

The msdata2 data package for proteomics benchmarking

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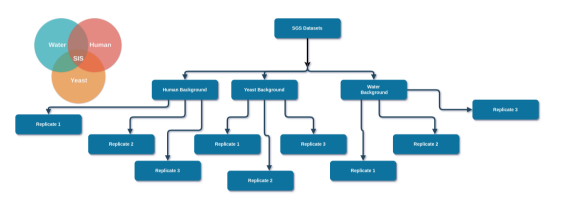
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Summary

The msdata2 package contains a set of published quantification/identification proteomics datasets. The purpose of the package is to provide standard and curated datasets to facilitate the benchmarking of proteomics workflows.

The SGS Datasets

The first dataset in msdata2 is a SWATH experiment processed with OpenSWATH output from SGS dataset. The SWATH-MS Gold Standard (SGS) dataset consists of 90 SWATH-MS runs of 422 synthetic stable isotopelabeled standard (SIS) peptides in ten different dilution steps (1, 2, 4, 8, ..., 512 times), spiked into three protein backgrounds of varying complexity (water, yeast and human), acquired in three technical replicates [1].



The SGS dataset was manually annotated, resulting in 342 identified and quantified peptides with three or four transitions each. In total, 30,780 chromatograms were inspected and 18,785 were annotated with one true peak group, whereas in 11,995 cases no peak was detected. The data were processed and converted into MSnSet objects [2] (see below and on the right):

Slot	Information
assayData	quantitative matrix with XIC values
phenoData	sample metadata
featureData	feature metadata (identification data, peptides sequences,)
experimentData	experimental methods and general annotations
processingData	processing information and log

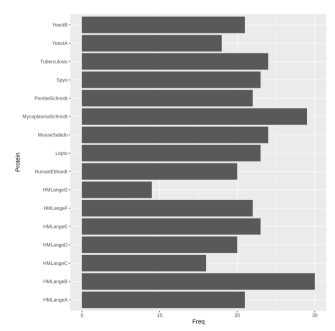


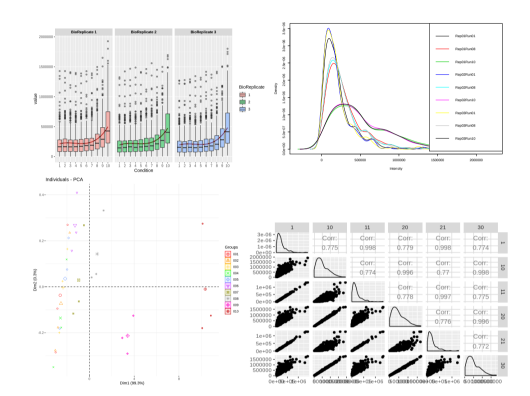
Figure 1: Unique peptide count for each protein in the human background.

Conclusion and Future Work

Our package msdata2 will provide formatted labelled and label-free quantification and raw data for proteomics. In the future, we will also include large raw MS data, making use of the ExperimentHub cloud infrastructure. Our goal is for msdata2 to become a benchmarking tools on different computational proteomics workflows.

Data exploration

Data visualization on the MSnSet with Human background. We can notice the good consistency between replicates. The principal component analysis confirms good separation for runs 10 to 7 (up to dilution 8x). Further dilutions become much more difficult to tell apart.



Data preparation

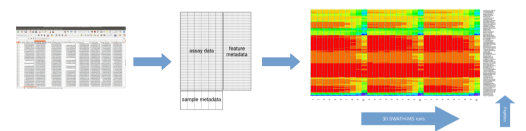


Figure 2: The OpenSWATH outputs for SGS dataset (human background) are cleaned, formatted and compressed as MSnSet object in msdata2 [2].

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References

- $[1] \ \ R\"{o}st, H.L. \ \ et \ \ al. \ \ (2014): \ \ OpenSWATH \ \ enables \ \ automated, \ targeted \ \ analysis$
- of data-independent acquisition ms data. Nat. Biotechnol., 32, 219–223.
 [2] Gatto L, Lilley K (2012): MSnbase an R/Bioconductor package for isobaric tagged mass spectrometry data visualization, processing and quantitation. Bioinformatics, 28, 288-289.