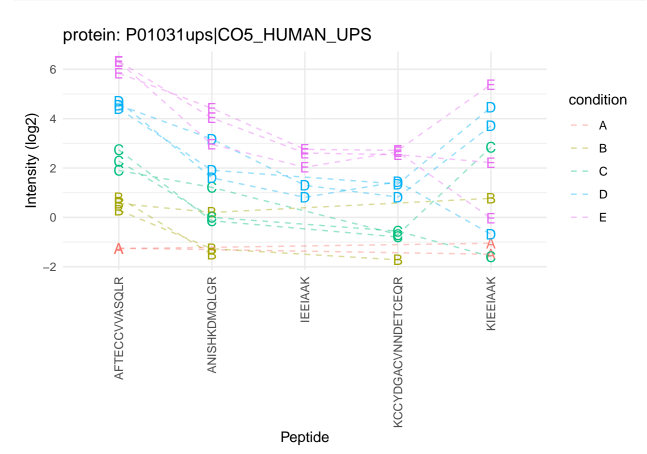


## 2.6 Pseudo replication

```
prot <- "P01031ups|C05_HUMAN_UPS"
data <- pe[["peptideNorm"]][
    rowData(pe[["peptideNorm"]])$Proteins == prot,
    colData(pe)$lab=="lab3"] %>%
    assay %>%
    as.data.frame %>%
    rownames_to_column(var = "peptide") %>%
    gather(sample, intensity, -peptide) %>%
    mutate(condition = colData(pe)[sample,"condition"]) %>%
    na.exclude
sumPlot <- data %>%
    ggplot(aes(x = peptide, y = intensity, color = condition, group = sample, label = condition), show.legeom_text(show.legend = FALSE) +
    theme_minimal() +
    theme(axis.text.x = element_text(angle = 90, vjust = 0.5, hjust = 1)) +
```

```
xlab("Peptide") +
ylab("Intensity (log2)") +
ggtitle(paste0("protein: ",prot))
```

```
sumPlot +
geom_line(linetype="dashed",alpha=.4)
```



- Sources of variability in plot:
  - Between treatment variability
  - Between sample variability
  - Between peptide variability
  - within sample variability
- Multiple peptides from same protein in a sample
- Peptide intensities in the same sample are correlated: Pseudo replication

## $\rightarrow$ Summarization!

- Strong peptide effect
- Unbalanced peptide identification

#### 2.6.1 Illustration on subset of CPTAC study: A vs B comparison in lab 3

#### 2.6.1.1 LFQ Click to see background and code

1. Import data

```
proteinsFile <- "https://raw.githubusercontent.com/statOmics/PDA21/data/quantification/cptacAvsB_lab3/p
ecols <- grep("LFQ\\.intensity\\.", names(read.delim(proteinsFile)))

peLFQ <- readQFeatures(
   table = proteinsFile, fnames = 1, ecol = ecols,
   name = "proteinRaw", sep = "\t"
)

cond <- which(
   strsplit(colnames(peLFQ)[[1]][1], split = "")[[1]] == "A") # find where condition is stored

colData(peLFQ)$condition <- substr(colnames(peLFQ), cond, cond) %>%
   unlist %>%
   as.factor
```

2. Preprocessing

3. Modeling and Inference

#### 2.6.1.2 Median & robust summarization Click to see background and code

1. Import Data

```
peptidesFile <- "https://raw.githubusercontent.com/stat0mics/SGA2020/data/quantification/cptacAvsB_lab3
ecols <- grep(
   "Intensity\\.",
   names(read.delim(peptidesFile))
)

peAB <- readQFeatures(
   table = peptidesFile,
   fnames = 1,
   ecol = ecols,
   name = "peptideRaw", sep="\t")

cond <- which(
   strsplit(colnames(peAB)[[1]][1], split = "")[[1]] == "A") # find where condition is stored

colData(peAB)$condition <- substr(colnames(peAB), cond, cond) %>%
   unlist %>%
   as.factor
```

