# Standardised and reproducible analysis of mass spectrometry-based single-cell proteomics data Replication of the SCoPE2 analysis by Specht et al. 2019

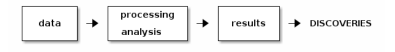
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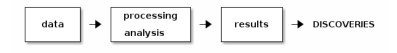
18 August 2020

# Outline

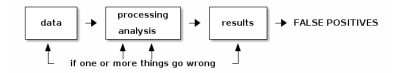
Expectation



Expectation



Reality



▶ **Replication-based development** agreement between the developer, the data producer and the user.

Replication is the first step to define sound data infrastructure and principled analysis.

- SCoPE2 quantifies thousands of proteins x thousands single-cells
- Full protocole available
- ► Full analysis script and data available

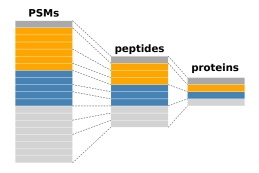
- Contribute a standardised and principled data and analysis that is broadly applicable.
- Open, transparent and reproducible computational infrastructure to further improve data analysis and interpretation.
- ▶ R and Bioconductor (?) offer an ideal environment to attain these goals.

Implemented in the scp package.

# Outline

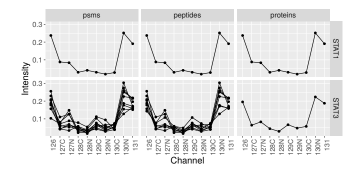
QFeatures: data framework dedicated to manipulate and process MS-based quantitative data.





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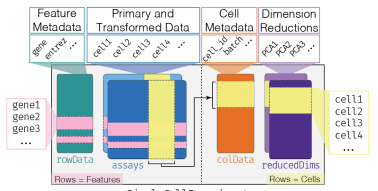


# Data infrastructure: SingleCellExperiment<sup>2,3</sup>

scp package

SingleCellExperiment: provides dedicated framework for single-cell data analysis.



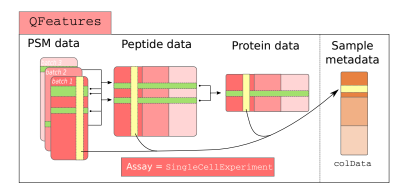


SingleCellExperiment

<sup>&</sup>lt;sup>2</sup>Lun and Risso (2020)

<sup>&</sup>lt;sup>3</sup>Amezquita et al. (2019)

$$scp = SingleCellExperiment + QFeatures$$



AND functions dedicated to processing and analyzing SCP data.

Load data scp package

# Load the SCoPE2 dataset called specht2019v2 4

```
1 library(scpdata)
2 data("specht2019v2")
```

#### Dataset overview

```
show(specht2019v2)
```

```
An instance of class QFeatures containing 179 assays:
[1] 190222S_LCA9_X_FP94AA: SingleCellExperiment with 2823 rows and 11 col...
[2] 190222S_LCA9_X_FP94AB: SingleCellExperiment with 4297 rows and 11 col...
[3] 190222S_LCA9_X_FP94AC: SingleCellExperiment with 4956 rows and 11 col...
[177] 191110S_LCB7_X_APNOV16plex2_Set_9: SingleCellExperiment with 4626 r...
[178] peptides: SingleCellExperiment with 9208 rows and 1018 columns
[179] proteins: SingleCellExperiment with 2772 rows and 1018 columns
```

Tabular data (generated by MaxQuant, ProteomeDiscoverer, ...) can be converted to QFeatures using the readSCP() function.

<sup>&</sup>lt;sup>4</sup>Specht et al. (2019)

Metadata scp package

Sample metadata common to **all assays** are stored in a single table, the colData

Set	Channel	SampleType	lcbatch	sortday	digest
190222S_LCA9_X_FP94AA	RI1	Carrier	LCA9	s8	N
190222S_LCA9_X_FP94AA	RI2	Reference	LCA9	s8	N
190222S_LCA9_X_FP94AA	RI3	Unused	LCA9	s8	N
190222S_LCA9_X_FP94AA	RI4	Macrophage	LCA9	s8	N
190222S_LCA9_X_FP94AA	RI5	Macrophage	LCA9	s8	N
190222S_LCA9_X_FP94AA	RI6	Macrophage	LCA9	s8	N
•••					

1. Load data

#### PSM data

```
[1] 190222S_LCA9_X_FP94AA: SingleCellExperiment with 2823 rows and 11 columns
[2] 190222S_LCA9_X_FP94AB: SingleCellExperiment with 4297 rows and 11 columns
[3] 190222S_LCA9_X_FP94AC: SingleCellExperiment with 4956 rows and 11 columns
...
[177] 191110S_LCB7_X_APNOV16plex2_Set_9: SingleCellExperiment with 4626 rows and 16 columns
```

1. Load data

#### PSM data

```
[1] 190222S_LCA9_X_FP94AR: SingleCellExperiment with 2823 rows and 11 columns
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...
[177] 191110S_LCB7_X_APNOV16plex2_Set_9: SingleCellExperiment with 4626 rows and 16 columns
```

- 2. PSM filtering
- 3. Expression channel by reference channel division
- 4. PSM to peptides aggregation
- Joining sets
- 6. Single cells filtering based on median CV
- 7. Normalization
- 8. Removal of highly missing peptides
- 9. Log-transformation

