

How to analyse single-cell proteomics data and
focus on the underlying biology?
Your results are only as good as your method and
software.

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Mass spectrometry-based single-cell proteomics (SCP) has become a credible player in the single-cell omics arena thanks to substantial technical improvements that have pushed the boundaries of sensitivity and throughput. But what should one do once the precious data have been acquired, often at great cost? Reviewing the SCP literature doesn't provide much help, as every lab tends to run their own in-house, either overly complex or unrealistically trivial and undocumented analysis pipeline. When facing complex data, best is to start with simpler but principled analyses approaches, such as the sciplainer method. The goal of sciplainer is to move the tension point from how to process SCP data to explain it in the light of the biological question. In this talk, I will use SCP to illustrate how to approach, as a bioinformatician, complex data and its underlying biological complexity, emphasising the role of research software engineering and computational science.

Slides: <https://lgatto.github.io/pub/2025CompSysBio.pdf>

Your results are only as good as your method and software.

My analysis is only as good as the explanation and the software to go with it.

Prof Susan Holmes

Is a computational researcher coding doing research?

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Better Software, Better Research

Software Sustainability Institute

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What is good software? What is data analysis?

*Your results are only as good as your method, software
and your users.*

Outline

Methods, software and users

Single-cell proteomics: introduction

SCP data/analysis - round 1

Computational challenges

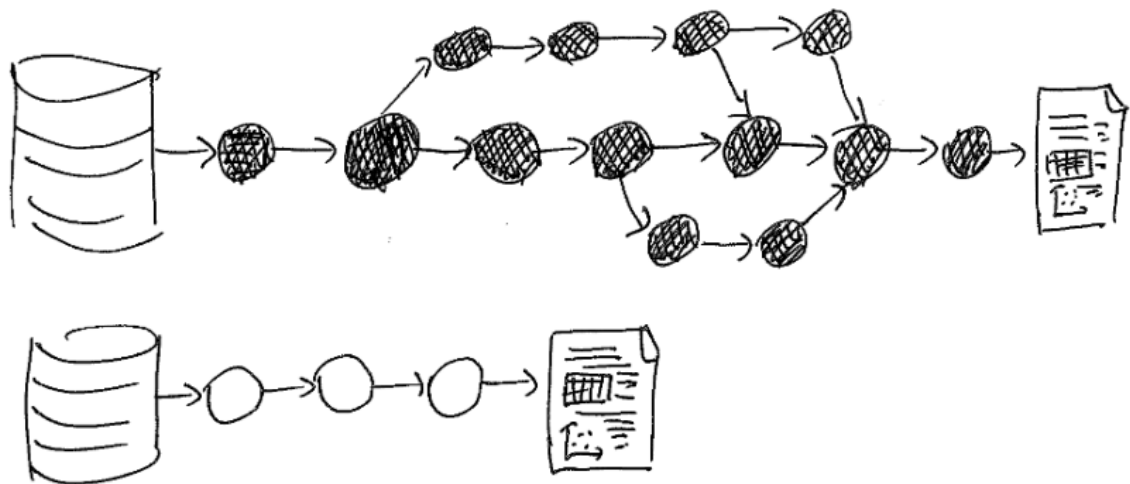
A principled approach to SCP data analysis - round 2

Implementation - `scp` and `scplainer`

Conclusions

What is a good data analysis?

Simpler is better



- ▶ Data analysis should be as simple as possible, but no simpler.
- ▶ Data analysis should be as complex as needed, but not more complex.

Software for data analysis

Software for data analysis

- ▶ Compose simple pipelines when possible
- ▶ Compose more complex pipelines when necessary
- ▶ Enable transparency and reproducibility

Users!

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1. knowledgeable in MS-based (single-cell) proteomics
2. has basic knowledge of data analysis
3. some R (or Python, ...) experience

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Single-cell technologies unravel cellular heterogeneity

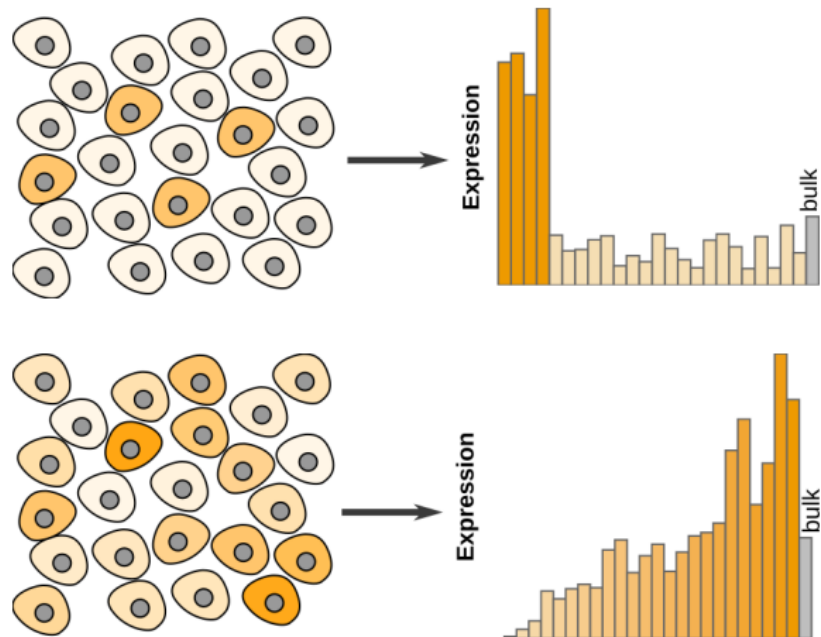


Figure: Cell types and cell states, subpopulation identification, differentiation trajectories (in the absence of known markers).