2. Preprocessing

```
rowData(peAB[["peptideRaw"]])$nNonZero <- rowSums(assay(peAB[["peptideRaw"]]) > 0)

peAB <- zeroIsNA(peAB, "peptideRaw") # convert 0 to NA

peAB <- logTransform(peAB, base = 2, i = "peptideRaw", name = "peptideLog")

peAB <- filterFeatures(peAB, ~ Proteins %in% smallestUniqueGroups(rowData(peAB[["peptideLog"]])$Protein

peAB <- filterFeatures(peAB, ~ Reverse != "+")

peAB <- filterFeatures(peAB, ~ Potential.contaminant != "+")

peAB <- filterFeatures(peAB, ~ nNonZero >=2)

nrow(peAB[["peptideLog"]])
```

## [1] 7011

```
name = "proteinMedian",
fun = matrixStats::colMedians)

peAB <- aggregateFeatures(peAB,
    i = "peptideNorm",
    fcol = "Proteins",
    na.rm = TRUE,
    name = "proteinRobust")</pre>
```

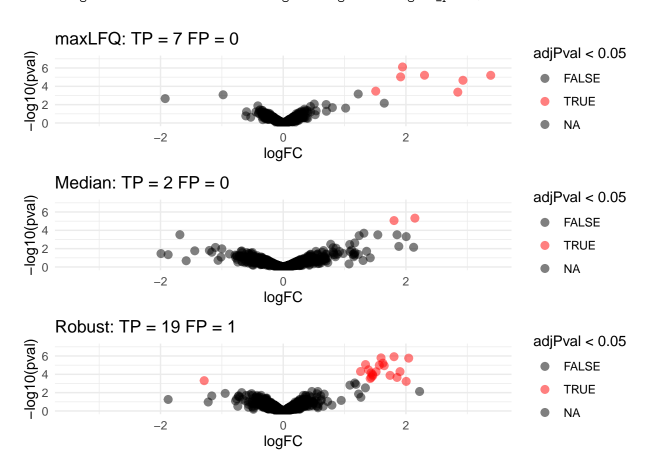
```
3. Modeling and inference
peAB <- msqrob(object = peAB, i = "proteinMedian", formula = ~condition)</pre>
L <- makeContrast("conditionB=0", parameterNames = c("conditionB"))</pre>
peAB <- hypothesisTest(object = peAB, i = "proteinMedian", contrast = L)</pre>
peAB <- msqrob(object = peAB, i = "proteinRobust", formula = ~condition)</pre>
peAB <- hypothesisTest(object = peAB, i = "proteinRobust", contrast = L)</pre>
volcanoMedian <- ggplot(rowData(peAB[["proteinMedian"]])$conditionB,</pre>
                  aes(x = logFC, y = -log10(pval), color = adjPval < 0.05)) +
  geom_point(cex = 2.5) +
  scale_color_manual(values = alpha(c("black", "red"), 0.5)) +
  theme_minimal() +
  ggtitle(paste0("Median: TP = ",sum(rowData(peAB[["proteinMedian"]])$conditionB$adjPval<0.05&grepl(row.
volcanoRobust<- ggplot(rowData(peAB[["proteinRobust"]])$conditionB,</pre>
                  aes(x = logFC, y = -log10(pval), color = adjPval < 0.05)) +
  geom_point(cex = 2.5) +
  scale_color_manual(values = alpha(c("black", "red"), 0.5)) +
  theme_minimal() +
  ggtitle(paste0("Robust: TP = ",sum(rowData(peAB[["proteinRobust"]])$conditionB$adjPval<0.05&grepl(row.
ylims \leftarrow c(0,
           ceiling(max(c(-log10(rowData(peLFQ[["protein"]])$conditionB$pval),
               -log10(rowData(peAB[["proteinMedian"]])$conditionB$pval),
               -log10(rowData(peAB[["proteinRobust"]])$conditionB$pval)),
               na.rm=TRUE))
)
xlims <- max(abs(c(rowData(peLFQ[["protein"]])$conditionB$logFC,</pre>
               rowData(peAB[["proteinMedian"]])$conditionB$logFC,
               rowData(peAB[["proteinRobust"]])$conditionB$logFC)),
               na.rm=TRUE) * c(-1,1)
compBoxPlot <- rbind(rowData(peLFQ[["protein"]])$conditionB %>% mutate(method="maxLFQ") %>% rownames_to
      rowData(peAB[["proteinMedian"]])$conditionB %% mutate(method="median")%% rownames_to_column(var
      rowData(peAB[["proteinRobust"]])$conditionB%% mutate(method="robust")%% rownames_to_column(var=
      mutate(ups= grepl(protein,pattern="UPS")) %>%
    ggplot(aes(x = method, y = logFC, fill = ups)) +
    geom_boxplot() +
    geom_hline(yintercept = log2(0.74 / .25), color = "#00BFC4") +
    geom_hline(yintercept = 0, color = "#F8766D")
```

