COMPUTATIONAL PROTEOMICS AND METABOLOMICS

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7. Peptide Identification I – Database Search
LEARNING UNIT 7A
PEPTIDE DATABASE SEARCH

- Peptide fragmentation
- Database search concepts
Peptide Identification

Why can we identify peptides from tandem MS spectra?

- **Goal: identify sequence**
- **Tandem MS**
  - Sequence consists of the **same 20 building blocks** (amino acids)
  - CID: peptide breaks preferentially along the **backbone**
  - Peptide **fragment ions correspond to prefixes and suffixes** of the whole peptide sequences
  - Complete ion series (ladders) reveal the sequence via mass differences of adjacent fragment ions
Peptide Identification

• **Issues**
  - Spectra are incomplete – ions are missing
  - Missing information makes it very hard to reconstruct full sequence

• **Database search**
  - Not all sequences occur in a proteome – only a fraction of sequence space is used
  - Try to find those sequences that match the ions present in the spectrum

Consensus spectrum: PeptideAtlas id 829036
Product ion generation

- A peptide of length n can potentially give rise to a,b,c and x,y,z ions. This example shows the fragments that can be produced between amino acids $R_m$ and $R_{m+1}$
- This nomenclature for fragment ions was first proposed by Roepstorff and Fohlman in 1984


Steen and Mann. Nature Reviews, Molecular Cell Biology, Vol. 5 2004
Ion Series

\[
[\text{H}_2\text{N}-\text{CH-CO-}\left(\text{NH}-\text{CH-CO}\right)_x\text{NH}-\text{CH-COOH} + \text{H}]^+ \\
\]

\[
\text{H}_-(\text{HN}-\text{CH-CO})_{n-1}^-\text{NH}^-\text{CH}^+ \\
\text{R}_n \\
a_n \\
\]

\[
\text{H}_-(\text{HN}-\text{CH-CO})_{n-1}^-\text{NH}^-\text{CH-C}=\text{O}^+ \\
\text{R}_n \\
b_n \\
\]

\[
\text{H}_-(\text{HN}-\text{CH-CO})_{n-1}^-\text{NH}^-\text{CH-CO}^-\text{NH}_3 \\
\text{R}_n \\
c_n \\
\]

\[
\text{R}_n \\
\]

\[
\text{R}_n \\
x_n \\
\]

\[
\text{R}_n \\
y_n \\
\]

\[
\text{R}_n \\
z_n \\
\]

b/y ions in CID

CID fragmentation predominately produces b and y ions

Note: $y_i$ ion is also called the *sister fragment* of the $b_{n-i}$ ion and vice versa
Ion Series - Example

- For simplicity we will consider theoretical spectra for the artificial (tryptic) peptide TESTPEPTIDEK
- For singly charged ion fragments, only one of the sister fragments will be observed
Ion Series - Example

• If the same peptide was multiply charged, the charges are usually distributed across the product ions
• Tandem spectrum then usually contains both sister ions and also doubly charged product ions
Ion Series - Example

• Theoretically, one can also observes a, c, x and z ions
Ion Series - Example

- Theoretically, one also observes a, c, x and z ions
- abc and xyz ions are called backbone ions.

This spectrum contains all theoretical backbone ions of charge 1-2 (theoretically generated for TESTPEPTIDEK)
Neutral Losses

- Besides backbone ions, we also observe the precursor ions and precursor ions with **neutral losses**

- **Neutral losses** most frequently occur as
  - **water loss** ($\text{H}_2\text{O}: -18.011 \text{ Da}$) on S, T, D and E
  - **ammonia loss** ($\text{NH}_3: -17.027 \text{ Da}$) on R, K, N and Q
  - **loss of phosphoric acid** ($\text{H}_3\text{PO}_4:-98 \text{ Da}$) on S, T and Y

- Neutral losses are uncharged fragments, but result in an additional charged ion with $\text{mass}_{\text{ion}} - \text{mass}_{\text{neutral}}$

- The problem of very intense ions resulting from neutral losses of precursor ions can be overcome by triggering an additional fragmentation
Internal Fragments

- **Internal fragments** result from double backbone fragmentation. Usually, these are formed by a combination of $b$-type and $y$-type ions, and consist of five residues or less.

