# Mass spectrometry-based proteomics

## Laurent Gatto

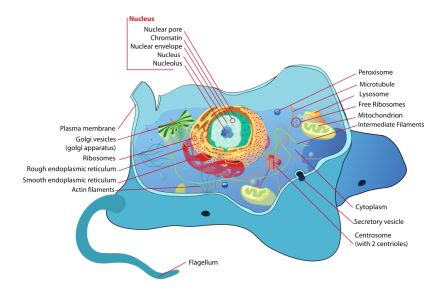
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## Biological information flow

There are three main biological entities that respectively store information, act as data intermediates, and the functional units, and the figure below show how information flows between these three levels.

**DNA**, that lives in the nucleus of cells, is the central information storage mechanism, and encodes the blueprint of the functional units as genes. DNA is **transcribed** into **messenger RNA** (mRNA), that relocalises outside the nucleus and is further processed into its mature *exon*-only form after removal of the non-coding *introns* sequences. Finally, the mRNA is translated by the ribosomial machinery into **proteins** directly into the endoplasmic reticulum (ER) where they are then redirected to their final destination.



In addition to the standard information worflow where DNA is transcribed into RNA that itself is translated into proteins, information flow, there is also reverse transcription, that generates (complementary) DNA from and RNA molecule, as well as replication of

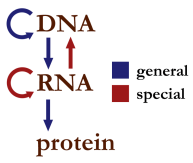


Figure 1: Information flow in biological systems (Source \*Central dogma of biology\* on Wikipedia).

Figure 2: Sub-cellular structure of an animal cell (Source \*Cell biology\* on Wikipedia).

DNA (during cell division) and RNA molecules.

# Why studying proteins

Proteins as the functional units in all living organisms, and they are highly dynamic. The caterpillar and the resulting butterfly have the same genome. The complement of all the expressed proteins, termed the proteome is however very different.



Figure 3: The metamorphosis from a caterpilar to a monarch butterfly. (Image from Phys.prg)

There are different modalities of the proteome that are of interest. In addition to the presence or amount of protein in a biological samples, it is also important to study the interactions between proteins forming protein-protein complexes, the presence of posttranscriptional modification (such as, for example, phosphorylations), the rate at which proteins are produced and degrated, or where the proteins reside inside a cell.

The technique of choice to study proteins in a high throughput way is mass spectrometry.

## Setup

We are going to use the Bioconductor (Huber et al. 2015) MSnbase package (L. Gatto and Lilley 2012), which can be install with the BiocManager package, available from CRAN. If BiocManager isn't available on your computer, install it with:

#### install.packages("BiocManager")

Now, install MSnbase and its dependencies with

BiocManager::install("MSnbase")

For additional information on how to analyse mass spectrometrybased proteomics data, refer to (Gatto and Christoforou 2014) and (L. Gatto 2019), or explore the the proteomics- and mass spectrometryrelated packages on the Bioconductor page



Figure 4: The 'MSnbase' package.

# *How does mass spectrometry work?*

Mass spectrometry (MS) is a technology that separates charged molecules (ions) based on their mass to charge ratio (M/Z). It is often coupled to chromatography (liquid LC, but can also be gas-based GC). The time an analytes takes to elute from the chromatography column is the retention time.

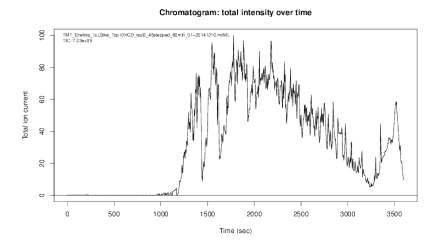


Figure 5: A chromatogram, illustrating the total amount of analytes over the retention time.

An mass spectrometer is composed of three components:

- 1. The source, that ionises the molecules: examples are Matrixassisted laser desorption/ionisation (MALDI) or electrospray ionisation. (ESI)
- 2. The analyser, that separates the ions: Time of flight (TOF) or Orbi-
- 3. The *detector* that quantifies the ions.

When using mass spectrometry for proteomics, the proteins are first digested with a protease such as trypsin. In mass shotgun proteomics, the analytes assayed in the mass spectrometer are peptides.

Often, ions are subjected to more than a single MS round. After a first round of separation, the peaks in the spectra, called MS1 spectra, represent peptides. At this stage, the only information we possess about these peptides are their retention time and their mass-to-charge (we can also infer their charge be inspecting their isotopic envelope, i.e the peaks of the individual isotopes, see below), which is not enough to infer their identify (i.e. their sequence).

In MSMS (or MS2), the settings of the mass spectrometer are set automatically to select a certain number of MS1 peaks (for example 20). Once a narrow M/Z range has been selected (corresponding to one high-intensity peak, a peptide, and some background noise), it is fragmented (using for example collision-induced dissociation (CID), higher energy collisional dissociation (HCD) or electron-transfer dissociation (ETD)). The fragment ions are then themselves separated in the analyser to produce a MS2 spectrum. The unique fragment ion pattern can then be used to infer the peptide sequence using de novo sequencing (when the spectrum is of high enough quality) of using a search engine such as, for example Mascot, MSGF+, ..., that will match the observed, experimental spectrum to theoratical spectra (see details below).