## 2.6.1.3 Comparison summarization methods

## Warning: Removed 746 rows containing missing values (geom\_point).

## Warning: Removed 166 rows containing missing values (geom\_point).

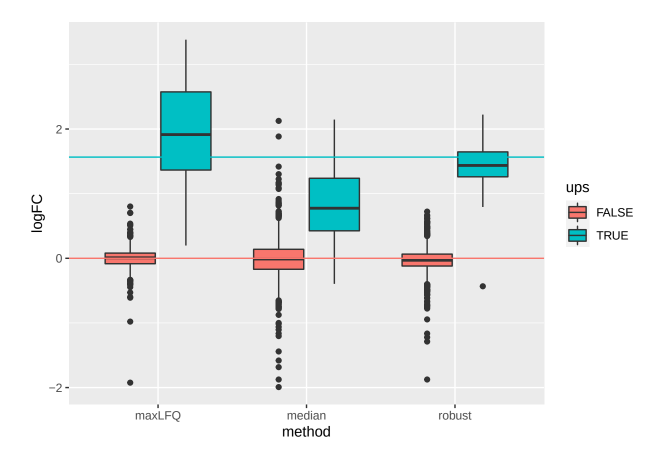
## Warning: Removed 167 rows containing missing values (geom\_point).



• Robust summarization: highest power and still good FDR control:  $FDP = \frac{1}{20} = 0.05$ .

## compBoxPlot

## Warning: Removed 1079 rows containing non-finite values (stat\_boxplot).



- Median: biased logFC estimates for spike-in proteins
- maxLFQ: more variable logFC estiamtes for spike-in proteins

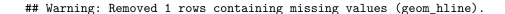
## 2.6.2 Median summarization

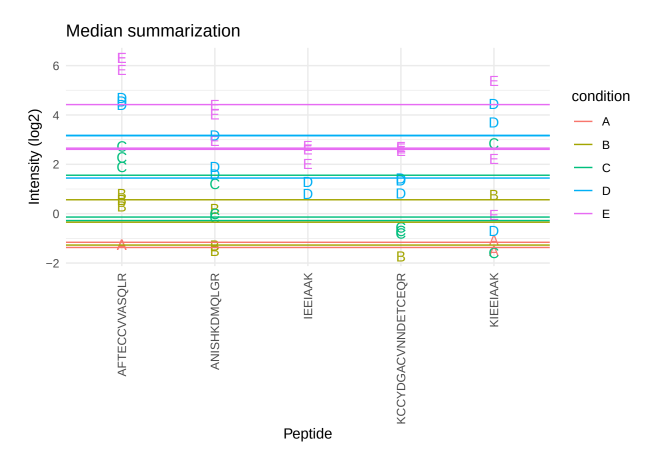
We first evaluate median summarization for protein P01031ups |CO5\_HUMAN\_UPS. Click to see code to make plot

```
dataHlp <- pe[["peptideNorm"]][
    rowData(pe[["peptideNorm"]]) Proteins == prot,
    colData(pe) lab=="lab3"] %>% assay

sumMedian <- data.frame(
    intensity= dataHlp
        %>% colMedians(na.rm=TRUE)
,
    condition= colnames(dataHlp) %>% substr(12,12) %>% as.factor )

sumMedianPlot <- sumPlot +
    geom_hline(
    data = sumMedian,
    mapping = aes(yintercept=intensity,color=condition)) +
    ggtitle("Median summarization")</pre>
```





- The sample medians are not a good estimate for the protein expression value.
- Indeed, they do not account for differences in peptide effects
- Peptides that ionize poorly are also picked up in samples with high spike-in concentration and not in samples with low spike-in concentration
- This introduces a bias.