- **Immonium ions** are a special case of internal fragments. They are composed of a single side chain formed by a combination of $\alpha$-type and $y$-type fragmentation.
Noise in Tandem Spectra

- In addition to the various types of ions, there is also noise in tandem spectra

Blank run

One isolated peptide

With noise

Without noise

Freitas and Xu, BMC Bioinformatics. 2010, 11:436
Ion Types – Summary

• Due to different fragmentation efficacies and different response factors, fragment ions will have different intensities

• These intensities can be predicted using machine learning techniques and appropriate fragmentation models, however, most search engines do not include intensity information, but only the masses

• In general, a simple peptide search engine should consider $b$ and $y$ type ions, doubly charged $b$ and $y$ type ($b^{2+}, y^{2+}$) ions and optionally $b^{-\text{NH}_3}, y^{-\text{NH}_3}$ and $b^{-\text{H}_2\text{O}}, y^{-\text{H}_2\text{O}}$
Database Search – Overview

LC-MS/MS experiment

Experimental spectra

Fragment m/z values

Compare

Sequence db

Theoretical spectra

Theoretical fragment m/z values values from suitable peptides

Score hits

1. QRESTATDILQK 18.77
2. EIEEDSLEGLKK 14.78
3. GIEDDLMLIKK 12.63

Q9NSC5|HOME3_HUMAN Homer protein homolog 3 - Homo sapiens (Human)
MSTAREQPSTRAHVYGPDATKRNWPAGKHALT+SYFDATNYPYRISIGKAAXNSTVPTMKFTYKSYQSGWSSBANGTV YGLOFAEORQHTQAEKGPVEKAEARLARKSQSGSTLPAL GLASHQVVPPIVSAAPGEGKRFQSDADQAIQTERRXK LSEGIEGVEQVHEAFTPALSDPNKVLAGDAEAANAAAGWRR QLEAQRUAERLQRQVAELEAGAAESEVTGETEGQGGISQL EQLEALVQKTHQSFQTYXTSTGSPREAELARFERTQDQVQDL ETRENELEQLEGERLRERRERRMRVQRAQLDQVSG FELSEUREGLARLAEAAP
**Database Search – Key Steps**

1. **Extract all sequence candidates** (usually tryptic) from the database matching the precursor mass of the MS\(^2\) spectrum with a given error tolerance

2. **Generate theoretical spectrum** for each of the candidate sequences

3. **Align** the theoretical spectra to the experimental spectrum

4. **Score** the alignment

5. **Report all peptide-spectrum matches** above a certain score threshold
Step 1. Generate Candidates

- Given: Experimental spectrum $S$
- Task: Identify the correct sequence for $S$ from a given protein database

1. Define the search space for $S$ for a given mass tolerance $d$:
   - $m_{\text{prec}}$ is the mass of the precursor ion of spectrum $S$
   - From the database, extract all peptide sequences with mass $m_{\text{cand}}$ given that
     \[ |m_{\text{prec}} - m_{\text{cand}}| \leq d \]
   - This set of candidates is defined as the search space for spectrum $S$ and denoted as $\Omega_S$
Step 2: Generate Theoretical Spectra

- 1\textsuperscript{st} option: extract all masses from the MS\textsuperscript{2} spectrum
- 2\textsuperscript{nd} option: try to model fragment ion intensities
Step 3. Align Spectra

The theoretical spectrum $T$, generated from a sequence $\ p_i \in \Omega_S \ $
Step 3: Align Spectra

2. Compare theoretical spectra for all \( p_i \in \Omega_S \) to the experimental spectrum \( S \)

\[
p_1 \in \Omega_S \\
p_2 \in \Omega_S \\
\ldots \\
p_n \in \Omega_S
\]
Step 4: Scoring of peptide candidates