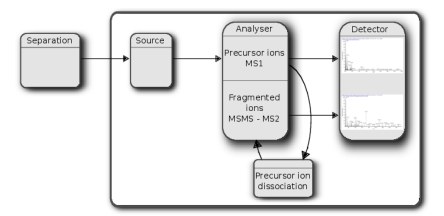


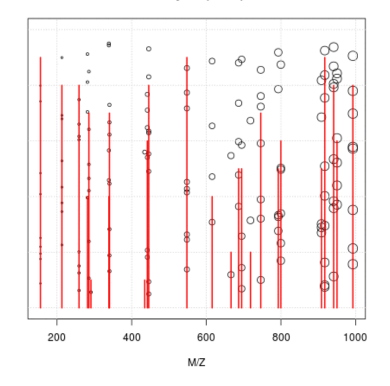
Figure 6: Schematics of a mass spectrometer and two rounds of MS.

The animation below show how 25 ions different ions (i.e. having different M/Z values) are separated throughout the MS analysis and are eventually detected (i.e. quantified). The final frame shows the hypothetical spectrum.

The figures below illustrate the two rounds of MS. The spectrum on the left is an MS1 spectrum acquired after 21 minutes and 3 seconds of elution. 10 peaks, highlited by dotted vertical lines, were selected for MS2 analysis. The peak at M/Z 460.79 (488.8) is highlighted by a red (orange) vertical line on the MS1 spectrum and the fragment spectra are shown on the MS2 spectrum on the top (bottom) right figure.

The figures below represent the 3 dimensions of MS data: a set of spectra (M/Z and intensity) of retention time, as well as the interleaved nature of MS1 and MS2 (and there could be more levels) data.

# Analyser (10/10)



Analytes

Figure 7: Separation and detection of ions in a mass spectrometer.

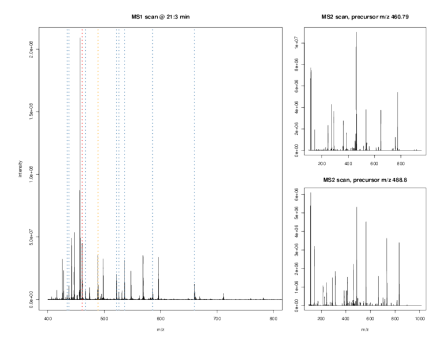
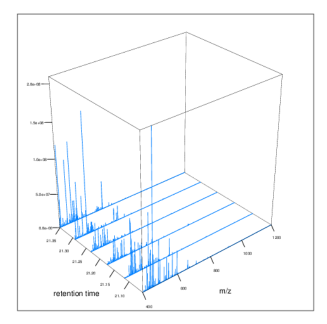
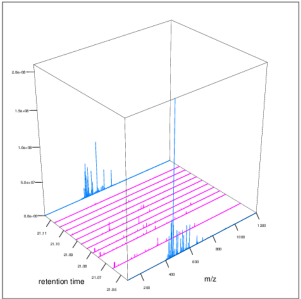


Figure 8: Parent ions in the MS1 spectrum (left) and two sected fragment ions MS2 spectra (right).





Reading and accessing MS data

Let's read a very small raw MS data file into R using the readMSData from the MSnbase package. The file that we are going to load is also available in the package.

1. Load the MSnbase package

```
library("MSnbase")
```

## MS level(s): 2

2. Get the path to the dummyiTRAQ.mzXML file

```
rawf <- dir(system.file(package = "MSnbase", dir = "extdata"),</pre>
    full.name = TRUE, pattern = "mzXML$")
basename(rawf)
## [1] "dummyiTRAQ.mzXML"
3. Read it in using the readMSData function.
x <- readMSData(rawf)</pre>
Х
## MSn experiment data ("MSnExp")
## Object size in memory: 0.18 Mb
## - - - Spectra data - - -
```

Figure 9: MS1 spectra (blue) over retention time (left). MS2 spectra (pink) interleaved between two MS1 spectra (right),

```
## Number of spectra: 5
## MSn retention times: 25:1 - 25:2 minutes
## - - - Processing information - - -
## Data loaded: Mon Mar 25 08:52:13 2019
   MSnbase version: 2.9.3
## - - - Meta data - - -
## phenoData
##
     rowNames: dummyiTRAQ.mzXML
##
    varLabels: sampleNames
    varMetadata: labelDescription
##
## Loaded from:
    dummyiTRAQ.mzXML
##
## protocolData: none
## featureData
##
     featureNames: F1.S1 F1.S2 ... F1.S5
##
       (5 total)
##
     fvarLabels: spectrum
     fvarMetadata: labelDescription
## experimentData: use 'experimentData(object)'
```

The object that is returned by readMSData is of class MSnExp, that can store, access and manipulate raw MS data. Note that here we are focusing on MS-based proteomics data, but this also applied to MS-based metabolomics data.