#### Peptide data

[178] peptides: SingleCellExperiment with 9208 rows and 1018 columns

Load data

#### PSM data

```
[1] 190222S_LCA9_X_FP94AR: SingleCellExperiment with 2823 rows and 11 columns
[2] 190222S_LCA9_X_FP94AC: SingleCellExperiment with 4957 rows and 11 columns
[3] 190222S_LCA9_X_FP94AC: SingleCellExperiment with 4956 rows and 11 columns
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#### Peptide data

[178] peptides: SingleCellExperiment with 9208 rows and 1018 columns

- 9. Peptides to proteins aggregation
- 10. Normalization
- 11. Imputation
- 12. Batch correction

#### Protein data

1. Load data

#### PSM data

```
[1] 190222S_LCA9_X_FP94AR: SingleCellExperiment with 2823 rows and 11 columns
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[178] peptides: SingleCellExperiment with 9208 rows and 1018 columns

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- 12. Batch correction

#### Protein data

# Outline

Filter out features based on the feature metadata

Example: filter out reverse hits. The filter is applied to the Reverse field in the feature metadata

# Data filtering: compute QC metrics

#### Some QC metrics are not compued by MaxQuant:

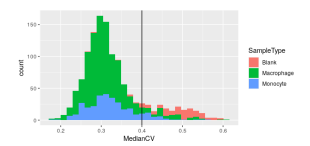
- ► Sample to carrier ratio: discard samples with intensities higher than expected
- ▶ Peptide FDR: expected proportion of features wrongly assigned to a given peptide
- Cell median coefficient of variation: reliability of the protein quantification in a cell.

### Example:

## Source code in scp

# Data filtering: plot QC metrics

## QC metrics are stored in the data set for plotting or subsetting



Feature aggregation = combine features into a higher-level structure.



Example: aggregate peptides to proteins

- Combine the quantitative data from multiple peptides to a single protein
- Store the relationship between the protein and the aggregated peptides

O's can be either **biological** or **technical** zero. They are better relaced by NA's.

Remove highly-missing features (e.g. >= 99 %)

## Impute missing data

Common data transformation can easily be applied such as **log-transformation** or **normalization**.

Example: *log*<sub>2</sub>-transformation:

**Custom function** can be applied to the data set, for example batch correction using 'ComBat'. Three-step procedure:

1. Extract the assay to process

```
1 x <- specht2019v2[["proteins"]]
```

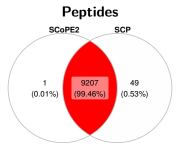
2. Apply the custom function

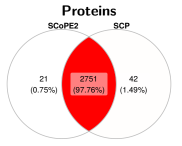
```
2 assay(x) <- ComBat(assay(x), ...)
```

3. Add the processed assay in the dataset

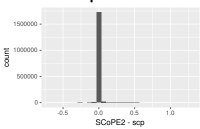
```
3 addAssay(specht2019v2, x, name = "proteins_batch_corrected")
```

# Outline

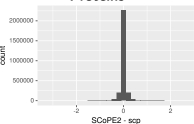








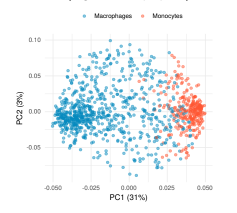
## **Proteins**

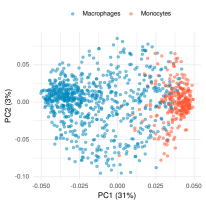


## Weighted PCA on the protein data

## SCoPE2 (Figure 3a in preprint)

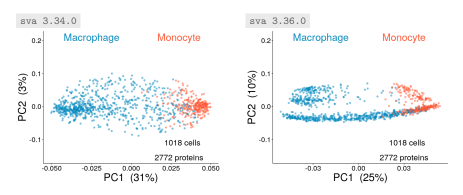
#### scp





Good software development includes continuous maintenance and improvement **BUT** might impact reproducibility

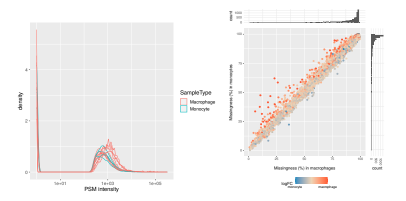
Example: batch correction using 2 versions of the ComBat algorithm (sva package)



Documenting software version is essential for reproducible work

# Outline

- MS-based single cell proteomics: young field, with many challenges and great progess. scp to address the need for principled and reproducible data analysis.
- scp isn't specific to SCoPE2/TMT data, applicable to other LF protocols such as nanoPOTS (Williams et al. (2020); Cong et al. (2020)).
- scp and SingleCellExperiment: same infrastructure for single cell proteomics and RNA sequencing.
- ► Tool for novel computational developments.



#### Resources

- scp: http://UClouvain-CBIO.github.io/scp
- scpdata: coming soon
- QFeatures: http://rformassspectrometry.org
- ► SingleCellExperiment : Bioconductor
- ▶ Slides: http://bit.ly/2020SCP under CC-BY SA

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## Acknowledgements

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- Fonds National de la Recherche Scientifique (FNRS)
- ► Thank you for your attention

# References I