

# Tutorial Overview

## Outline

In this series of 6 tutorials we will learn how to use various tools for the analysis of DIA data (see figure 1). These tutorials can be grouped into 3 conceptual stages in the analysis workflow (red sections in figure 1):

### *Library Generation*

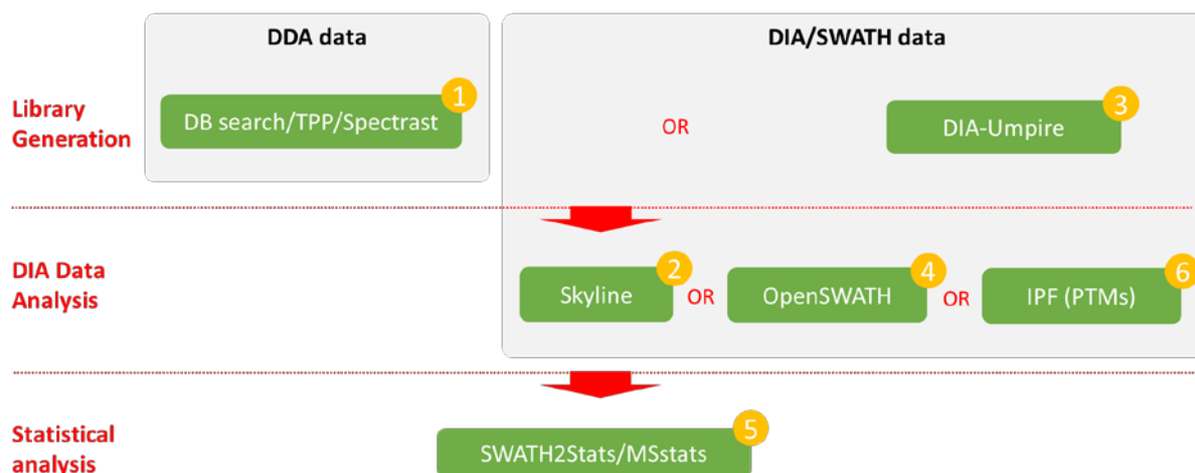
The first component required for analysis of DIA data is library generation. We will do this using two different approaches. In tutorial 1 we will perform a standard database search<sup>1-5</sup> of DDA data generated from the samples of interest to build a spectral library from which we will generate peptide query parameters to later analyze the DIA data. In tutorial 3, we will similarly build a spectral library, however, this will be done directly from the DIA data using DIA-Umpire<sup>6</sup>. Note that in a real experiment you would only need to do one of these approaches but we will do both for illustration.

### *DIA data analysis*

We will use Skyline<sup>7</sup> (tutorial 2) or OpenSWATH<sup>8</sup> / PyProphet<sup>9</sup> / TRIC<sup>10</sup> (tutorial 4) to perform peptide-centric analysis of the DIA data using the libraries we have built in the first part. Again, here we are showing 2 alternatives here for analysing the DIA data but under normal circumstances you would perform analyses with one or other of these tools (or related tools). We will also analyse a separate phosphopeptide data set using IPF<sup>11</sup> (inference of peptidofoms) extension of the OpenSWATH workflow (tutorial 6).

### *Statistical analysis*

In tutorial 5, the output from the DIA data analysis will be subjected to comparative statistical analysis using the R package MSstats<sup>12</sup>. To prepare the data for MSstats analysis we will use another R package, SWATH2Stats<sup>13</sup>, to perform some filtering and reformatting.



**Figure 1 – DIA/SWATH Course tutorial overview**

In addition to these tutorials where you will follow a written step-by-step procedure, we will have a walk-through session where Brendan MacLean, principle Skyline developer, will demonstrate the analysis of another data set in Skyline online with participants following his analysis.

## Datasets

The data we will analyze come from samples reconstituted to match those used in the LFQBenchmark study<sup>14</sup> where quantitative benchmarking samples were created by mixing proteomes of 3 organisms in defined ratios (figure 2). The DIA (SWATH-MS) data were acquired in technical triplicate on a QqTOF instrument (6600 TripleTOF, AB Sciex) using a 64 variable width window precursor isolation scheme, or on a QqOrbitrap instrument (Q-Exactive Plus) using a 19 variable width window precursor isolation scheme. Similarly, DDA data were acquired by analyzing these samples for the purpose of library generation. By using these data sets we have a ground truth regarding the quantitative differences between samples A and B. This will allow us to determine whether our analysis is performing well or not. You can choose whether you want to perform the data analysis for the TripleTOF or Q-Exactive data set.