### class(x)

```
## [1] "MSnExp"
## attr(,"package")
## [1] "MSnbase"
```

4. We can find out how many spectra are available in that data using the function length. Full MS acquisitions would contain hundreds of thousands spectra.

#### length(x)

```
## [1] 5
```

5. We can use various accessor function to get the MS level of these spectra, their retention time, or the M/Z and intensity of the precursor peaks of the ion corresponding to the MS2 spectra.

#### msLevel(x)

```
## F1.S1 F1.S2 F1.S3 F1.S4 F1.S5
      2
            2
                  2
                        2
                              2
##
```

### rtime(x)

F1.S1 F1.S2 F1.S3 F1.S4 ## 1501.35 1501.59 1501.85 1502.07 1502.31

### precursorMz(x)

F1.S1 F1.S2 F1.S3 F1.S4 ## F1.S5 ## 645.3741 546.9586 645.3741 716.3405 437.8040

### precursorIntensity(x)

## F1.S1 F1.S2 F1.S3 F1.S4 F1.S5 ## 47659400 26356100 23432400 24854800 7052960

6. We can also extract individual spectra using [[ and plot them.

### x[[3]]

## Object of class "Spectrum2"

Precursor: 645.3741

Retention time: 25:2

## Charge: 2 ## MSn level: 2

Peaks count: 2125

Total ion count: 150838188

# **plot**(x[[3]])

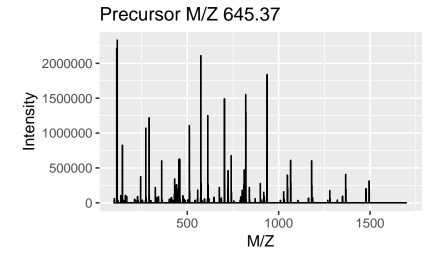


Figure 10: Visualisation of the 3rd MS spectrum in our small test data set.

#### **Exercise**

For the rest of this tutorial, we will be using a slightly larger dataset (still tiny compared to full acquisitions) that is distributed with the MSnbase package. Load it as shown below and compute the number of spectra available in that dataset, their MS level, and the retention time range over which these spectra have been acquired.

```
data(itragdata)
length(itragdata)
## [1] 55
unique(msLevel(itragdata))
## [1] 2
formatRt(range(rtime(itraqdata)))
## [1] "19:9" "50:18"
```

This object also contains additional metadata for each spectrum, that can be accessed, as a data.frame, with fData.

# Identification

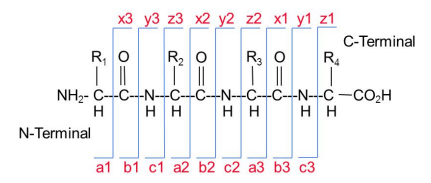
The raw data is still a long way of obtaining biologically relevant proteomics data. The first step to obtain proteomics data is to identify the peptides that have been acquired in the MS. Peptide identification work by comparing expected and observed spectra. As shown below, when a precursor peptide ion is fragmented in a CID cell, it breaks at specific bonds, producing sets of peaks (a, b, c and x, y, z) that can be predicted.

It is thus possible to calculate the expected set of fagment peaks for a given peptide, such as SIGFEGDSIGR below.

### calculateFragments("SIGFEGDSIGR")

##	mz	ion	type	pos	Z	seq
## 1	88.03931	b1	b	1	1	S
## 2	201.12337	b2	b	2	1	SI
## 3	258.14483	b3	b	3	1	SIG
## 4	405.21324	b4	b	4	1	SIGF
## 5	534.25583	b5	b	5	1	SIGFE
## 6	591.27729	b6	b	6	1	SIGFEG
## 7	706.30423	b7	b	7	1	SIGFEGD
## 8	793.33626	b8	b	8	1	SIGFEGDS

Figure 11: Peptide fragmentation.