Single-cell technologies

	FC	scRNA-Seq	SCP
features	10	10^4	10^3
cells	10^6	10^4	10^3
samples	10 - 100	1 - 10	1 ...
	sample/cell throughput	feature throughput	functional

Single-cell proteomics

	FC	scRNA-Seq	SCP
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samples	10 - 100	1 - 10	1 ...
	sample/cell throughput	feature throughput	functional

- ▶ RNA → intention vs. Protein → action
- ▶ Inference of direct regulatory interactions with minimal assumptions ([Slavov, 2022](#); [Hu et al., 2023](#)).
- ▶ Post-translational modifications

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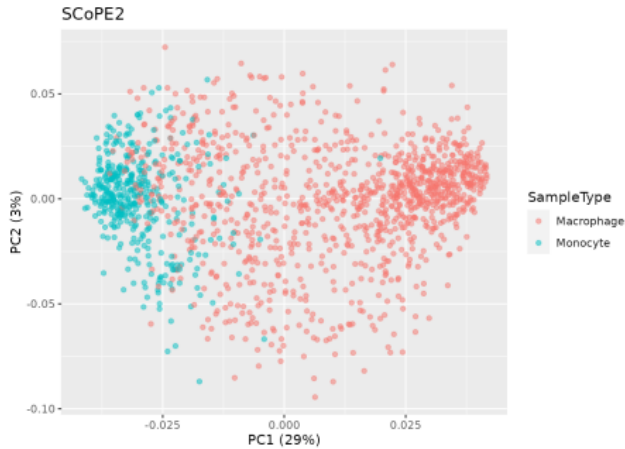
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Material (1)

The SCoPE2 dataset

- ▶ Seminal dataset published by [Specht et al. \(2021\)](#)
- ▶ 1096 macrophages, 394 monocytes (after QC)
- ▶ 9354 peptides, 3042 proteins
- ▶ **Pre-print, data and code available since 2019**



Methods

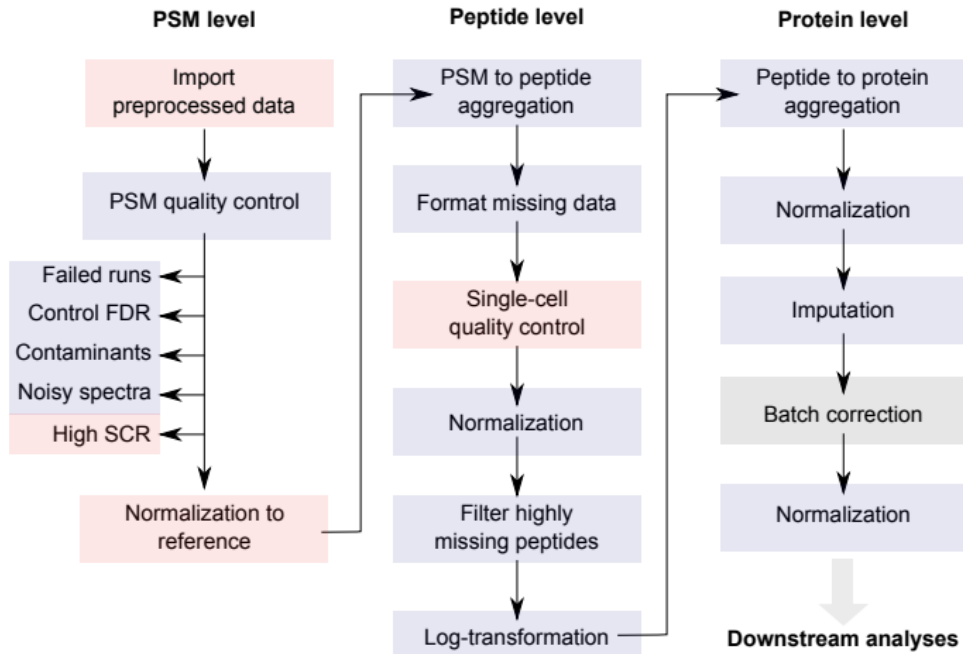


Figure: Overview of the key steps performed in the SCoPE2 pipeline (Vanderaa and Gatto, 2021). Blue boxes: QFeatures. Red boxes: scp. Gray box: sva::ComBat.

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Challenge 1: batch effects

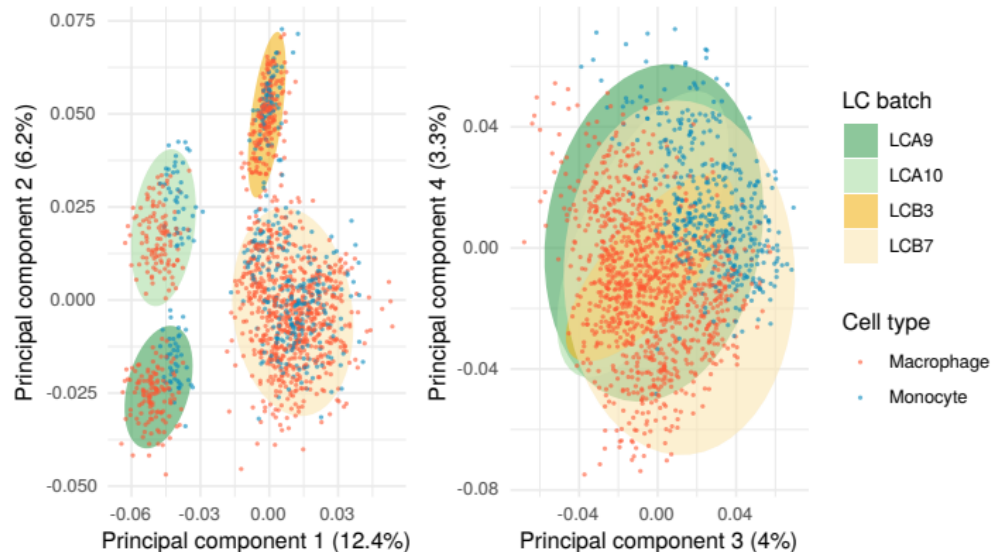


Figure: PCA for the first four components. Each point represents a single-cell and is colored according to the corresponding cell type ([Vanderaa and Gatto, 2021](#)).