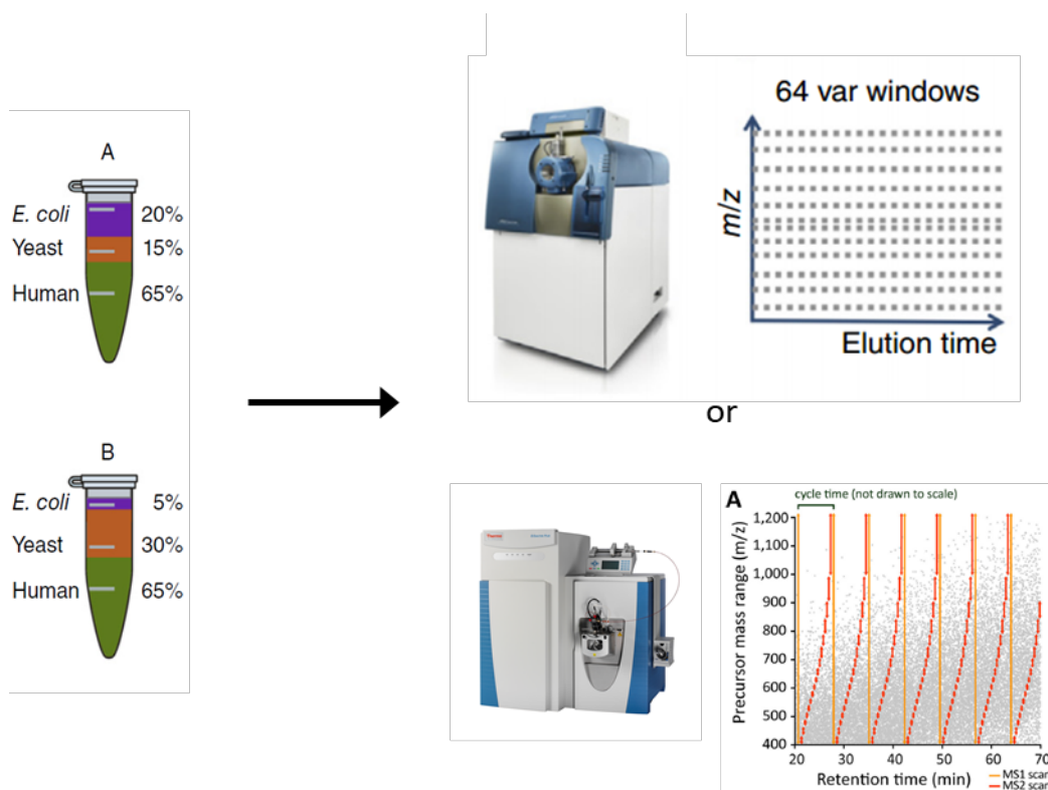
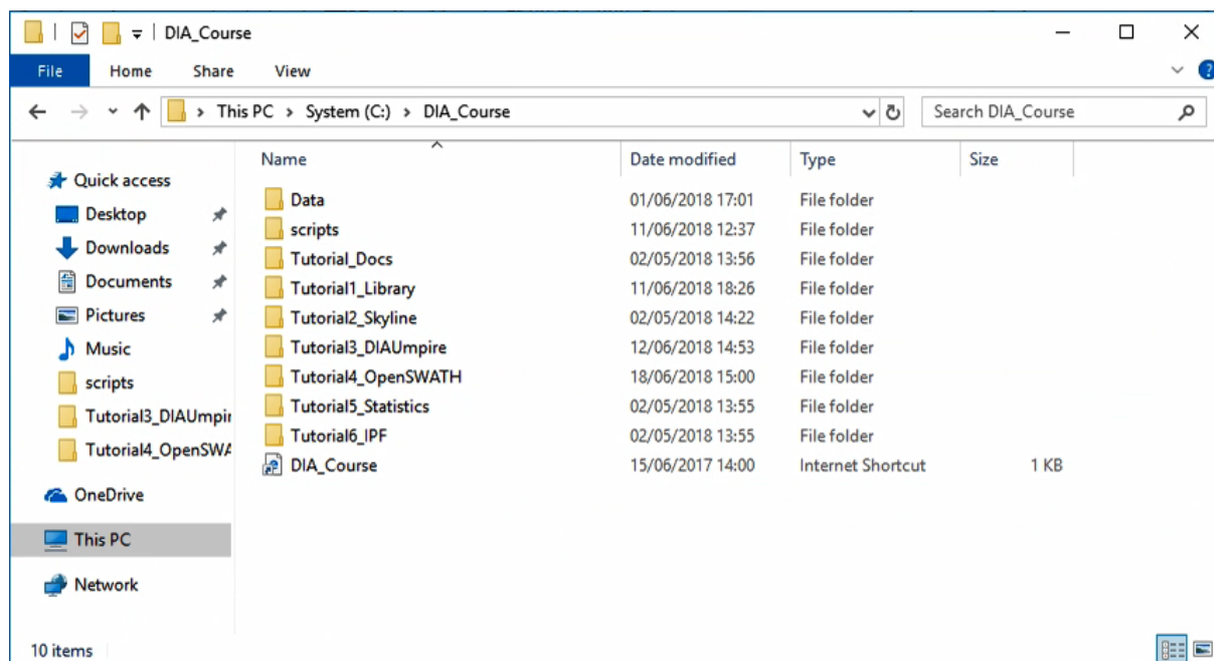