Biemann, K Methods Enzymol (1990) 193 886-887

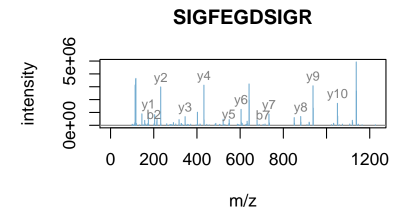
```
## 9
       906.42032
                                      SIGFEGDSI
                      b9
                            b
                                 9 1
       963.44178
                                10 1 SIGFEGDSIG
## 10
                    b10
                            b
       175.11895
                                 1 1
                                                R
## 11
                     у1
                     y2
                                 2 1
## 12
        232.14041
                            У
                                               GR
## 13
       345.22447
                                 3 1
                                             IGR
                     y3
                            У
## 14
       432.25650
                                 4 1
                                            SIGR
                     y4
                            У
                                 5 1
## 15
       547.28344
                     y5
                                           DSIGR
                            У
       604.30490
                     y6
                                 6 1
## 16
                                          GDSIGR
                            У
## 17
       733.34749
                                 7 1
                     y7
                                         EGDSIGR
                            У
## 18
       880.41590
                                 8 1
                     y8
                                        FEGDSIGR
                            У
## 19
       937.43736
                     y9
                            У
                                 9 1
                                     GFEGDSIGR
## 20 1050.52142
                                10 1 IGFEGDSIGR
                    y10
                            У
                                 9 1
## 21
       873.42266
                    b9_
                                       SIGFEGDSI
                           b_{-}
       930.44412 b10_
                                10 1 SIGFEGDSIG
## 22
                           b_{-}
       514.28579
                                 5 1
## 23
                    y5_
                                           DSIGR
                           y_{-}
## 24
       571.30725
                                 6 1
                    y6_
                           y_{-}
                                          GDSIGR
                                 7 1
## 25
       700.34984
                    y7_
                           y_{-}
                                         EGDSIGR
## 26
       847.41825
                    y8_
                                 8 1
                                        FEGDSIGR
                           y_{-}
## 27
       904.43971
                    y9_
                                 9 1
                                      GFEGDSIGR
                           \mathbf{y}_{-}
## 28 1017.52377 y10_
                                10 1 IGFEGDSIGR
                           y_{-}
        142.12130
                                 1 1
   29
                    y1_{-}
                                                R
                           y_{-}
## 30
       199.14276
                    y2_
                           y_{-}
                                 2 1
                                               GR
       312.22682
                                 3 1
                                              IGR
## 31
                    y3_
                           y_{-}
       399.25885
## 32
                   y4_
                           y_{-}
                                 4 1
                                            SIGR
```

The last step is to compare obseved and expected peaks. If there is a good match, the MS2 spectrum is assigned the peptide sequence.

```
itraqdata2 <- pickPeaks(itraqdata, verbose = FALSE)</pre>
```

```
s <- "SIGFEGDSIGR"
plot(itraqdata2[[14]], s, main = s)
```

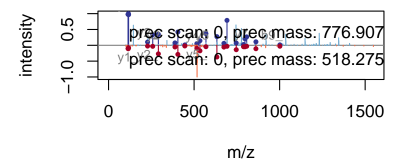
Figure 12: Matching observed and expected peaks.



It is also possible to plot 2 spectra to compare them directly.

```
plot(itraqdata2[[25]], itraqdata2[[28]], sequences = rep("IMIDLDGTENK",
    2))
```

Figure 13: Direct comparison of 2 MS2 spectra.



In a full experiment, all possible peptides from the known (or relevant) proteome of interest (such as databases that can be downloaded from the UniProt site1) are compared to the millions of observed

<sup>&</sup>lt;sup>1</sup> The Universal Protein Resource (UniProt) is a freely and accessible comprehensive resource for protein sequence and annotation data.

spectra.

From the list of identified petides, it is then necessary to infer the most propable proteins that were present in the biological sample.

### **Exercise**

Plot the 44th spectrum of the itragdata2 experiment. The sequence can be accessed in the feature metadata with

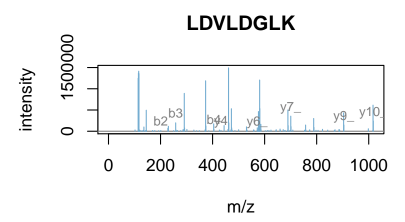
Statistical challenge The matching between millions of observed and possible spectra causes real challenges due to the large search space and the risk of false positives. See [@Kall:2008] for futher reading.

```
fData(itraqdata2)$PeptideSequence[[44]]
```

```
## [1] LDVLDGLK
```

## 47 Levels: AADALLLK ... VWVVEGSK

plot(itraqdata2[[44]], s, main = fData(itraqdata2)\$PeptideSequence[[44]])



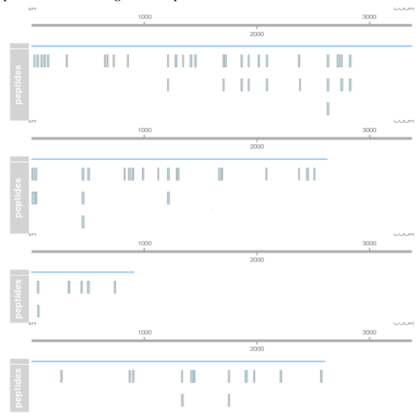
## Quantitation

The last step of MS data processing is to quantify peptide abundances in the biological samples. The table below summarises the different possibilites depending whether the proteins or peptides are labelled, and whether the quantitation is performed in MS1 or MS2.

