## A Practical Beginner's Guide to Proteomics

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### **Abstract**

Proteomics is the large-scale study of protein structure and function from biological systems. "Shotgun proteomics" or "bottom-up proteomics" is the prevailing strategy, in which proteins are hydrolyzed into peptides that are analyzed by mass spectrometry. Proteomics studies can be applied to diverse studies ranging from simple protein identification to studies of protein-protein interactions, post-translational modifications, and protein stability. To enable this range of different experiments, there are diverse strategies for proteome analysis. The nuances of how proteomics workflows differ may be challenging to understand for new practitioners. Here, we provide a comprehensive tutorial of different proteomics methods. Our tutorial covers all necessary steps starting from protein extraction and ending with biological interpretation. We expect that this work will serve as a basic resource for new practitioners of the field of shotgun or bottom-up proteomics.

### Introduction

Proteomics is the large scale study of protein structure and function. Proteins are translated from mRNAs that are transcribed from the genome. Although the genome encodes potential cellular functions and states, the study of proteins is necessary to truly understand biology. Currently, proteomic studies are facilitated by mass spectrometry, although alternative methods are being developed.

Modern proteomics started around the year 1990 with the introduction of soft ionization methods that enabled, for the first time, transfer of large biomolecules into the gas phase without destroying them [[1]][2]. Shortly afterward, the first computer algorithm for matching peptides to a database was introduced [3]. Another major milestone that allowed identification of over 1000 proteins were actually improvements to chromatography [4]. As the volume of data exploded, methods for statistical analysis transitioned use from the wild west to modern informatics based on statistical models [5] and the false discovery rate [6].

Two strategies of mass spectrometry-based proteomics differ fundamentally by whether proteins are cleaved into peptides before analysis: "top-down" and "bottom-up". Bottom-up proteomics (also refered to as shotgun proteomics) is defined by the hydrolysis of proteins into peptide pieces [7]. Therefore, bottom-up proteomics does not actually measure proteins, but must infer their presence [8]. Sometimes proteins are infered from only one peptide sequence representing a small fraction of the total protein sequence predicted from the genome. In contrast, top-down proteomics attempts to measure all proteins intact [9]. The potential benefit of top-down proteomics is the ability to measure proteoforms [10]. However, due to analytical challenges, the depth of protein coverage that is achievable by top-down proteomics is less than the depth that is achievable by bottom-up proteomics.

In this tutorial we focus on the bottom-up proteomics workflow. The most common version of this workflow is generally comprised of the following steps. First, proteins in a biological sample must be extracted. Usually this is done by denaturing and solubilizing the proteins while disrupting DNA and tissue. Next, proteins are hydrolyzed into peptides, usually using a protease like trypsin. Peptides from proteome hydrolysis must be purified. Most often this is done with reversed phase chromatography cartridges or tips. The peptides are then almost always separated by liquid chromatography before they are ionized and introduced into a mass spectrometer. The mass spectrometer then collects precursor and fragment ion data from those peptides. The data analysis is usually the rate limiting step. Peptides must be identified, and proteins are inferred and quantities are assigned. Changes in proteins across conditions are determined with statistical tests, and results must be interpreted in the context of the relevant biology.

There are many variations on this workflow. The wide variety of experimental goals that are achievable with proteomics technology leads to a wide variety of potential proteomics workflows. Even choice is important and every choice will affect the results. In this tutorial, we cover all of the required steps in detail to serve as a tutorial for new proteomics practioners.

- 1. Types of experiments enabled by proteomics
- 2. Protein extraction
- 3. proteolysis
- 4. Isotopic Labeling
- 5. Enrichments
- 6. Peptide purification
- 7. Mass Spectrometry
- 8. Peptide Ionization

- 9. Data Acquisition10. Basic Data Analysis11. Biological Interpretation12. Experimental considerations and design

### **Types of Experiments**

A wide range of questions are addressable with proteomics technology, which translates to a wide range of variations of proteomics workflows. Sometimes identifying what proteins are present is desired, and sometimes the quantities of as many proteins as possible are desired. Proteomics experiments can be both qualitative and quantitative.

### **Qualitative experiments**

- Identifying proteins
- Identifying post translational modifications
- Identifying protein isoforms

### **Quantitative experiments**

- Protein abundance changes
- Phosphoproteomics
- Glycoproteomics
- Structural techniques (XL-MS, HDX-MS, FPOP, protein-painting, LiP-MS, radical footprinting, ion mobility)
- Protein stability and small molecule binding (Thermal proteome profiling, TPP, or cellular thermal shift assay, CETSA)
- Protein-protein interactions (PPIs): AP-MS, APEX, BioID

### **Protein Extraction**

First, proteins must be isolated from the sample matrix. Because some proteins alter other proteins, the goal is to simultaneously solubilize and denature proteins. This is achieved with a combination of salt and chaotropic agent.