Figure 2 – DIA/SWATH Course data overview (adapted from<sup>14,15</sup>)

## Directory Structure

All of the data files you need to perform the analysis are already loaded onto the virtual machines we will use for the analysis in the directory “C:\DIA\_Course” (a copy of all data with the same structure is provided on your USB stick). In this directory you will find individual folders for each of the 5 tutorials which contain some files specific to each tutorial. The DDA and DIA data you need are in the “Data” directory along with some other general files that will be required. There is also a directory called “backup”. This directory contains all of the intermediate and final files from the completed tutorials. In case you experience some problems in a given section, you should be able to use the files in here to continue the tutorials successfully.



## Appendix 1 – Software used in the tutorials

In the information email sent before the course we included a complete list of software and dependencies that had been installed on the virtual machines for the purpose of the tutorials. Here we summarize again the tools required and provide links where they can be downloaded with detailed instructions for installation.

- Trans Proteomic Pipeline

Link:

<http://tools.proteomecenter.org/wiki/index.php?title=Software:TPP>

[http://tools.proteomecenter.org/wiki/index.php?title=Windows\\_Installation\\_Guide](http://tools.proteomecenter.org/wiki/index.php?title=Windows_Installation_Guide)

A broad collection of software tools from the Institute for Systems Biology, Seattle focused primarily on DDA analysis of various types. This software is relatively easy to install on Windows (requires pre-installation of Perl and some MSVC/.Net Frameworks components) and is supported on Mac and Linux. The components we are using: Comet, PeptideProphet, InterProphet, SpectraST

- Skyline

Link:

<https://skyline.ms/wiki/home/software/Skyline/page.view?name=default>

Skyline is a freely-available and open source Windows client application for building Selected Reaction Monitoring (SRM) / Multiple Reaction Monitoring (MRM), Parallel Reaction Monitoring (PRM - Targeted MS/MS), Data Independent Acquisition (DIA/SWATH) and targeted DDA with MS1 quantitative methods and analyzing the resulting mass spectrometer data. Skyline is very well tested and supported and is easy to use due to a rich graphical user interface.

- DIA-Umpire

Link:

<http://diaumpire.sourceforge.net/>

DIA-Umpire is an open source Java program for computational analysis of data independent acquisition (DIA) mass spectrometry-based proteomics data. It enables untargeted peptide and protein identification and quantitation using DIA data.

- OpenSWATH / PyProphet / TRIC

Link:

<http://openswath.org/en/latest/>

The OpenSWATH Workflow enables targeted data analysis of data-independent acquisition (DIA) or SWATH-MS proteomic data. OpenSWATH is part of, and distributed with, the OpenMS (<http://www.openms.de/>) project. PyProphet and TRIC are Python tools. These tools require Python and several other dependencies to be installed. A tutorial focused specifically on these tools has recently been published<sup>16</sup>.

- SWATH2stats

Link:

<https://bioconductor.org/packages/release/bioc/html/SWATH2stats.html>

SWATH2stats is an R package intended to transform SWATH data from the OpenSWATH software into a format readable by other statistics packages while performing filtering, annotation and FDR estimation. SWATH2stats requires R and is distributed by the Bioconductor project (<https://bioconductor.org/>)

- MSstats

Link: <http://msstats.org/msstats-2/>

MSstats is an open-source R package for statistical relative quantification of proteins and peptides in global, targeted and data-independent proteomics. It requires R and is also available from the Bioconductor project (<https://bioconductor.org/>).

- RStudio

Link: <https://www.rstudio.com/products/rstudio/download/>

RStudio is an integrated development environment (IDE) for R. It includes a console, syntax-highlighting editor that supports direct code execution, as well as tools for plotting, history, debugging and workspace management. In this course we use this environment for executing R commands but also as an interface to Git bash terminal for all of the command tools.