	Label-free	Labelled
MS <sub>1</sub>	XIC	SILAC, 15N
MS2	Counting	iTRAQ, TMT

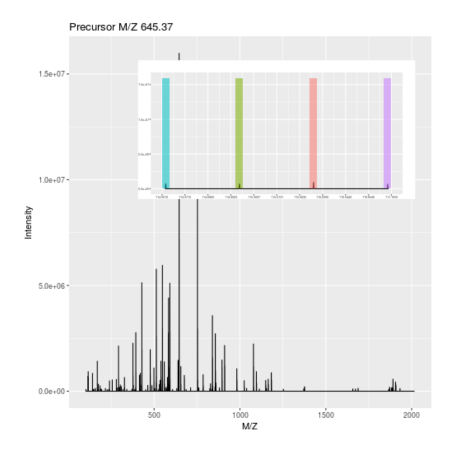
### Label-free MS2: Spectral counting

In spectral counting, on simply counts the number of quantified peptides that are assigned to a protein.



### Labelled MS2: Isobaric tagging

Isobaric tagging refers to the labelling using isobaric tags, i.e. chemical tags that have the same mass and hence can't be distinguish by the spectrometer. The peptides of different samples (4, 6, 10 or 11) are labelled with different tags and combined prior to mass spectrometry acquisition. Given that they are isobaric, all identical peptides, irrespective of the tag and this the sample of origin, are co-analysed, up to fragmentation prior to MS2 analysis. During fragmentation, the isobaric tags fall of, fragment themselves, and result in a set of sample specific peaks. These specific peaks can be used to infer samplespecific quantitation, while the rest of the MS2 spectrum is used for identification.



Label-free MS1: extracted ion chromatograms

In label-free quantitation, the precursor peaks that match an identified peptide are integrated of retention time and the area under that extracted ion chromatogram is used to quantify that peptide in that sample.

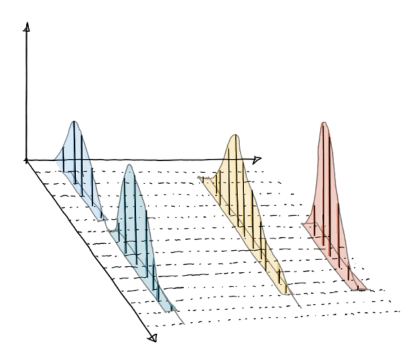


Figure: credit Johannes Rainer.

# Labelled MS1: SILAC

In SILAc quantitation, sample are grown in a medium that contains heavy amino acids (typically arginine and lysine). All proteins gown in this heavy growth medium contain the heavy form of these amino acids. Two samples, one grown in heavy medium, and one grown in normal (light) medium are then combined and analysed together. The heavy peptides precursor peaks are systematically shifted compared to the light ones, and the ratio between the height of a heavy and light peaks can be used to calculate peptide and protein foldchanges.

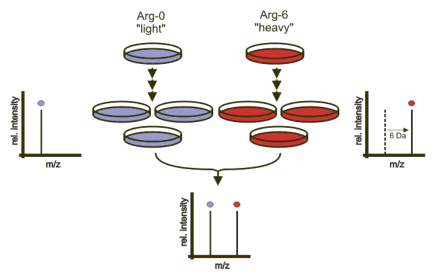


Figure: credit Wikimedia Commons.

### Exercise

As its name implies, the itraqdata is an iTRAQ-based isobar quantitation experiment. We can visualise the reporter peaks as follows:

plot(itraqdata[[14]], reporters = iTRAQ4, full = TRUE)

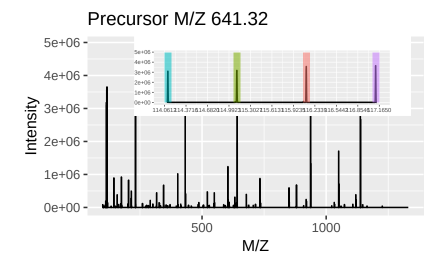


Figure 14: Visualisation of the iTRAQ reporter peaks.

We can quantify these four peaks with the quantify method, to produce and object of class MSnSet containing quantitation data. The quantitation values can be accessed with exprs. This data also contains feature metadata that can be accessed with the fData function.

```
msnset <- quantify(itragdata, method = "trap",</pre>
    reporters = iTRAQ4)
msnset
## MSnSet (storageMode: lockedEnvironment)
## assayData: 55 features, 4 samples
    element names: exprs
## protocolData: none
## phenoData
    sampleNames: iTRAQ4.114 iTRAQ4.115
##
       iTRAQ4.116 iTRAQ4.117
##
    varLabels: mz reporters
    varMetadata: labelDescription
##
## featureData
##
    featureNames: X1 X10 ... X9 (55
##
       total)
##
    fvarLabels: spectrum ProteinAccession
##
       ... collision.energy (15 total)
##
     fvarMetadata: labelDescription
## experimentData: use 'experimentData(object)'
## Annotation: No annotation
## - - - Processing information - - -
## Data loaded: Wed May 11 18:54:39 2011
## Updated from version 0.3.0 to 0.3.1 [Fri Jul 8 20:23:25 2016]
## iTRAQ4 quantification by trapezoidation: Mon Mar 25 08:52:15 2019
## MSnbase version: 1.1.22
head(exprs(msnset))
##
       iTRAQ4.114 iTRAQ4.115 iTRAQ4.116
        1347.6158 2247.3097 3927.6931
## X1
## X10
       739.9861 799.3501 712.5983
## X11 27638.3582 33394.0252 32104.2879
## X12 31892.8928 33634.6980 37674.7272
## X13 26143.7542 29677.4781 29089.0593
## X14 6448.0829 6234.1957 6902.8903
##
      iTRAQ4.117
## X1
        7661.1463
## X10
        940,6793
## X11 26628.7278
## X12 37227.7119
## X13 27902.5608
## X14 6437.2303
head(fData(msnset))
```

