Hexicon 2
Quick start guide and user manual

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1 Quickstart

This section is for the impatient who want to see what Hexicon 2 does to their data. We shall assume that you have a complete and functional copy of Hexicon 2 binaries as well as the required runtime libraries for your operating system (cf. section 2.1). Furthermore, you have all data at hand, know your experimental parameters and all you want to do is get results – fast. Be warned that some tinkering with parameter settings and downstream postprocessing are required for optimal results.

1.1 Prerequisites

Hexicon 2 is a reference-based bottom-up workflow which can run using one reference LC-MS map, at least one map of deuterated protein and the corresponding protein sequences. More formally, the files you need are:

- Reference dataset, one mzXML file per LC-MS map, line spectra
- Deuterated data, one mzXML file per LC-MS map (at least one), line spectra
- Protein sequence of your experimental construct
- Protein sequences of all significant other components in the mixture (if present)

While most parameter settings of Hexicon 2 are only needed for fine-tuning of the workflow, it is essential that you know some parameters concerning data acquisition. This includes:

- Instrument resolution (FWHM)
- Instrument calibration accuracy
- Retention time range during which peptides elute

1.2 Walkthrough

Data Import

Hexicon 2 opens with the data import window (Figure 1). Hit Add to load your LC-MS maps. After selecting a map in mzXML format, a dialog will pop up and ask you for incubation time and group number. The group number is used to identify replicates: maps with the same incubation time and group number will be treated as experimental replicates. Enter 0 as incubation time and group number of the reference.

For subsequent deuterated datasets, enter the respective D$_2$O incubation time in seconds and assign the same (arbitrary) group number to each set of replicate maps. The example in Figure 1 shows two deuterated maps which constitute replicates of the 15 sec incubation time point, hence they belong to the same group.

If you have an appropriately formatted MS/MS search report (cf. section 2.2), it is strongly recommended that you provide Hexicon 2 with this information. A protein sequence in plain text
format always needs to be specified. If you used a 100% labeling control, e.g., by labeling in denaturing buffer, load that dataset and enter an arbitrary group number and time point higher than for your longest-incubated actual sample. Check the box next to **Use last time point as 100% D reference**. When the location of all datasets has been specified, proceed to the protein mixture filter by clicking **Next**.

**Tip**

For optimal performance, some tinkering with parameter settings may be required. For your convenience and for better reproducibility, the locations of the input files can be saved and restored using **Save** and **Open**, respectively.
Protein Mixture Filter

This step is only required if your experiment contains nonnegligible amounts of more than one protein. If this is not the case, proceed to the extraction parameters with Next.

Hexicon 2 is designed for the analysis of one protein of interest at a time. If other proteins are present in the mixture at significant amounts (such that peptides originating from these proteins are detectable in the spectra), you need to provide their sequences as background set. Do this with Add for each protein other than the protein of interest.

The optional reference map which can be loaded at the bottom of this window is currently not used (build 12 JUN 2014) as it provided no measurable improvement of sensitivity in our test data.

Parameter Settings

See section 2.4 for a detailed description of the available parameters. Only a few of these values must be changed for Hexicon to produce useful data given that you are using a common LC-MS experimental setup:

The **Retention Time Range** is crucial for sensitive feature extraction. It is highly discouraged to run Hexicon on the entire map. Much better performance can be achieved by entering the time range at which your peptides elute from the chromatographic column. Make sure that the time window not only accommodates the reference peptides but also the deuterated ones and do not worry about adding a minute or so at each side of the window.

An estimate of the actual instrument Resolution (FWHM) ($\pm 10\%$) is required for Hexicon to get an estimate of the attainable mass precision. More importantly, if your detector tends to drift over the course of an experiment, you should calibrate it regularly (at least before the first and after the last injection, to know which errors to expect) and record the mass deviation. Enter the largest relative mass deviation ($\frac{\Delta mz}{mz}$ in ppm) that you expect for any of the analyzed measurements as **Calibration Error (ppm)**.

**Tip**

If calibration logs are not available and you cannot reproduce calibration errors experimentally, have a look at the precursor mass values reported for MS/MS sequenced peptides and compare them to theoretically expected masses. This corresponds to the delta M field of MASCOT reports.

The **Export Settings** are intuitive and should be adjusted. In addition to the specified output file, Hexicon will create a number of other files in the same directory, including a runtime log (Hexicon.log) and a subdirectory (deuteration), in which measured deuteration distributions for each identified peptide are stored. Is is therefore recommended to use one dedicated directory for the result output of each Hexicon run.
**Tip**

Save parameter settings in the Hexicon result output directory for documentation and reproducibility. The resulting .hxpar file is a human-readable plain text file containing all parameter settings.

If all parameters are set, proceed to the **Run** tab with **Next**.

**Run Hexicon**

If no error message is displayed in the status monitor, start Hexicon with **Start Hexicon Workflow**. An approximate progress bar and the status monitor showing the contents of the Hexicon.log file as it is created keep you informed about the progress and provide you with key statistics. When the run has finished, a csv file containing one feature per line, as well as a YAML file will be created. The YAML file can be read by hxviewer, Hexicon’s result browser, which is the recommended way of postprocessing and viewing data.
2 Hexicon Core

This section describes the core of the Hexicon 2 workflow, upstream requirements, parametrization, the structure of the output as well as recommendations for quality control and downstream processing.