Challenge 2: missing data

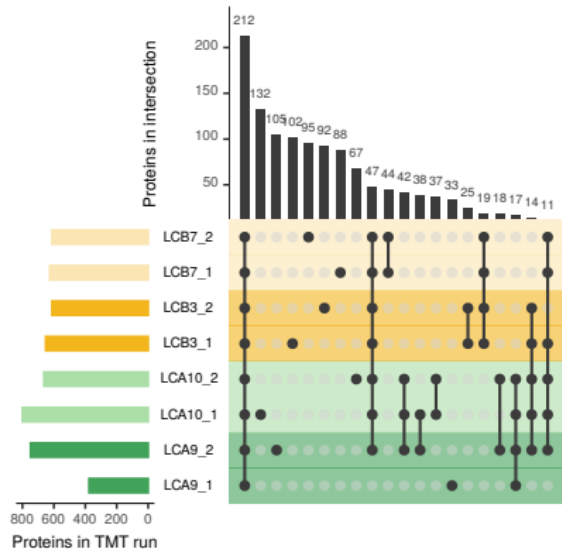
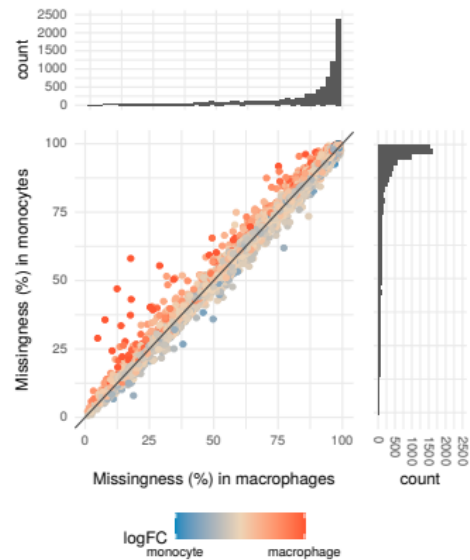


Figure: Missing data is the consequence of biological and technical components (Vanderaa and Gatto, 2021, 2023b).

Challenge 3: 1 + 2

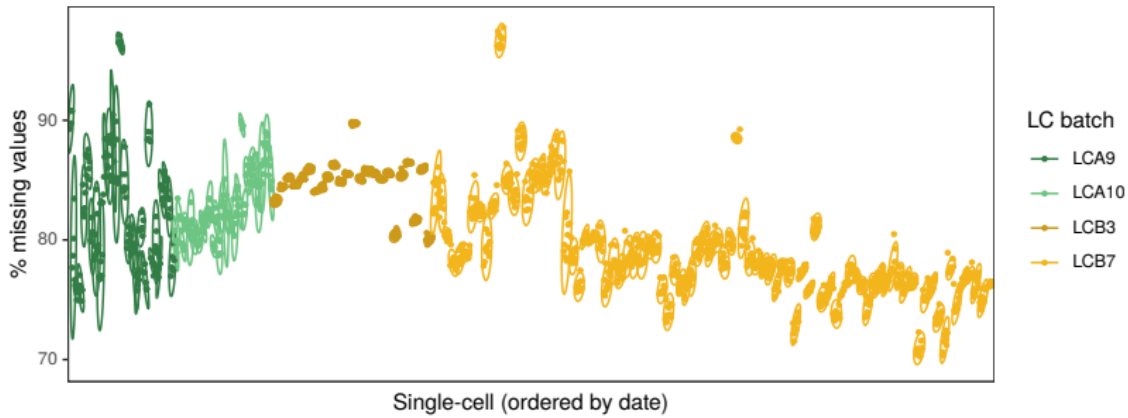


Figure: Influence of batch on data missingness ([Vanderaa and Gatto, 2021](#)).

Data analyses review

- ▶ How do researchers process their data?
- ▶ How do they deal with batch effects?
- ▶ How do they deal with missing data?

Systematic review

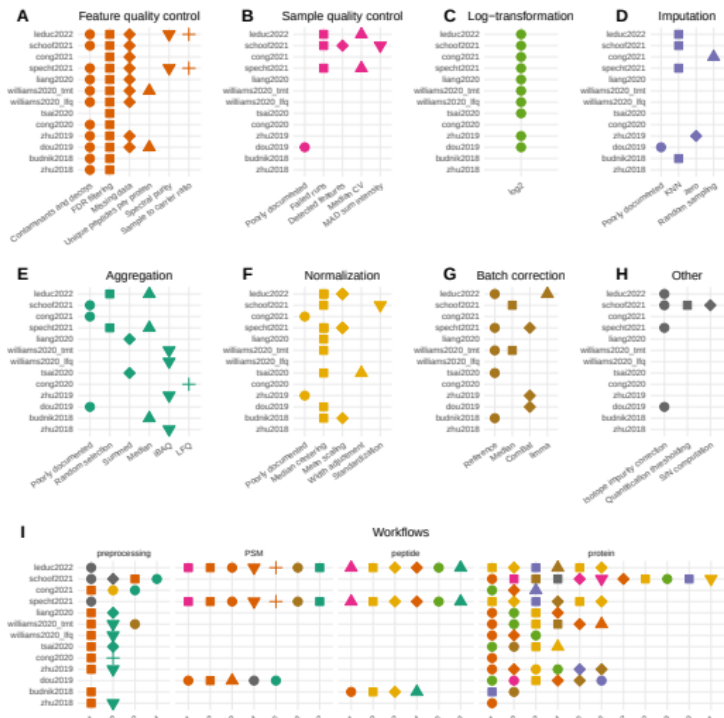


Figure: SCP.replication: systematic reproduction/replication of published SCP studies using the **scp** package - **one workflow per paper/lab..** (Vanderaa and Gatto, 2023a).