##		spectrum Pro	oteinAcces	ssion	
##	X1	1		BSA	
##	X10	10	ECA	1422	
##	X11	11	ECA	44030	
##	X12	12	ECA	\3882	
##	X13	13	ECA	A1364	
##	X14	14	ECA	N0871	
##				ProteinD	escription
##	X1		k	oovine ser	rum albumin
##	X10	glucose-1-pl	nosphate d	cytidylylt	ransferase
##	X11	50S	ribosoma	l subunit	protein L4
##	X12		cha	aperone pr	otein DnaK
##	X13	succin	yl-CoA syr	nthetase a	alpha chain
##	X14		NADP-der	oendent ma	alic enzyme
##		PeptideSeque	ence file	[dx retent	ion.time
##	X1	NYO	QEAK	1	1149.31
##	X10	VTLVDTGEHSM	TGGR	1	1503.03
##	X11	SI	PIWR	1	1663.61
##	X12	TAID	DALK	1	1663.86
##	X13	SII	LINK	1	1664.08
##	X14	DFEVVNNES	SDPR	1	1664.32
##		precursor.m	z precurso	or.intensi	ty charge
##	X1	520.783	3	34496	)20 2
##	X10	573.953	9	78494	120 3
##	X11	401.7392	2	412536	500 2
##	X12	567.8339	9	235495	500 2
##	X13	488.3269	9	130252	200 2
##	X14	782.871	5	184050	000 2
##		peaks.count	tio	ionCour	nt ms.level
##	X1	1922	26413754	4 2641375	54 2
##	X10	1376	24482283	1 2448228	31 2
##	X11	1571	231075934	4 23107593	34 2
##	X12	2397	247323187	7 24732318	37 2
##	X13	2574	207247502	2 20724750	)2 2
##	X14	1829	115317275	5 11531727	<sup>7</sup> 5 2
##		acquisition	.number co	ollision.e	energy
##	X1		2		40
##	X10		11		40
##	X11		12		40
##	X12		13		40
##	X13		14		40
##	X14		15		40

# Quantitative data processing

In our examples, we not have processing data for the 55 peptides and 4 samples. In this data, there is only 1 missing value, corresponding to an absent reporter peak. We are going to simply drop that feature.

```
table(is.na(exprs(msnset)))
##
## FALSE TRUE
     219
msnset <- filterNA(msnset)</pre>
```

In MS1 label-free experiments, given that each sample is acquired independently, the proportion of missing values can be as high several tens of percent. In such situations, removing rows with missing values isn't possible at all. Imputation is possible, albeit tricky, as different mechanisms can be responsible for missing value that appear either at random or not at random (Lazar et al. 2016).

Next, we aggregate the spectrum-level quantitation values into protein-level data using the median and the combineFeatures function:

```
prots <- combineFeatures(msnset, fcol = "ProteinAccession",</pre>
   method = "median")
head(exprs(prots))
##
          iTRAQ4.114 iTRAQ4.115 iTRAQ4.116
## BSA
            1347.616 2247.310
                                3927.693
## ECA0172 17593.548 18545.620 19361.837
## ECA0435 4923.628 5557.818 5775.203
## ECA0452 1524.148 1399.897 1547.218
## ECA0469 1069.945 1035.689 1029.420
## ECA0621 1101.062
                      1124.167 1140.093
##
          iTRAQ4.117
## BSA
           7661.1463
## ECA0172 18328.2365
## ECA0435 5079.2952
## ECA0452 1563.2299
## ECA0469
           999.6957
## ECA0621 1191.8055
```

Following on from here, many data processing such as normalisation, non-specific filtering, and hypothesis testing is very similar to other omics data.

# Applications in statistical learning

In this section, I illustrate a use case of mass spectrometry-based proteomics to infer sub-cellular protein localisation and the application of statistical learning. This section requires the pRoloc and pRolocdata packages (Gatto et al. 2014; Breckels et al. 2016). Both packages can be installed with the BiocManager::install function, as illustrated above.

```
library("pRoloc")
library("pRolocdata")
```

The hyperLOPIT2015 data contains localisation data for 5032 proteins in mouse embryonic stem cells (Christoforou et al. 2016). The protein information is collected along a 20 fraction density gradient - our data matrix has dimensions 5032 rows and 20 columns. We use PCA to easily visualise it in 2 dimensions.

```
data(hyperLOPIT2015)
## set colours
setStockcol(paste0(getStockcol(), 80))
## produce PCA plot
plot2D(hyperLOPIT2015)
```

Each dot on the PCA plot represents a single protein. The protein is either of unknown localisation, and represented by a grey circle, or is of known localisation (and called an organelle marker) and is coloured according to its expected sub-cellular localisation.