- There are numerous tools for the comparison of theoretical and experimental candidate peptides.
- The main difference of search engines is the implementation of the scoring schemes (resulting in differences in runtime and performance).
- However, conceptually all search engine algorithms are based on fragment ion comparison.
- In the following, we will discuss:
Real-life implementations are more complex and often require additional pre-/post-processing and depend on a large number of parameters.
LEARNING UNIT 7B
SEARCH ENGINES

- X!Tandem
- Sequest
- Other search engines
Database Search Engines

• Dozens of different database search tools are currently being used
• Common to these tools are the fundamentals describe in Learning Unit 7A
• Tools differ with respect to
  • Spectrum pre-processing
  • Scoring of peptide-spectrum matches
  • Post-processing of peptide-spectrum matches
  • Score statistics
  • Speed
• Results for the same dataset will differ between search engines!
X!Tandem

- X!Tandem
  - is a popular open-source database search engine
  - is fast
  - has been published in various versions including multiple refinements to the core algorithm sketched here (latest version: X!Tandem Sledgehammer, 2013)

Original reference
- http://www.thegpm.org/tandem/instructions.html
Find overlapping masses

To find overlapping masses, a maximal **fragment mass tolerance** window needs to be set (for ion traps this is usually 0.5 Da)
X!Tandem’s dot product

- Reduce the experimental spectrum to only those peaks that match peaks in the theoretical spectrum
- Calculate dot product ($dp$) (using ion intensities and the number of matching ions)

$$dp = \sum_{i=0}^{n} I_i P_i$$

*Intensities from experimental spectrum*  
$I_i$ ... fragment ion intensities  
*Predicted or not in theoretical spectrum*  
$P_i \in \{0, 1\}$
Survival function and e-value

- Let $x$ represent the dot product score for the experimental spectrum $S$ and the theoretical spectrum $T \in \Omega$.
- $p(x)$ is calculated from the frequency histograms (counts of PSMs per score bin).
- With $f(x)$, the number of PSMs that are given the score $x$, $p(x)$ is calculated as $p(x) = f(x)/N$ with $N$ being the total number of PSMs.

Example of a frequency histogram
The survival function, $s(x)$, for a discrete stochastic score probability distribution, $p(x)$ is defined as

$$s(x) = P(X > x) = \sum_{X > x} p(x)$$

where $P(X > x)$ is the probability to have a greater value than $x$ by random matches in a database.
Survival function and e-value

- With the survival function $s(x)$, we can calculate the E-value $e(x)$, indicating the number of PSMs that are expected to have scores of $x$ or better

$$e(x) = n s(x)$$

where $n$ is the number of sequences in $\Omega_S$

- Now, each PSM can be ranked according to $e(x)$
The hyperscore (HS) is calculated by multiplying with factorials of the number of assigned b and y ions. The use of the factorials is based on the hypergeometric distribution that is assumed for matches of product ions.
X!Tandem Hyperscore

- If $p(x)$ is now plotted as a function of their log(hyperscores), the valid PSM is much better separated from the bulk of incorrect assignments.
Distribution of “Incorrect” Hits

Adapted from Interpreting MS/MS Proteomics Results by Brian Searls
Estimate Likelihood (E-Value)

![Hyper Score vs Log(# of Matches) Diagram]

Adapted from Interpreting MS/MS Proteomics Results by Brian Searls
Estimate Likelihood (E-Value)

Hyper Score

Log (# of Matches)

0 10 20 30 40 50 60 70

Expected Number Of Random Matches

Best Hit

Adapted from Interpreting MS/MS Proteomics Results by Brian Searls
Estimate Likelihood (E-Value)

Score of 60 has 1/10 chance of occurring at random.