#### 2.6.3 Mean summarization

$$y_{ip} = \beta_i^{\text{sample}} + \epsilon_{ip}$$

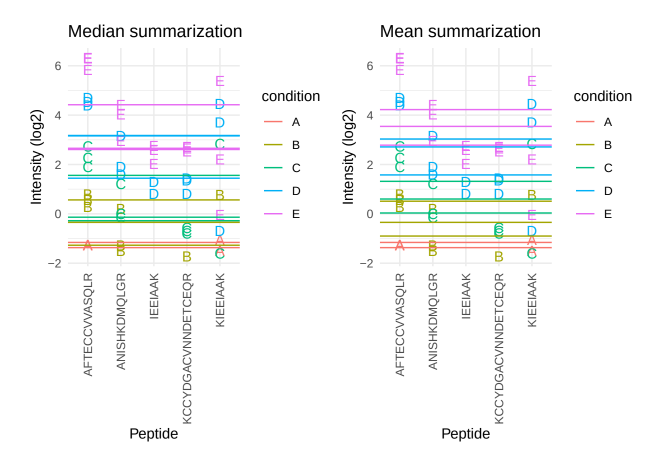
```
sumMeanMod <- lm(intensity ~ -1 + sample,data)

sumMean <- data.frame(
  intensity=sumMeanMod$coef[grep("sample",names(sumMeanMod$coef))],
  condition= names(sumMeanMod$coef)[grep("sample",names(sumMeanMod$coef))] %>% substr(18,18) %>% as.fac
```

```
sumMeanPlot <- sumPlot + geom_hline(
  data = sumMean,
  mapping = aes(yintercept=intensity,color=condition)) +
  ggtitle("Mean summarization")</pre>
```

```
grid.arrange(sumMedianPlot, sumMeanPlot, ncol=2)
```

## Warning: Removed 1 rows containing missing values (geom\_hline).



#### 2.6.4 Model based summarization

We can use a linear peptide-level model to estimate the protein expression value while correcting for the peptide effect, i.e.

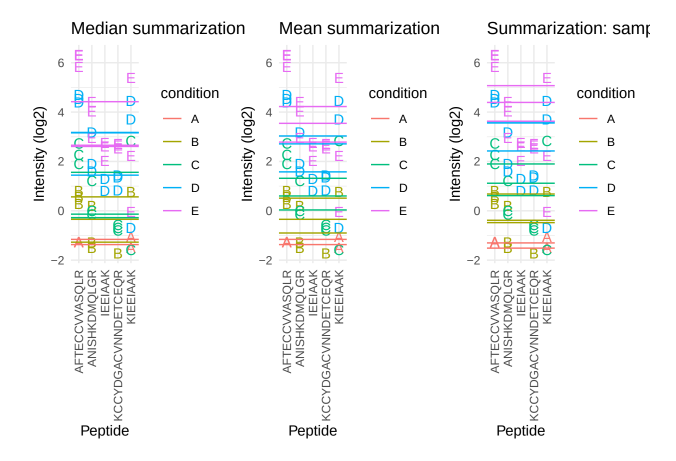
$$y_{ip} = \beta_i^{\text{sample}} + \beta_p^{peptide} + \epsilon_{ip}$$

```
sumMeanPepMod <- lm(intensity ~ -1 + sample + peptide,data)
sumMeanPep <- data.frame(
  intensity=sumMeanPepMod$coef[grep("sample",names(sumMeanPepMod$coef))] + mean(data$intensity) - mean(
  condition= names(sumMeanPepMod$coef)[grep("sample",names(sumMeanPepMod$coef))] %>% substr(18,18) %>%
```

```
fitLmPlot <- sumPlot + geom_line(
   data = data %>% mutate(fit=sumMeanPepMod$fitted.values),
   mapping = aes(x=peptide, y=fit,color=condition, group=sample)) +
   ggtitle("fit: ~ sample + peptide")
sumLmPlot <- sumPlot + geom_hline(
   data = sumMeanPep,
   mapping = aes(yintercept=intensity,color=condition)) +
   ggtitle("Summarization: sample effect")</pre>
```

```
grid.arrange(sumMedianPlot, sumMeanPlot, sumLmPlot, nrow=1)
```

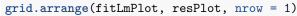
## Warning: Removed 1 rows containing missing values (geom\_hline).