Problem

- ▶ Complex data, many alternative pipelines.
- ▶ **Different pipelines produce different results** (see [Vanderaa and Gatto \(2023a\)](#)).
- ▶ Little control/understanding of the implications of what is done to the data.

Problem

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Solution: a principled approach

- ▶ KISS (*Keep it simple stupid!*), as simple as possible.
- ▶ Use what we know to **model** our data.
- ▶ Control what we do, **quantify** effects.

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Given that we aren't sure about the effect of data processing...

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Let's start with **minimally processed data**

- ▶ Remove low quality precursors and cells
- ▶ Aggregate from precursors into peptides
- ▶ \log_2 -transform
- ▶ Remove features with *too many* NAs
- ▶ No imputation

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And use ANOVA–simultaneous component analysis (ASCA)-like methods ([Thiel et al., 2017](#)).

(1) Linear modelling

$$y = \beta_0 + \beta_1 \times group + \epsilon$$

$$y = \beta_0 + \beta_1 \times group + \beta_i \times batch_i + \epsilon$$

(2) Quantify the effects' contributions

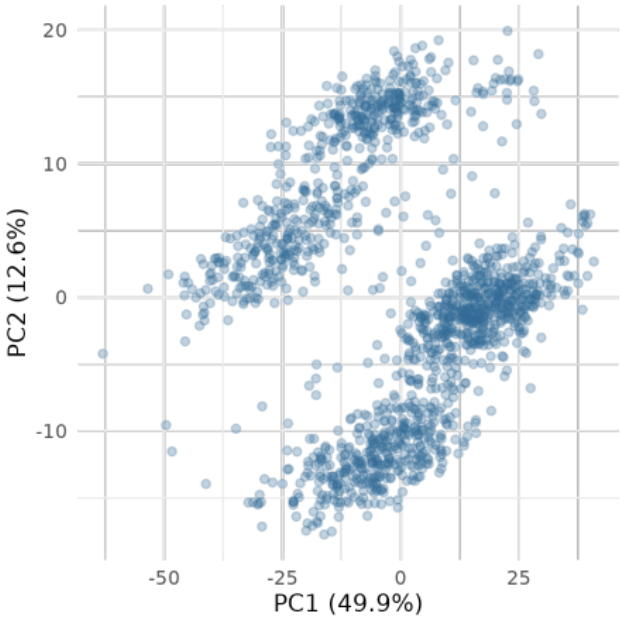
(3) Principal Component Analysis

On **effect + residual** matrices (of dimensions *features* \times *samples*).

Material (2)

The nPOP dataset

- ▶ Data from Leduc et al. (2022)
- ▶ nano-ProteOmic sample Preparation
- ▶ 877 monocytes, 878 melanoma cells
- ▶ 19374 peptides, 3348 proteins
- ▶ **Availability of data and code**



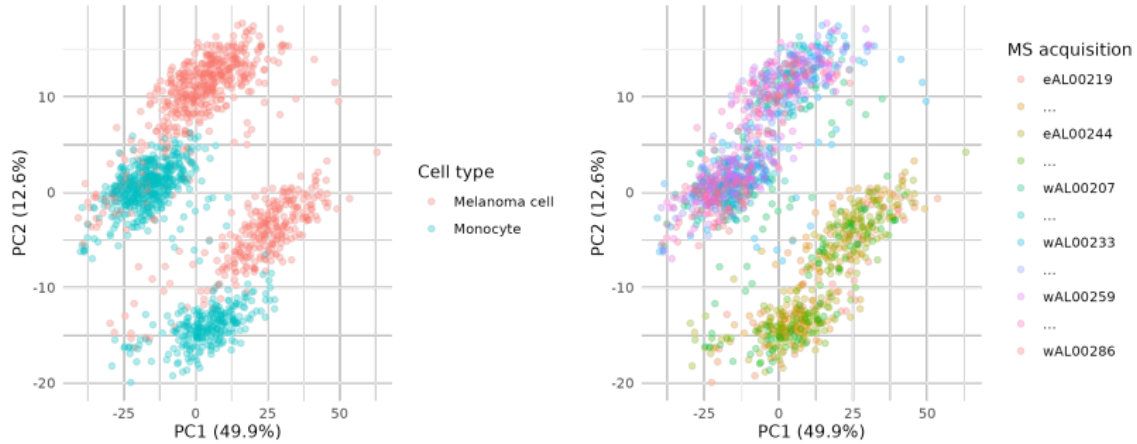


Figure: Melanoma cells and monocytes (left) acquired across multiple acquisition batches (right) (Leduc et al., 2022).