Adapted from Interpreting MS/MS Proteomics Results by Brian Searls.
Sequest

Sequest – Cross Correlation

- Sum all the peaks that overlap between theoretical and experimental spectrum
- This score is called cross-correlation
Sequest – Autocorrelation
Sequest – $X_{\text{corr}}$ Score

- By shifting the spectra, the assumption is that the peaks should not overlap. The spectra are displaced by $x$ Da
- The peaks that overlap upon spectra shifting are used to calculate the autocorrelation

- Sequest reports $X_{\text{corr}}$ scores

$$X_{\text{corr}} = \frac{\text{Cross}_{\text{corr}}}{\text{Average(\text{Auto}_{\text{corr}})}_{-75 \leq x \leq 75}}$$

for displacement $x$ [Da] $\in \{-75, 75\}$

Sequest – $\Delta C_n$ Score

- $X_{corr}$ scores can be calculated for every theoretical spectrum in the search space $\Omega_S$ for an experimental spectrum $S$
- Additionally to the $X_{corr}$ score, Sequest also calculates the $\Delta C_n$ score for the top scoring PSM (best $X_{corr}$)
- This score measures how good the best score is in relation to the second best

$$\Delta C_n = \frac{X_{cross 1} - X_{cross 2}}{X_{cross 1}}$$
Other Search Engines

  - Mascot is one of the most popular search engines
  - Commercial software
  - Algorithmic details have never been published
  - Mascot calculates $p$-values for all candidates in the search space and ranks the output according to these $p$-values

- Phenyx
  - Commercial software

- InsPecT
  - Very fast open-source search engine
  - Designed for the identification of posttranslational modification

- Myrimatch
  - Open source
  - Tabb et al., J Proteome Res. 6(2) 654-61. 2007 Feb
Search Settings

- OpenMS offers TOPP tools for the most common search engines
- .ini files allow to adjust the parameters
- This is an example for X! Tandem settings for analyzing LTQ-Orbitrap data
Mass Tolerance Settings

• Mass tolerance settings:
  • Easy to estimate when knowing the instrument, calibration runs
  • Precursor tolerance determines search space
    • should be stringent, but broad enough to have several entries per search space (e.g., for E-value calculation)
    • 5-10 ppm is commonly used for data acquired on well-calibrated Orbitrap instruments
  • Product (or fragment) tolerance determines the number of theoretical fragment ions that can be matched to the experimental spectrum
    • again, should be stringent, but also provide enough flexibility for statistical assessment (e.g., drawing the Poisson distribution in the OMSSA algorithm)
    • 0.5 Da is commonly used for data recorded by ion traps (e.g. LTQ)
Charge State and Missed Cleavages

**Charge state**

- Frequently, the mass spectrometer is set to only fragment features with charge $> 1$
- If you know your data is restricted to several charge states (e.g., for your mass spectrometric settings), you can save time by not looking at these

**Missed cleavages**

- Sometimes, proteases don’t cleave perfectly
- 1 or 2 missed cleavages should be allowed, but be careful since the number of missed cleavages increases your search space sizes!
Modifications

The modification settings mostly depend on the sample preparation

Fixed modifications

- **Carbamidomethylation of cysteins** is used as fixed modification in most experiments, since proteins are usually subjected to a DL-Dithiothreitol (DTT) treatment to reduce disulfide bonds built by cysteins. To protect the liberated −SH the samples are treated with iodoacetamide. This leads to a stable modification of cysteins.

\[ \text{-CH}_2\text{-SH} + \text{I-CH}_2\text{-CONH}_2 \rightarrow \text{-CH}_2\text{-S-CH}_2\text{-CONH}_2 \]

CYS residue   Iodacetamide   S-caramidomethylated CYS

- A fixed modification on amino acid X replaces the original amino acid X during database search
Modifications

The modification settings mostly depend on the sample preparation

Variable modifications

- Variable modifications should be set if you know that a subset of the amino acids are modified. Routinely oxidation of methionine should be set as variable modifications. During the electrospray ionization Met residues frequently react with the oxygen in the ionization source environment.
- Note that variable modifications are considered as additional amino acids and impact search space size drastically.

http://ionsource.com/Card/MetOx/metox.1.gif
Variable Modifications

Intuitively...

- More variable modifications should discover more peptides
- Large parts of the proteome are modified

However...