In the next figure, we have trained an support vector machine (SVM) model and classified proteins of unknown localisation using an SVM model. The size of the dots are scaled according to the classifier score and only assignments corresponding to a false discovery rate of 5% have been considered.

```
sz <- exp(fData(hyperLOPIT2015)$svm.score) - 1</pre>
plot2D(hyperLOPIT2015, fcol = "final.assignment",
    cex = sz)
```

### Session information

```
## R version 3.5.3 Patched (2019-03-11 r76221)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 18.04.2 LTS
## Matrix products: default
```



Figure 15: The 'pRoloc' package.

Figure 16: Mouse stem cell spatial proteomics data from Christoforou et al.

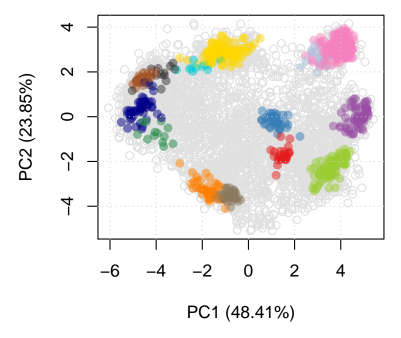
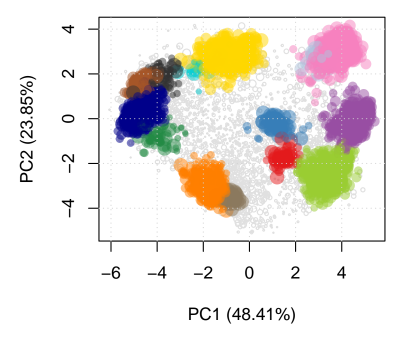


Figure 17: New sub-cellular assignment after using support vector machine classifier.



```
## BLAS: /usr/lib/x86_64-linux-gnu/libf77blas.so.3.10.3
## LAPACK: /usr/lib/x86_64-linux-gnu/atlas/liblapack.so.3.10.3
##
## locale:
  [1] LC_CTYPE=en_US.UTF-8
## [2] LC_NUMERIC=C
## [3] LC_TIME=fr_FR.UTF-8
## [4] LC_COLLATE=en_US.UTF-8
## [5] LC_MONETARY=fr_FR.UTF-8
## [6] LC_MESSAGES=en_US.UTF-8
## [7] LC_PAPER=fr_FR.UTF-8
## [8] LC_NAME=C
## [9] LC_ADDRESS=C
## [10] LC_TELEPHONE=C
## [11] LC_MEASUREMENT=fr_FR.UTF-8
## [12] LC_IDENTIFICATION=C
##
## attached base packages:
## [1] stats4
                parallel stats
                                     graphics
## [5] grDevices utils
                           datasets
                                    methods
## [9] base
##
## other attached packages:
## [1] pRolocdata_1.20.0
## [2] pRoloc_1.22.2
## [3] BiocParallel_1.16.6
## [4] MLInterfaces_1.62.0
## [5] cluster_2.0.7-1
## [6] annotate_1.60.1
## [7] XML_3.98-1.19
## [8] AnnotationDbi_1.44.0
## [9] IRanges_2.16.0
## [10] MSnbase_2.9.3
## [11] ProtGenerics_1.14.0
## [12] S4Vectors_0.20.1
## [13] mzR_2.16.2
## [14] Rcpp_1.0.1
## [15] Biobase_2.42.0
## [16] BiocGenerics_0.28.0
## loaded via a namespace (and not attached):
     [1] plyr_1.8.4
##
     [2] igraph_1.2.4
     [3] lazyeval_0.2.2
##
```