$$y = \textcolor{blue}{MS \ acquisition} + \textcolor{blue}{TMT \ channel} + \textcolor{orange}{Cell \ type} + \epsilon$$

$$y = MS \ acquisition + TMT \ channel + Cell \ type + \epsilon$$

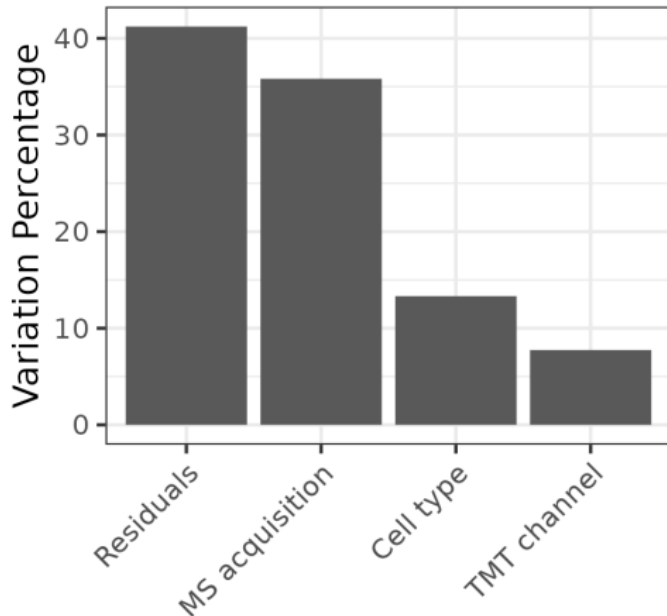


Figure: We are now in a position to **quantify known and unknown**

effects: percentages of explained variances of our explained (known) and unexplained (residuals) effects. NB: low biological variance \neq low quality!

PCA on effect matrices

$$y = \textcolor{red}{MS \ acquisition} + TMT \ channel + Cell \ type + \epsilon$$

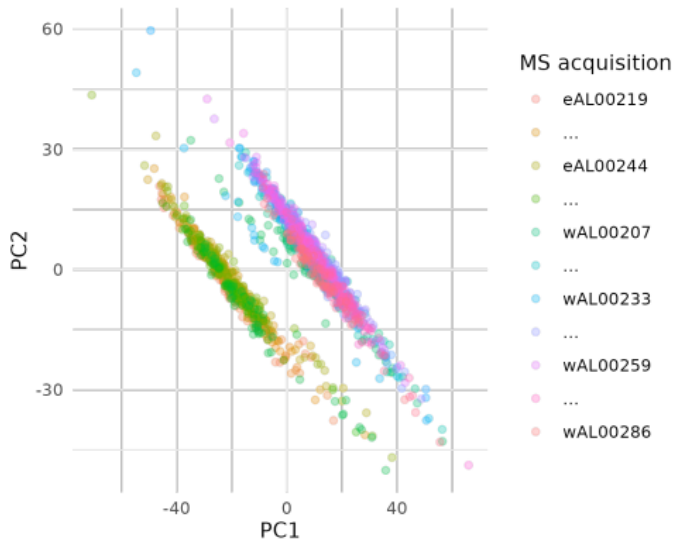
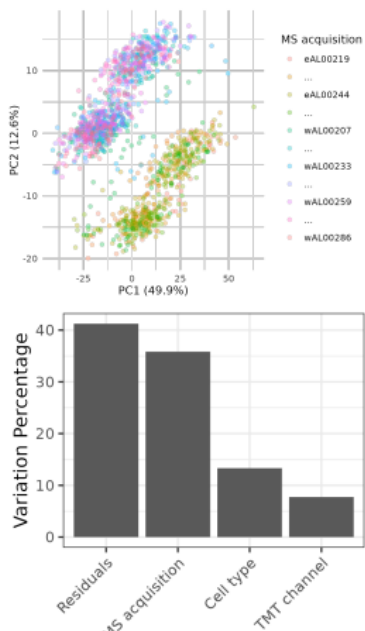


Figure: PCA on the **MS acquisition** effect matrix.

PCA on effect matrices

$$y = MS \text{ acquisition} + TMT \text{ channel} + Cell \text{ type} + \epsilon$$

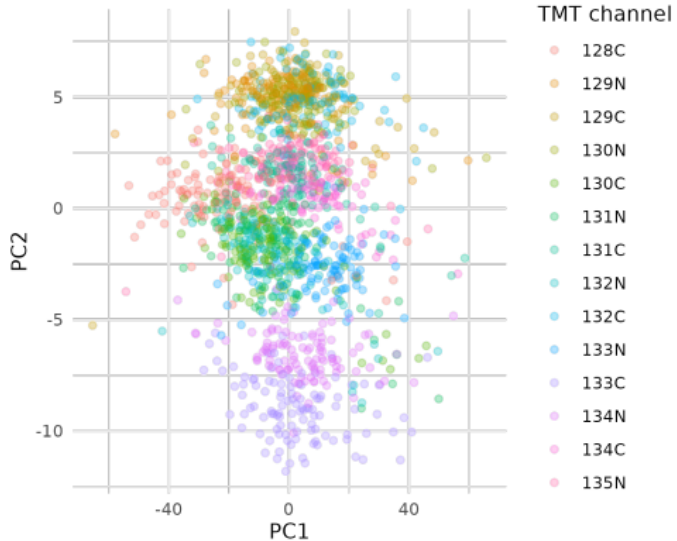
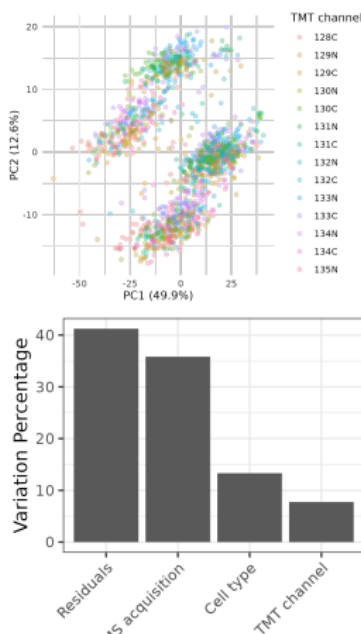


Figure: PCA on the **TMT channel** effect matrix.