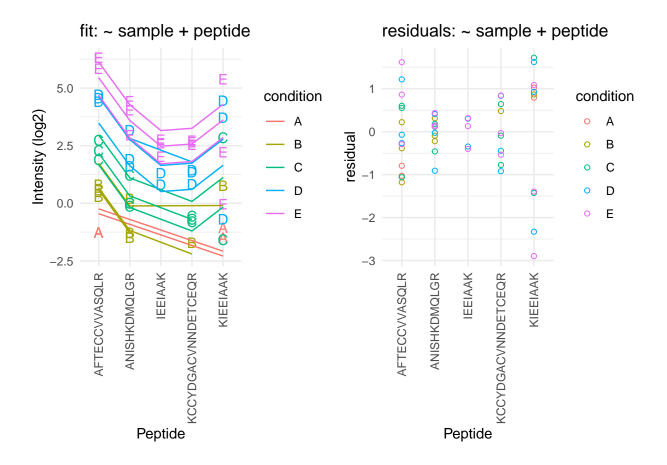


- By correcting for the peptide species the protein expression values are much better separated an better reflect differences in abundance induced by the spike-in condition.
- Indeed, it shows that median and mean summarization that do not account for the peptide effect indeed overestimate the protein expression value in the small spike-in conditions and underestimate that in the large spike-in conditions.
- Still there seem to be some issues with samples that for which the expression values are not well separated according to the spike-in condition.

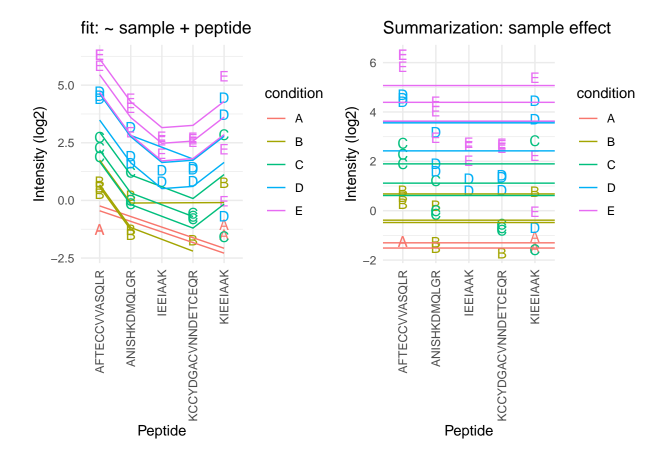
A residual analysis clearly indicates potential issues:

```
resPlot <- data %>%
  mutate(res=sumMeanPepMod$residuals) %>%
  ggplot(aes(x = peptide, y = res, color = condition, label = condition), show.legend = FALSE) +
  geom_point(shape=21) +
  theme_minimal() +
  theme(axis.text.x = element_text(angle = 90, vjust = 0.5, hjust = 1)) +
  xlab("Peptide") +
  ylab("residual") +
  ggtitle("residuals: ~ sample + peptide")
```





```
grid.arrange(fitLmPlot, sumLmPlot, nrow = 1)
```



- The residual plot shows some large outliers for peptide KIEEIAAK.
- Indeed, in the original plot the intensities for this peptide do not seem to line up very well with the concentration.
- This induces a bias in the summarization for some of the samples (e.g. for D and E)

## 2.6.5 Robust summarization using a peptide-level linear model

$$y_{ip} = \beta_i^{\text{sample}} + \beta_p^{peptide} + \epsilon_{ip}$$

• Ordinary least squares: estimate  $\beta$  that minimizes

$$OLS: \sum_{i,p} \epsilon_{ip}^2 = \sum_{i,p} (y_{ip} - \beta_i^{\text{sample}} - \beta_p^{\text{peptide}})^2$$