- ## [4] splines\_3.5.3
- ## [5] ggvis\_0.4.4
- [6] crosstalk\_1.0.0 ##
- ## [7] ggplot2\_3.1.0
- ## [8]  $digest_0.6.18$
- [9] foreach\_1.4.4 ##
- [10] htmltools\_0.3.6 ##
- [11] viridis\_0.5.1
- ## [12] gdata\_2.18.0
- ## [13] magrittr\_1.5
- [14]  $memoise_1.1.0$
- [15] doParallel $_1.0.14$ ##
- ## [16] mixtools\_1.1.0
- ## [17] sfsmisc\_1.1-3
- ## [18] limma\_3.38.3
- ## [19] recipes\_0.1.5
- ## [20] gower\_0.2.0
- ## [21] rda\_1.0.2-2.1
- ## [22] lpSolve\_5.6.13
- ## [23] prettyunits\_1.0.2
- ## [24] colorspace\_1.4-1
- ## [25] blob\_1.1.1
- ## [26] xfun\_0.5
- ## [27] dplyr\_0.8.0.1
- ## [28] crayon\_1.3.4
- ## [29] RCurl\_1.95-4.12
- ## [30] hexbin\_1.27.2
- ## [31] genefilter\_1.64.0
- ## [32] impute\_1.56.0
- ## [33] survival\_2.43-3
- ## [34] iterators\_1.0.10
- ## [35] glue\_1.3.1
- ## [36] gtable\_0.2.0
- ## [37] ipred\_0.9-8
- ## [38] zlibbioc\_1.28.0
- ## [39] kernlab\_0.9-27
- ## [40] Rhdf5lib\_1.4.2
- ## [41] prabclus\_2.2-7
- ## [42] DEoptimR\_1.0-8
- ## [43] scales\_1.0.0
- ## [44] vsn\_3.50.0
- ## [45] mvtnorm\_1.0-10
- ## [46] DBI\_1.0.0
- ## [47] viridisLite\_0.3.0

- ## [48] xtable\_1.8-3
- ## [49] progress\_1.2.0
- ## [50] bit\_1.1-14
- ## [51] proxy\_0.4-23
- ## [52] mclust\_5.4.3
- ## [53] preprocessCore\_1.44.0
- ## [54] lava\_1.6.5
- ## [55] prodlim\_2018.04.18
- ## [56] sampling\_2.8
- ## [57] htmlwidgets\_1.3
- ## [58] httr\_1.4.0
- ## [59] threejs\_0.3.1
- ## [60] FNN\_1.1.3
- ## [61] RColorBrewer\_1.1-2
- ## [62] fpc\_2.1-11.1
- ## [63] modeltools\_0.2-22
- ## [64] pkgconfig\_2.0.2
- ## [65] flexmix\_2.3-15
- ## [66] nnet\_7.3-12
- ## [67] caret\_6.0-81
- ## [68] tidyselect\_0.2.5
- ## [69] labeling\_0.3
- ## [70] rlang\_0.3.2
- ## [71] reshape2\_1.4.3
- ## [72] later\_0.8.0
- ## [73] munsell\_0.5.0
- ## [74] mlbench\_2.1-1
- ## [75] tools\_3.5.3
- ## [76] LaplacesDemon\_16.1.1
- ## [77] generics\_0.0.2
- ## [78] RSQLite\_2.1.1
- ## [79] pls\_2.7-0
- ## [80] evaluate\_0.13
- ## [81] stringr\_1.4.0
- ## [82] mzID\_1.20.1
- ## [83] yaml\_2.2.0
- ## [84] tufte\_0.4
- ## [85] ModelMetrics\_1.2.2
- ## [86] knitr\_1.22
- ## [87] bit64\_0.9-7
- ## [88] robustbase\_0.93-4
- ## [89] randomForest\_4.6-14
- ## [90] purrr\_0.3.2
- ## [91] dendextend\_1.10.0

- ## [92] ncdf4\_1.16.1
- ## [93] nlme\_3.1-137
- ## [94] whisker\_0.3-2
- ## [95] mime\_0.6
- ## [96] formatR\_1.6
- ## [97] biomaRt\_2.38.0
- ## [98] compiler\_3.5.3
- ## [99] e1071\_1.7-1
- ## [100] affyio\_1.52.0
- ## [101] tibble\_2.1.1
- ## [102] stringi\_1.4.3
- ## [103] lattice\_0.20-38
- ## [104] trimcluster\_0.1-2.1
- ## [105] Matrix\_1.2-15
- ## [106] gbm\_2.1.5
- ## [107] pillar\_1.3.1
- ## [108] BiocManager\_1.30.4
- ## [109] MALDIquant\_1.19.2
- ## [110] data.table\_1.12.0
- ## [111] bitops\_1.0-6
- ## [112] httpuv\_1.5.0
- ## [113] R6\_2.4.0
- ## [114] pcaMethods\_1.74.0
- ## [115] affy\_1.60.0
- ## [116] hwriter\_1.3.2
- ## [117] promises\_1.0.1
- ## [118] gridExtra\_2.3
- ## [119]  $codetools_0.2-16$
- ## [120] MASS\_7.3-51.1
- ## [121] gtools\_3.8.1
- ## [122] assertthat\_0.2.1
- ## [123] rhdf5\_2.26.2
- ## [124] withr\_2.1.2
- ## [125] diptest\_0.75-7
- ## [126] hms\_0.4.2
- ## [127] timeDate\_3043.102
- ## [128] grid\_3.5.3
- ## [129] rpart\_4.1-13
- ## [130] coda\_0.19-2
- ## [131] class\_7.3-15
- ## [132] rmarkdown\_1.12
- ## [133] segmented\_0.5-3.0
- ## [134] lubridate\_1.7.4
- ## [135] shiny\_1.2.0

#### ## [136] base64enc\_0.1-3

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