PCA on effect matrices

$$y = MS \text{ acquisition} + TMT \text{ channel} + \text{Cell type} + \epsilon$$

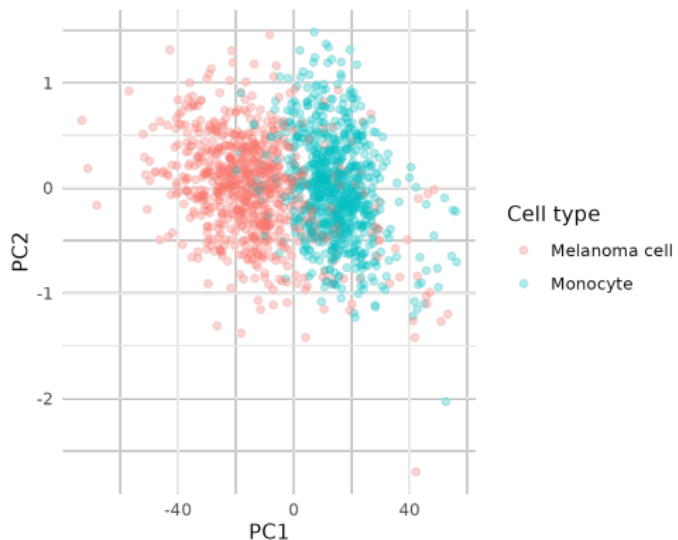
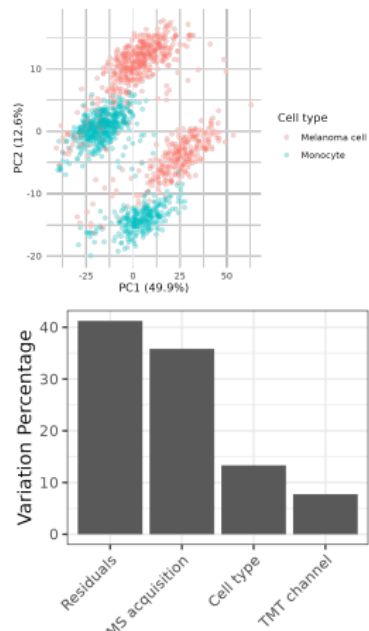


Figure: PCA on the **Cell type** effect matrix.

PCA on effect matrices

$$y = MS \text{ acquisition} + TMT \text{ channel} + Cell \text{ type} + \epsilon$$

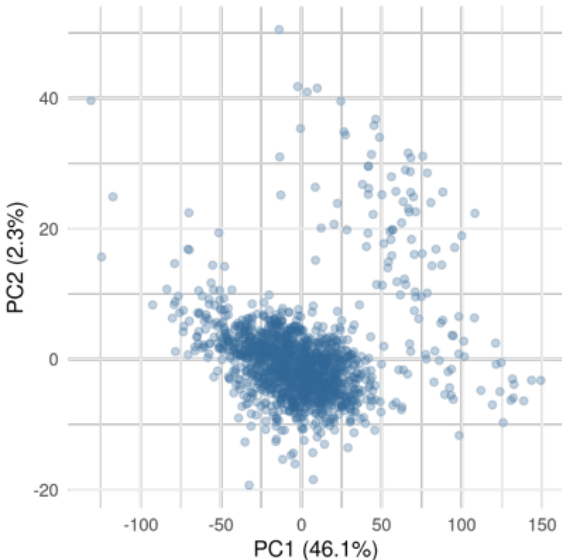
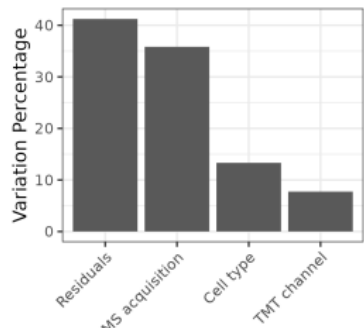
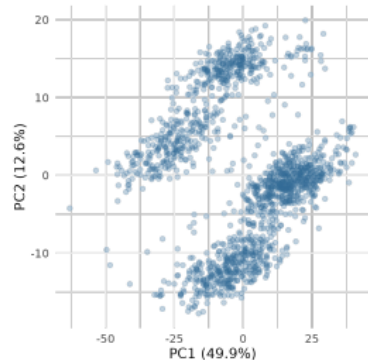
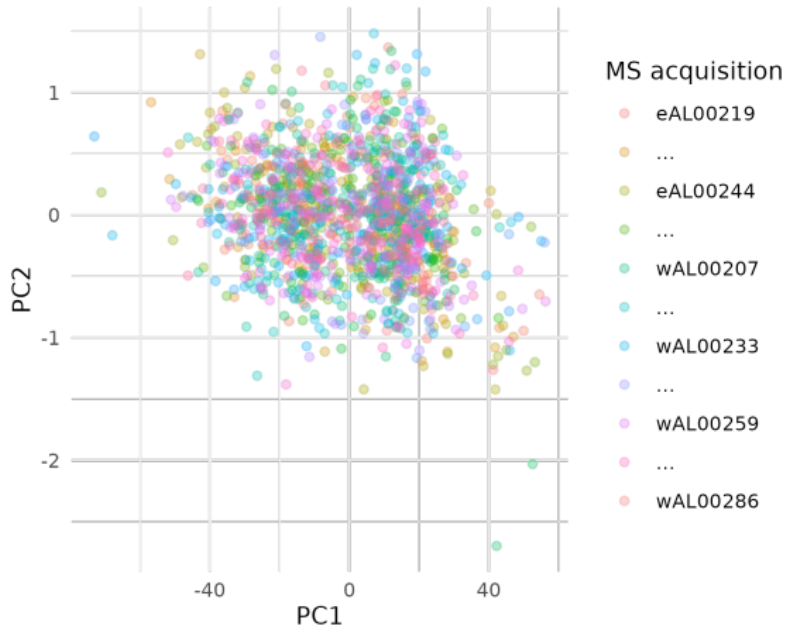


Figure: PCA on the **residuals** effect matrix.