- More ‘amino acids’: increase the search space (combinatorial explosion)
- Loss in sensitivity
- Variable modifications need to carefully chosen
LEARNING UNIT 7C
FDR ESTIMATION
Database Settings

• The database should contain all protein sequences that are potentially in the sample (e.g., all human proteins of your looking at proteomics data from human cell lines)

• From the database and the enzyme’s ‘cutting rule’ settings, the peptide candidates are calculated

• Apart from the expected proteins, the database should also contain common contaminants, such as trypsin (or other enzymes), keratins or BSA (bovine serum albumin, often used for instrument calibration)

• Databases can also be designed in a way to give an intuitive idea of false discovery rates -> target/decoy databases
Target-Decoy Databases

- Take the original protein sequences (target sequences) and reverse, pseudo-reverse, randomize or shuffle these sequences to create **decoy sequences**
- Spectra are either searched twice (first against target, then decoy database) or against the concatenated database (target + decoy)
- Decoy sequences are random sequences and should not be present in the sample
- PSMs against decoy proteins have to be **false positive identifications**

**Note**
- The decoy database design should provide equal numbers of decoy peptides as there are target peptides per search space (with randomized sequences this is hard to control)
- Ideally one should avoid large overlap between target and decoy peptides
## Target-Decoy Approach

<table>
<thead>
<tr>
<th>Peptide identification</th>
<th>Search engine score</th>
<th>TARGET/DECOY</th>
</tr>
</thead>
<tbody>
<tr>
<td>LCEVEEGKDKEVDVK</td>
<td>$s_1$</td>
<td>TARGET</td>
</tr>
<tr>
<td>YTAQVDAEEKEDVK</td>
<td>$s_2$</td>
<td>TARGET</td>
</tr>
<tr>
<td>IVADKDYSVTANSK</td>
<td>$s_3$</td>
<td>TARGET</td>
</tr>
<tr>
<td>TGIIEIKK</td>
<td>$s_4$</td>
<td>TARGET</td>
</tr>
<tr>
<td>DLGEEHFK</td>
<td>$s_5$</td>
<td>TARGET</td>
</tr>
<tr>
<td>TASSDTSEELNSQDSPK</td>
<td>$s_6$</td>
<td>DECOY</td>
</tr>
<tr>
<td>GAGGENEPPAAPEPR</td>
<td>$s_7$</td>
<td>TARGET</td>
</tr>
<tr>
<td>IKDPDAAKPEDWDDDR</td>
<td>$s_8$</td>
<td>TARGET</td>
</tr>
<tr>
<td>VDEVGGEALGR</td>
<td>$s_9$</td>
<td>TARGET</td>
</tr>
<tr>
<td>SEEQLKEEGIEYK</td>
<td>$s_{10}$</td>
<td>DECOY</td>
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<tr>
<td>LHVDPENFK</td>
<td>$s_{11}$</td>
<td>TARGET</td>
</tr>
<tr>
<td>FSTVAGESGSADTVRDP</td>
<td>$s_{12}$</td>
<td>TARGET</td>
</tr>
<tr>
<td>AEEDEILNR</td>
<td>$s_{13}$</td>
<td>DECOY</td>
</tr>
</tbody>
</table>

- PSMs are sorted by (deteriorating) score
- As the score gets worse, the likelihood of finding a decoy hit increases, likelihood for target hit decreases
- By choosing an appropriate score threshold, one can ensure a given false-discovery rate (FDR)
PSM Score Distribution

FPR = B/(B' + B)

FDR = B/A = \left( \sum_{i=1}^{A} PEP_i \right)/A

PEP = b/a
Target-Decoy Approach

Design decoy sequences

Although different decoy database designs produce very similar results, the most frequently used approaches are the reversed and pseudo-reversed decoy databases.