- 1. Choice of Lysis buffer
- Urea can cause chemical modifications
- 2. Sample type and homogenisation methods
- specialised sample preparation protocols for non-denaturing protein isolation (i.e. for LiP-MS, HDMX-MS etc)
- 4. chemicals to avoid: PEGs, detergents etc
- 5. removal of contaminations, Protein Precipitation
- detergent reomoval resins, S-TRAP (Protifi) etc
- 7. protein alkylation
- choices of reduction and alkylation reagents, TCEP/DTT/2BME, Chloroacetamide/iodoacetamide, nethyl maleimide

### **Proteolysis**

Proteolysis is the defining step that differentiates bottom-up or shotgun proteomics from top-down proteomics. Hydrolysis of proteins is extremely important because it defines the population of potentially identifyable peptides. Generally peptides between a length of 7-35 amino acid are considered useful for mass spectrometry analysis. Peptides that are too long are difficult to identify by tandem mass spectrometry, or may be lost during sample preparation due to Peptides that are too short are less likely to uniquely match to a single protein. There are many choices of enzymes and chemicals that hydrolyze proteins into peptides. This section summarizes potential choices and their strengths and weaknesses.

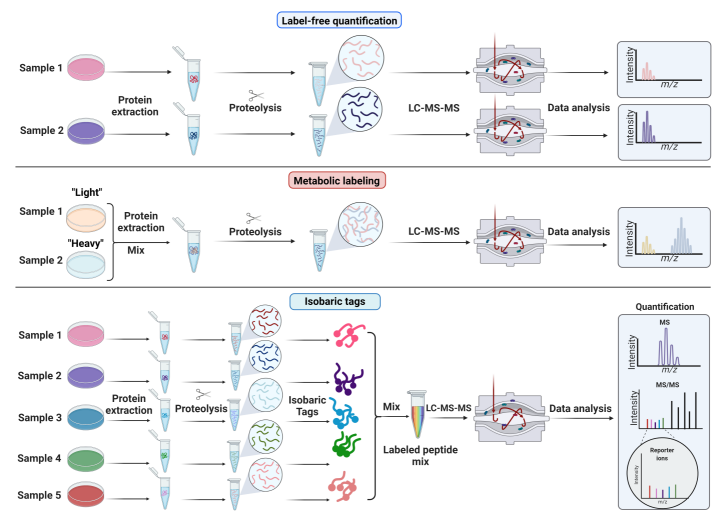
Trypsin is the most common choice of protease for proteome hydrolysis [11].

- 3. theoretical studies of proteolysis enzymes [12]
- 4. Challenges associated with alternative enzyme choices (non-specific and semi-specific enzymes)
- 5. Alternative enzyme choices (one paragraph each?) LysC
- 6. GluC
- 7. AspN
- 8. Alpha-lytic protease [13] and how it enables mapping human SUMO sites [14].
- 9. others?

### **Peptide and Protein Labeling**

Discussion of methods to isotopically label peptides or proteins that enable quantification

- 1. SILAC/SILAM
- 2. iTRAQ
- 3. TMT
- 4. dimethyl labeling



**Figure 1: Quantitative strategies commonly used in proteomics.** A) Label-free quantitation. Proteins are extracted from samples, enzymatically hydrolyzed into peptides and analyzed by mass spectrometry. Chromatographic peak areas from peptides are compared across samples that are analyzed sequentially. B) Metabolic labelling. Stable isotope labeling with amino acids in cell culture (SILAC) is based on feeding cells stable isotope labeled amino acids ("light" or "heavy"). Samples grown with heavy or light amino acids are mixed before cell lysis. The relative intensities of the heavy and light peptide are used to compute protein changes between samples. C) Isobaric or chemical labelling. Proteins are isolated separately from samples, enzymatically hydrolyzed into peptides, and then chemically tagged with isobaric stable isotope labels. These isobaric tags produce unique reporter mass-to-charge (m/z) signals that are produced upon fragmentation with MS/MS. Peptide fragment ions are used to identify peptides, and the relative reporter ion signals are used for quantification.