Does it work: negative control

Do we have any MS acquisition batch leftovers in the cell type effect?

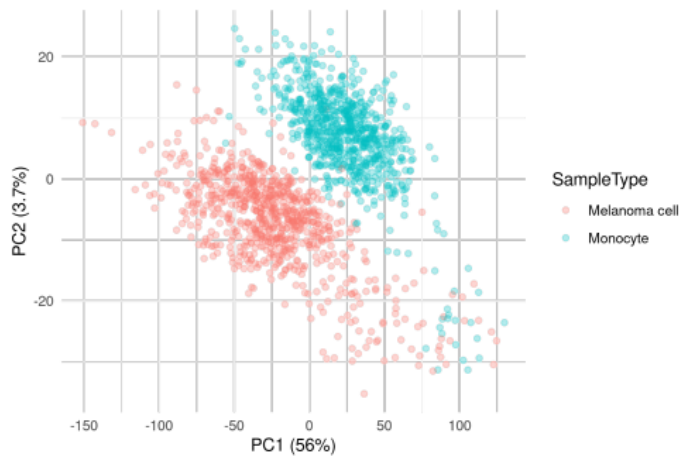
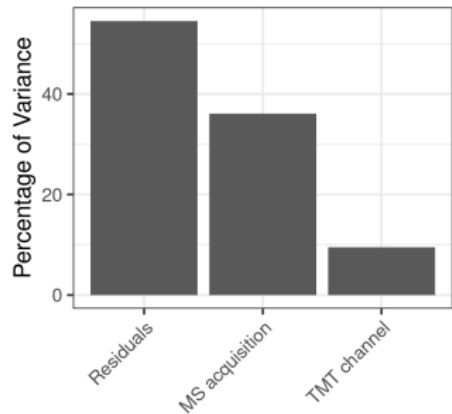


Does it work: positive control

$$y = \text{MS acquisition} + \text{TMT channel} + \epsilon$$

Does it work: positive control

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Does it work: new biology in the residuals

$$y = \text{MS acquisition} + \text{TMT channel} + \text{Cell type} + \epsilon$$

Does it work: new biology in the residuals

$$y = \text{MS acquisition} + \text{TMT channel} + \text{Cell type} + \epsilon$$

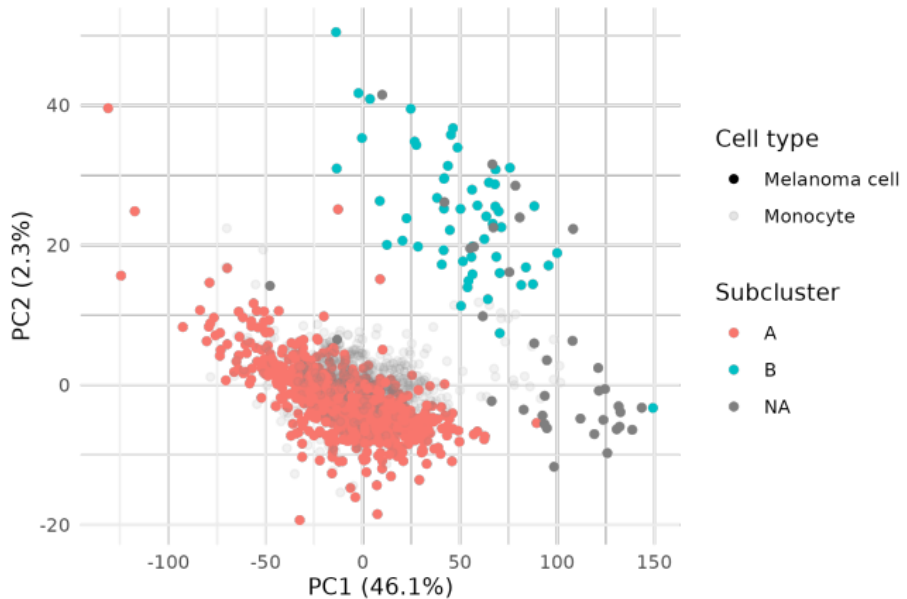


Figure: Melanoma subpopulations: transcriptomic signature associated with a cell state that is more likely to resist treatment by the cancer drug vemurafenib (clusters A and B from [Leduc et al. \(2022\)](#))