- mapDIA

Link: <http://mapdia.sourceforge.net/Main.html>

mapDIA is software for statistical analysis of differential expression using MS/MS fragment-level quantitative data from data independent acquisition (DIA) proteomics experiments. It offers a series of tools for essential data preprocessing, including a novel retention time-based normalization method and multiple peptide/fragment selection steps.

## Appendix 2 – Raw data file conversion and centroiding

The raw data (DIA and DDA) has been converted to the mzXML<sup>17</sup> format. The data has further been centroided, and the DIA data has had an absolute intensity threshold applied to reduce the file size for the purpose of the course. The commands to perform these steps, using the ProteoWizard<sup>18</sup> tool MSConvert, are given below (note: there is also a GUI version of this tool, MSConvertGUI, available if you prefer this to the command line).

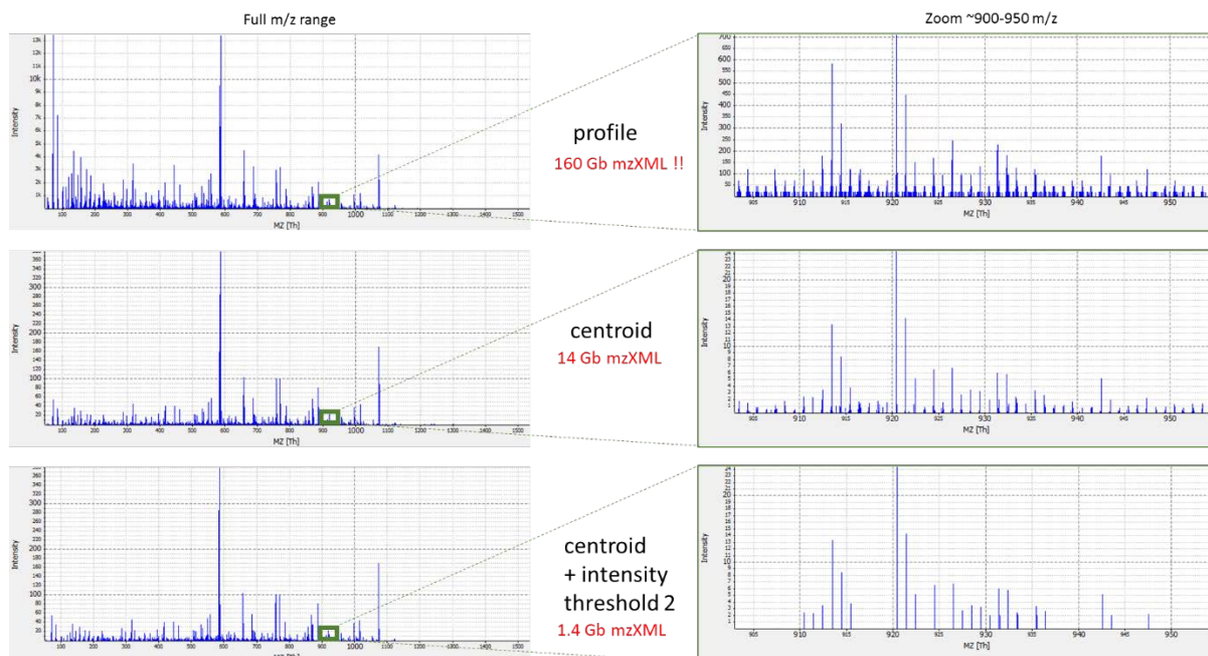
Note: While we do routinely analyze data either in profile or centroid mode, we would not typically apply the absolute intensity threshold. This thresholding has been applied for the purpose of the course to reduce the file size and reduce the analysis time. While the quality of the analysis should not be significantly affected by this step. Examples of how centroiding and thresholding affect the data are shown in Figure 3 below.

- Commands for converting to mzXML and centroiding

```
"c:\Program Files\ProteoWizard\ProteoWizard 3.0.10765\msconvert.exe"  
--mzXML --filter "peakPicking true 1-" *.wiff
```

- Commands for converting to mzXML, centroiding, and thresholding (filter order is important!)

```
"c:\Program Files\ProteoWizard\ProteoWizard 3.0.10765\msconvert.exe"  
--mzXML --filter "peakPicking true 1-" --filter "threshold absolute  
2 most-intense" *.wiff
```



**Figure 3 – effects of centroiding and intensity thresholding**

### References

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