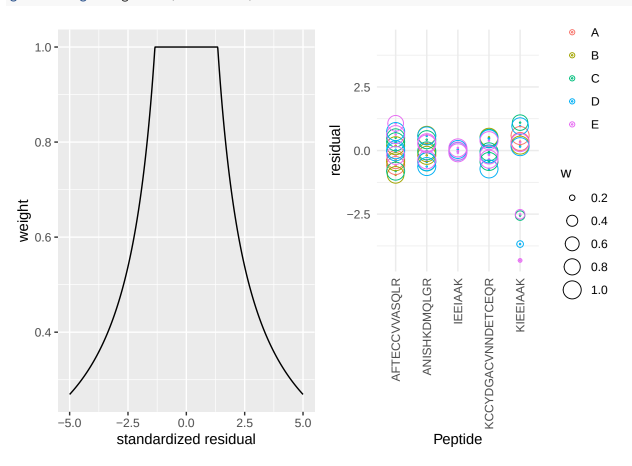
We replace OLS by M-estimation with loss function

$$\sum_{i,p} w_{ip} \epsilon_{ip}^2 = \sum_{i,p} w_{ip} (y_{ip} - \beta_i^{\text{sample}} - \beta_p^{\text{peptide}})^2$$

- Iteratively fit model with observation weights  $w_{ip}$  until convergence
- The weights are calculated based on standardized residuals

```
sumMeanPepRobMod <- MASS::rlm(intensity ~ -1 + sample + peptide,data)</pre>
resRobPlot <- data %>%
  mutate(res = sumMeanPepRobMod$residuals,
         w = sumMeanPepRobMod$w) %>%
  ggplot(aes(x = peptide, y = res, color = condition, label = condition, size=w), show.legend = FALSE) +
  geom_point(shape=21,size=.2) +
  geom_point(shape=21) +
  theme minimal() +
  theme(axis.text.x = element_text(angle = 90, vjust = 0.5, hjust = 1)) +
  xlab("Peptide") +
  ylab("residual") +
  ylim(c(-1,1)*max(abs(sumMeanPepRobMod$residuals)))
weightPlot <- qplot(</pre>
  seq(-5,5,.01),
  MASS::psi.huber(seq(-5,5,.01)),
  geom="path") +
  xlab("standardized residual") +
  ylab("weight")
```

## grid.arrange(weightPlot,resRobPlot,nrow=1)

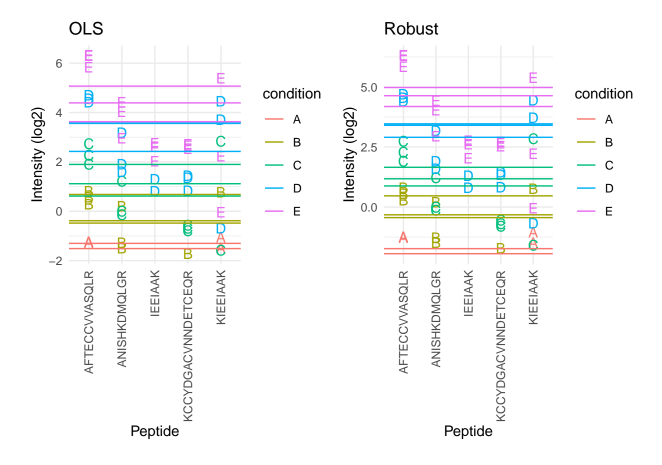


• We clearly see that the weights in the M-estimation procedure will down-weight errors associated with outliers for peptide KIEEIAAK.

```
sumMeanPepRob <- data.frame(
  intensity=sumMeanPepRobMod$coef[grep("sample",names(sumMeanPepRobMod$coef))] + mean(data$intensity) -
  condition= names(sumMeanPepRobMod$coef)[grep("sample",names(sumMeanPepRobMod$coef))] %>% substr(18,18

sumRlmPlot <- sumPlot + geom_hline(
  data=sumMeanPepRob,
  mapping=aes(yintercept=intensity,color=condition)) +
  ggtitle("Robust")</pre>
```

```
grid.arrange(sumLmPlot + ggtitle("OLS"), sumRlmPlot, nrow = 1)
```



• Robust regresion results in a better separation between the protein expression values for the different samples according to their spike-in concentration.