## **Peptide or Protein Enrichment**

### Protein enrichment (e.g. for protein protein interactions)

- APEX
- bioID
- bioplex

## **Peptide enrichment**

- antibody enrichments of modifications, e.g. lysine acetylation [15].
- TiO2 and Fe enrichment of phosphorylation
- Glycosylation
- SISCAPA

## **Methods for Peptide Purification**

- 1. Reverse phase including tips and cartridges
- 2. stage tips
- 3. in stage tip (iST)
- 4. SP2, SP3
- 5. s traps

## **Types of Mass Spectrometers used for Proteomics**

- 1. QQQ
- 2. Q-TOF
- 3. Q-Orbitrap
- 4. LTQ-Orbitrap
- 5. TOF/TOF
- 6. FT-ICR
- 7. types of ion mobility
- SLIM
- FAIMS
- traveling wave
- tims

## **Peptide Ionization**

The 2002 Nobel Prize in Chemistry was awarded to partially to John Fenn and Koichi Tanaka "for their development of soft desorption ionisation methods for mass spectrometric analyses of biological macromolecules" [16/].

#### MALDI

### **Electrospray Ionization**

## **Data Acquisition**

Data acquisition strategies for proteomics fall generally within targeted or untargeted, and they can depend on the data (data dependent acquisition or DDA) or be data independent (data-independent acquisition or DIA).

#### **DDA**

**Targeted DDA** 

**Untargeted DIA** 

DIA

**Targeted DIA** 

**Untargeted DIA** 

### **Analysis of Raw Data**

The goal of basic data analysis is to convert raw spectral data into identities and quantities of peptides and proteins that can be used for biologically-focused analysis. This step may often include measures of quality control, cross-run data normalization, quantification on different levels (precursor, peptide, protein), protein inference, PTM (post translational modification) localization and also first steps of data analysis, such as statistical hypothesis tests.

In typical bottom-up proteomics experiments, proteins are digested into peptides and further analyzed with LC-MS/MS systems. Peptides can have different PTMs and ionize differently depending on their length and amino acid distributions. Therefore, mass spectrometers often record different charge and modification states of one single peptide. The entity that is recorded on a mass spectrometer is usually referred to as a precursor ion (peptide with its modification and charge state). This precursor ion is fragmented and the precursor or peptide sequences are obtained though spectral matching. The quantity of a precursor is estimated with various methods. The measured precursor quantities are combined to generate a peptide quantity. Peptides are also often combined into a protein group through protein inference, which combines multiple peptide identifications into a single protein identification [17] [18]. Protein inference is still a challenge in bottom-up proteomics.

Due to the inherent differences in the data structures of DDA and DIA measurements, there exist different types of software that can facilitate the steps mentioned above. The existing software for DDA and DIA analysis can be further divided into freeware and non-freeware:

#### **DDA** freeware

Name	Publication	Website
MaxQuant	Cox and Mann, 2008[ <u>19</u> ]	<u>MaxQuant</u>
MSFragger	Kong et al., 2017[ <u>20</u> ]	<u>MSFragger</u>
Mascot	Perkins et al., 1999[ <u>21</u> ]	<u>Mascot</u>
MS-GF+	Kim et al., [ <u>22</u> ]	MS-GF±

#### **DIA freeware:**

Name	Publication	Website
MaxDIA	Cox and Mann, 2008[ <u>19</u> ]	<u>MaxQuant</u>
Skyline	MacLean et al., 2010[ <u>23</u> ]	<u>Skyline</u>
DIA-NN	Demichev et al., 2019[ <u>24</u> ]	<u>DIA-NN</u>

### **Targeted proteomics freeware:**

Name	Publication	Website
Skyline	MacLean et al., 2010[ <u>23</u> ]	<u>Skyline</u>

#### **DDA non-freeware:**

Name	Publication	Website
ProteomeDis coverer		<u>ProteomeDiscoverer</u>
Mascot	Perkins et al., 1999[ <u>21</u> ]	<u>Mascot</u>
Spectromine		<u>Spectromine</u>
PEAKS	Tran et al., 2018[ <u>25</u> ]	<u>PEAKS</u>

### **DIA non-freeware:**

Name	Publication	Website
Spectronaut	Bruderer et al., 2015[ <u>26</u> ]	<u>ProteomeDiscoverer</u>
PEAKS	Tran et al., 2018[ <u>25</u> ]	<u>PEAKS</u>

## **Analysis of DDA data**

Strategies for analysis of DIA data

**Targeted proteomics data analysis** 

**Quality control** 

**Statistical hypothesis testing** 

## **Biological Interpretation**

- 1. term enrichment analysis (KEGG, GO)
- 2. network analysis methods
- 3. structure analysis
- 4. isoform analysis
- 5. follow-up experiments

## **Experiment Design**

This section should discuss trade offs and balancing them to design an experiment. 1. constraints: Each experiment will have different constraints, which may include the number of samples needed for analysis, or desire to quantify a specific subset of proteins within a sample. 2. sample size 3. statistics 4. costs

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