Calculation of FDRs

• General equation for FDR calculation (see statistics lecture)

\[ FDR = \frac{FP}{FP+TP} \]

There are two ways how FDRs are calculated based on target-decoy search results:

• Käll et al. suggest \(^1\) (Käll et al., Proteome Res. 2008, 7, 29-34)

\[ FDR = \frac{\#\text{decoy}}{\#\text{target}} \]

• Zhang et al. suggest \(^2\) (Zhang et al., J Proteome Res 2007;6(9):3549-3557)

\[ FDR = \frac{2\#\text{decoy}}{\#\text{target}+\#\text{decoy}} \]

• OpenMS tool **FalseDiscoveryRate** uses the Käll metrics
LEARNING UNIT 7D
CONSENSUS IDENTIFICATION

• Overlap between search engines
• Consensus identification idea
• Scaffold
• OpenMS Consensus ID
Comparison of Search Engines

Simple experiment
- 18 protein mix, digest, measure
- Same dataset searched with three different search engines
- Equivalent search parameters

Result
- Overlap between search engines is rather limited
- Each search engine finds (correct) candidates none of the others finds

Idea
- Combine results from multiple search engines (consensus identification)

Searle et al., Journal of Proteome Research. 2008, 7, 245-253
Multiple search engines

- Majority voting
  - Reliability $\uparrow$ sensitivity $\downarrow$
- All peptide IDs
  - Reliability $\downarrow$ sensitivity $\uparrow$

- Combine search engine scores
  1. Scores are inherently different
  2. Different number of peptide candidates

- Combination approaches
  - Scaffold: Searle et al., J Proteome Res. 2008, 7, 245-253
Scaffold

Scaffold integrates search results from Sequest, Mascot and X!Tandem

1. Use mixture models to normalize different scores to probabilities
2. Calculate agreement score for each PSM across all search engines

\[ D = \text{PSM (Peptide spectrum matching)} \]

\[ D_{i,j} = \text{PSM: spectrum } i \text{ to peptide } j \]

p = probabilities for correct assignment (from mixture model)

\[
A_{i,j,k} = \sum_{k' \neq k} \begin{cases} 
0.05 \leq p(+|D_{i,j,k'}) < 0.5 & 0.0 \bigg| 0.05 \leq p(+|D_{i,j,k'}) < 0.5 & 0.5 \bigg| 1.0 
0.5 \leq p(+|D_{i,j,k'}) & 0.05 \leq p(+|D_{i,j,k'}) < 0.5 & 0.5 \bigg| 1.0 
0.5 \leq p(+|D_{i,j,k'}) & 0.5 \leq p(+|D_{i,j,k'}) \end{cases}
\]

\[ p(+|D, A) = \frac{p(A|+) p(+|D)}{p(A|+) p(+|D) + p(A|-) p(-|D)} \]
Scaffold Performance

We did not discuss the naïve max.

\[ \text{max} \left[ p(+ | D, A) \right] \]

**Graph:**
- **naïve max**
- **SEQUQUEST**
- **Mascot**
- **X! Tandem**

**Axes:**
- **Number of Identified Spectra**
- **Percent Error Rate**
ConsensusID integrates search results from OMSSA, Mascot and X!Tandem

1. Use mixture models to normalize different scorings to probabilities

Nahnsen et al., J Proteome Res. 2011 Aug 5;10(8):3332-43
## ConsensusID – Mixture Modeling

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Nahlsen et al., J Proteome Res. 2011 Aug 5;10(8):3332-43
### ConsensusID – Similarity Scoring

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Nahnsen et al., J Proteome Res. 2011 Aug 5;10(8):3332-43
ConsensusID - Consensus Score

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ConsensusID (p1) = $\frac{s_1(p_1) + \alpha s_2(p_i) + \beta s_3(p_j)}{(1 + \alpha + \beta)^2}$

ConsensusID (QRESTATDILQK) = $\frac{0.54 + 0.3 \cdot 0.96 + 1 \cdot 0.99}{(1 + 0.3 + 1)^2}$ = 0.34

Nahnsen et al., J Proteome Res. 2011 Aug 5;10(8):3332-43
ConsensusID Performance

LTQ-Orbitrap - high accuracy

error rates = false discovery rates

Nahnsen et al., J Proteome Res. 2011 Aug 5;10(8):3332-43
Materials

• Online Materials
  • Learning Unit 3B (statistics for FDR)
  • Learning Unit 7A, B, C, D

• Slides on peptide ID by Brian Searle
  • https://proteome-software.wikispaces.com/file/view/interpreting-MS-MS-proteomics-results.ppt
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