# 2.6.6 Comparison summarization methods

• maxLFQ

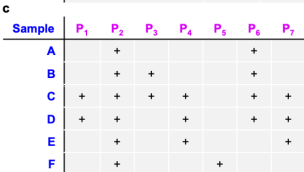
# **a** >P63208

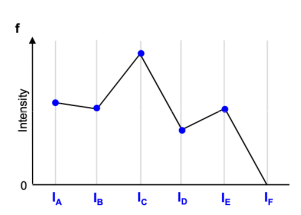
MPSIKLQSSDGEIFEVDVEIAKQSVTIKTMLEDLGMDDEGDD
DPVPLPNVNAAILKKVIQWCTHHKDDPPPPEDDENKEKRTDD
IPVWDQEFLKVDQGTLFELILAANYLDIKGLLDVTCKTVANM
IKGKTPEEIRKTFNIKNDFTEEEEAQVRKENQWCEEK

b **Peptide** Sequence Charge Mod. species 2 LQSSDGEIFEVDVEIAK P<sub>2</sub> 3 LQSSDGEIFEVDVEIAK 2  $P_3$ RTDDIPVWDQEFLK  $P_4$ TVANMIK 2  $P_5$ 2 TVANMIK Oxid. 3 **TPEEIRK** 2

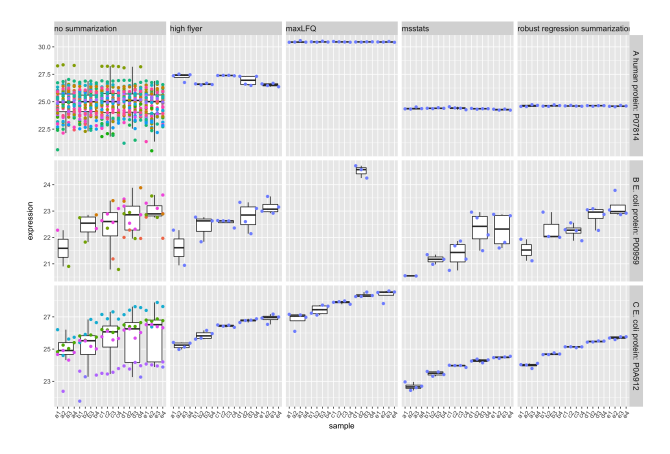
	A	В	C	D	E	F
F	r <sub>FA</sub>	r <sub>FB</sub>	r <sub>FC</sub>	r <sub>FD</sub>	r <sub>FE</sub>	
E	r <sub>EA</sub>	r <sub>EB</sub>	r <sub>EC</sub>	r <sub>ED</sub>		
D	r <sub>DA</sub>	r <sub>DB</sub>	r <sub>DC</sub>			
С	r <sub>CA</sub>	r <sub>CB</sub>				
В	r <sub>BA</sub>					
Α						
d						

•		
$r_{BA} = I_B / I_A$	$r_{CA} = I_C / I_A$	$r_{CB} = I_C / I_B$
$r_{DA} = I_D / I_A$	$r_{DB} = I_D / I_B$	$r_{DC} = I_D / I_C$
$r_{EC} = I_E / I_C$	$r_{ED} = I_E / I_D$	<i>I<sub>F</sub></i> = 0





- MS-stats also uses a robust peptide level model to perform the summarization, however, they typically first impute missing values
- Proteus high-flyer method: mean of three peptides with highest intensity



- (Sticker et al. 2020)
- doi: https://doi.org/10.1074/mcp.RA119.001624
- pdf

# References

Sticker, A., L. Goeminne, L. Martens, and L. Clement. 2020. "Robust Summarization and Inference in Proteome-wide Label-free Quantification." *Mol Cell Proteomics* 19 (7): 1209–19.