# Standardised and reproducible analysis of mass spectrometry-based single-cell proteomics data

Laurent Gatto, Christophe Vanderaa

Computational Biology Unit (CBIO), de Duve Institute, UCLouvain

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## Outline

#### Introduction

scp package

scp showcase

Replication results

Conclusion

MS-SCP: Mass spectrometry-based single-cell proteomics MS-SCP consist of shotgun proteomics at single-cell level

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

#### BUT

Lack of standardized analysis software

Provide a suite of software package dedicated to MS-SCP that fulfill:

- User-friendly
- Computationaly efficient
- Modularity: integrate other software packages
- Promote reproducibility
- Platform-independent
- Free of charge

R/Bioconductor is an ideal environment

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## Data infrastructure: QFeatures<sup>1</sup>

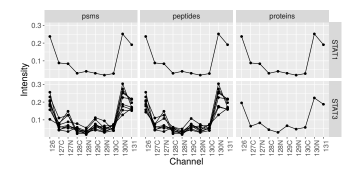
QFeatures: data framework dedicated to manipulate and process MS-based quantitative data.



# **PSMs** peptides proteins

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



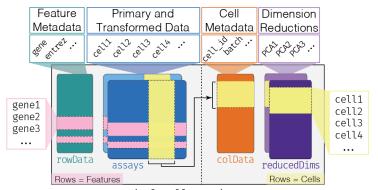


## Data infrastructure: SingleCellExperiment<sup>2,3</sup>

scp package

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

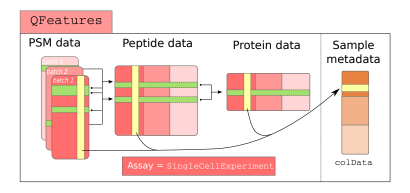




SingleCellExperiment



<sup>&</sup>lt;sup>2</sup>Lun and Risso (2020) <sup>3</sup>Amezquita et al. (2019)



Load data scp package

### Load the SCoPE2 dataset called specht2019v2

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

#### Dataset overview

1 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...
```

Tabular data (such as generated from MaxQuant, ProteomeDiscoverer, ...) can be read using the readSCP() function.

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

Metadata scp package

colData stores sample metadata for all assays in one table

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	<b>s</b> 8	N

## Analysis workflow

#### 1. Load data

#### PSM data

```
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```

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#### PSM data

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

#### Peptide data

[178] peptides: SingleCellExperiment with 9208 rows and 1018 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

#### Peptide data

1. Load data

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#### Peptide data

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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

## QC metrics (1)

Interesting metrics for MS-SCP quality control:

- ► Sample to carrier ratio: ratio of the carrier channel intensity signal over the sample channel intensity
- ▶ Peptide FDR<sup>4</sup>: expected rate of wrongly assigned features to a given peptide
- ► Cell median CV<sup>5</sup>: reliability of the protein quantification summarized over each cell.

#### Example:

#### Source code in scp

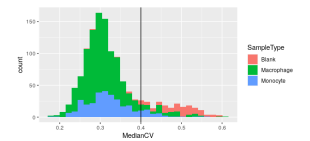


<sup>&</sup>lt;sup>4</sup>false discovery rate

<sup>&</sup>lt;sup>5</sup>coefficient of variation

## QC metrics (2)

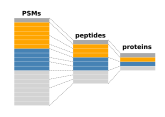
#### QC metrics are stored in the data set for plotting or subsetting



## Feature aggregation

#### Feature aggregation includes 2 steps:

- Combine the quantitative data from multiple features to a single aggregated features
- Store the relationship between the parent features and the aggregated features



#### Example: aggregate peptides to proteins

 ${\color{red}0}$  's can be either **biological** or **technical** zero. They are better relaced by  ${\color{red}NA}$  's.

Features containing too many missing data (e.g. >= 99 %) should be removed

#### Common data transformation can easily be applied:

- Normalization
- ► Log-transformation
- Imputation

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

Some custom function can be applied to the data set too. Example: batch correction using sva::ComBat. First, extract the data

```
1 sce <- specht2019v2[["proteins"]]</pre>
```

## Build the correction matrix and apply the ComBat algorithm

## Add the corrected protein to the dataset and keep feature relationships

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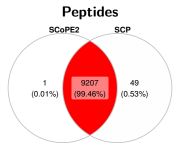
Introduction

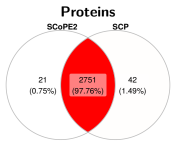
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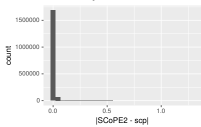
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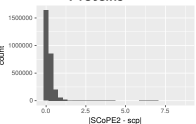




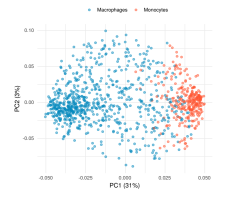




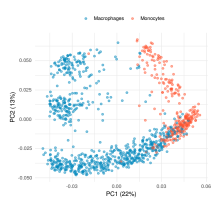
#### **Proteins**



#### SCoPE2



#### scp



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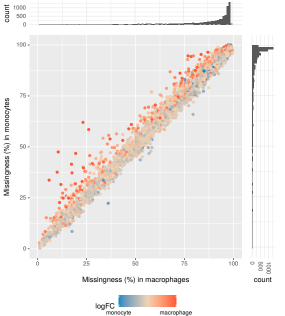
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Missingness Conclusion





- scp package suite provides a standardized environment for performing MS-SCP data analysis
- Flexibly reproduce existing analyses from different groups or protocoles (multiplex vs label free)

#### Advantages:

- ► Allow automation of the analysis
- Facilitate new computational developments
- Promotes reproducibility
- Increases field visibility
- Include other modalities: scRNA-Seq, ATAC-Seq, etc

Resources Conclusion

#### **Packages**

- ▶ scp: GitHub repository UClouvain-CBIO/scp
- scpdata: coming soon
- ► QFeatures: GitHub repository rformassspectrometry/QFeatures
- ► SingleCellExperiment : Bioconductor

#### SCoPE2 reproduction vignette

Available at...

#### Slides and source code

Available at...

Acknowledgements

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#### References I

Robert A Amezquita, Aaron T L Lun, Etienne Becht, Vince J Carey, Lindsay N Carpp, Ludwig Geistlinger, Federico Martini, Kevin Rue-Albrecht, Davide Risso, Charlotte Soneson, Levi Waldron, Hervé Pagès, Mike L Smith, Wolfgang Huber, Martin Morgan, Raphael Gottardo, and Stephanie C Hicks. Orchestrating single-cell analysis with bioconductor. *Nat. Methods*, pages 1–9, December 2019.

Laurent Gatto. QFeatures: Quantitative features for mass spectrometry data, 2020. URL https://github.com/RforMassSpectrometry/QFeatures. R package version 0.7.0.

Aaron Lun and Davide Risso. SingleCellExperiment: S4 Classes for Single Cell Data, 2020. R package version 1.10.1.