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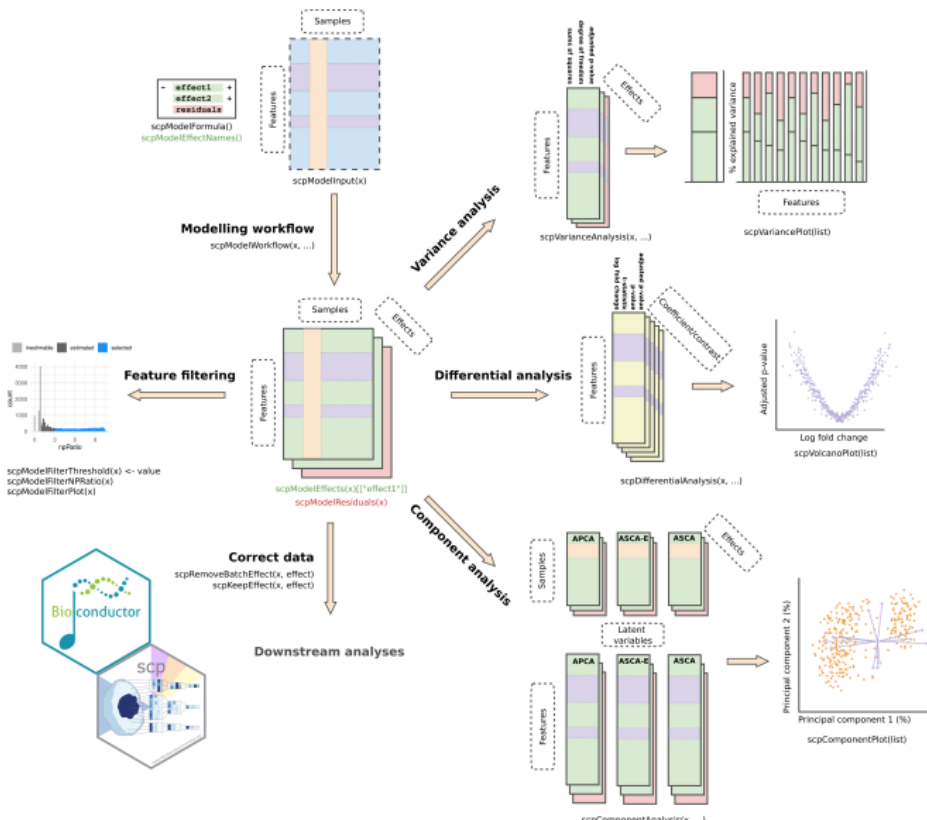


Figure: scplainer package - scplainer: using linear models to understand mass spectrometry-based single-cell proteomics data (Vanderaa and Gatto, 2025).

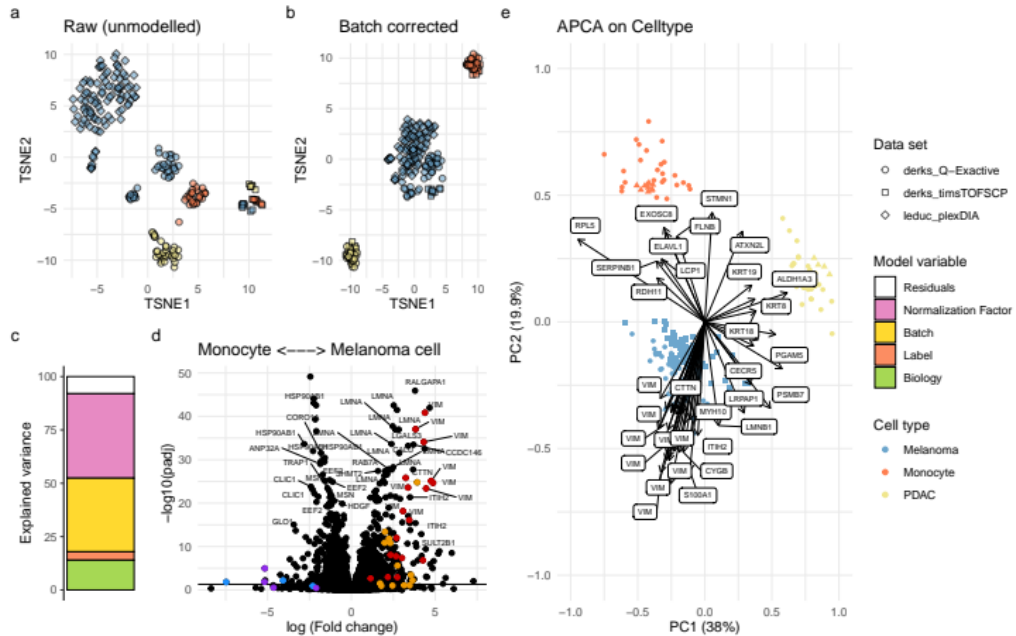


Figure: scplainer – variance, differential and component analysis, integration

What are best RSE practice?

1. ...
2. ...
3. ...

Our software

- ▶ <https://bioconductor.org/packages/QFeatures>
- ▶ <https://bioconductor.org/packages/scp>
- ▶ <https://bioconductor.org/packages/scpdata>

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- ▶ Many experimental and computational workflows. Different workflows → different results.
- ▶ We need a flexible and **principled computational approach** → control what we do, to guarantee the validity of our results.
- ▶ **Residuals** – what we don't know (yet), generally what we are most interested in.
- ▶ Showed component analysis, differential abundance, analysis of variance. Also clustering, trajectory analysis, ... based on the batch-corrected/normalised effect matrices.
- ▶ **Limitation:** multi-patient/condition designs - mixed effects (Sticker et al., 2020) and pseudo-bulking.

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- ▶ **Limitation:** multi-patient/condition designs - mixed effects (Sticker et al., 2020) and pseudo-bulking.
- ▶ Work openly and reproducibly! (Markowetz, 2015).
- ▶ Importance of the **experimental design** (Gatto et al., 2023).
- ▶ Better software, better methods, better research.

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- ▶ CBIO lab

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Discussion points

- ▶ Your results are only as good as your method, software and your users.
- ▶ Is a computational researcher coding doing research?
- ▶ What is good software? What is data analysis?
- ▶ Should all software meet the highest standard? Should every piece of research be 100% reproducible?
- ▶ What about LLM-generated code?

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