2.1 Prerequisites

You have most likely received Hexicon 2 binaries in a zipped folder. The archive contains the executables hexicon.exe and hxviewer.exe, the libraries ms++.dll, hxcore.dll, sashimi.dll, yamlcpp.dll, zlib.dll as well as Qt libraries (QtXml4.dll, QtSvg4.dll, QtGui4.dll, QtCore4.dll, imageformats/qsvg4.dll). The text file pepsintable.txt contains protease cleavage score definitions (cf. section 2.5).

It may be necessary to install Microsoft Visual Studio 2010 runtime libraries available free of charge from Microsoft.

2.2 Input Data

LC-MS Maps

Hexicon 2 was designed for the analysis of high-resolution liquid chromatography mass spectrometry data generated from high-resolution TOF or ion trap based instruments. Since the amount of data generated by such devices can be enormous and most of them provide sophisticated methods for peak picking, we leave this task to upstream processing by the user and start with line spectra.

The open *mzXML* format is currently the most widely used and most flexible open data format for mass spectrometry data. We are aware of the shortcomings of XML-based formats for data storage and plan to provide support for other data formats as new open standards are emerging.

In order to run Hexicon 2, you will need exactly one reference map containing undeuterated peptides, at least one map containing deuterated peptides, and the protein sequence of the construct used in your experiment.

Hexicon 2 is geared towards the analysis of continuous labeling experiments, i.e., the mass difference over D$_2$O incubation time is analyzed. Therefore, each LC-MS map needs one associated time point. Maps with the same time point and group number are treated as replicates, i.e. extracted deuteration values are averaged.

Relative deuteration centroid values can be corrected for back-exchange using a 100 % deuterated control sample measured under identical LC-MS conditions. You can provide such measurements as a set of additional maps. Give these maps an arbitrary group number and time point that’s higher than the group numbers and time points of all other maps, then check the box next to **Use last time point as 100 % D reference**.
MS-MS Report

Loading a MASCOT search report will greatly improve the quality of sequence assignment to peptides contained therein. Furthermore, it helps resolving ambiguous sequence assignments. The format is historically a copy of the peptide summary browser display, hence Hexicon 2 looks for a whitespace or tab-separated file containing the following fields:

- Query ID
- Observed m/z
- Mr (measured)
- Mr (theoretical)
- delta m
- missed cleavages
- Ion Score
- E-value
- Rank
- Unique (letter U to denote uniqueness)
- Peptide sequence

Only the values printed in **boldface** are used, hence dummy values can be inserted in other fields to create Hexicon 2 compatible MASCOT reports. Sequences must be provided in the format \( \text{X.YYYYY.X} \) with \( \text{YYYYY} \) being the peptide sequence. Residues flanking the cleavage site are not used in the current release (12 JUN 2014), however this behavior may change. We are planning on adding support for a simpler format for the specification of MS\(^2\) confirmed peptides in the next release.

Protein Sequences

The **Protein Sequence** field is mandatory. Load a plain text file containing the protein sequence corresponding to your experimental construct in one-letter code. Only standard proteinogenic amino acids are allowed, however you may define some exotic amino acids as post translational modifications, given that they contain only C,H,\(^2\)H,N,O,P,S and I atoms. Other atoms can likely be included, however some elements, e.g., Selenium, throw NITPICK’s Averagine-based isotope pattern model off.

It may be useful to add a **Reference Sequence** if you want to compare constructs of different length, containing mutations or just for the sake of using standardized sequence numbering. Positions of found peptides will be mapped to their reference position (using ungapped sequence
alignment) and the reference sequence will be shown instead of the construct sequence when generating protein-level graphical output, i.e., peptide maps and coverage histograms. Note that all graphical displays of Hexicon 2 will only allow comparing peptides of identical sequence regardless of the reference sequence.

**Post Translational Modifications**

Hexicon 2 allows definition of simple fixed post translational modifications. Once the protein sequence has been loaded, hit the **Modifications** button to open the corresponding dialog (Figure 2). You will need to define the modifications you want to apply to your protein in a YAML file which is structured as follows:

```yaml
elements map - must be the first node of the document and contain a map of all elements that you want to define and their mass number. The mass number is only used for identification of the element in our stoichiometry table - isotope distributions are pulled from an internal library.

Example:

```
elements:
  H: 1
  D: 2
  C: 12
  N: 14
  O: 16
  P: 30
  S: 32
  I: 53
```

---

![Figure 2: Hexicon 2 Post translational modifications dialog.](image)
This block can be followed by any number of nodes, each containing following tags:

- **modification**: the name of the modification;
- **short**: a short identifier for display in the peptide sequence;
- **composition**: a map of the elemental composition which will be added to the modified amino acid’s stoichiometry (e.g., H: -2 means removal of two hydrogen atoms);
- **applyTo**: List containing one-letter code of amino acids to which the defined modification can be applied. Example:

```plaintext
modification: Phosphorylation
short: p
composition:
   H: 2
   O: 3
   P: 1
applyTo:
   - Y
   - S
   - T
---
modification: Phosphopantetheinylation
short: ppt
composition:
   H: 21
   O: 6
   P: 1
   C: 11
   N: 2
   S: 1
applyTo:
   - S
```

Once modifications have been loaded, use the mouse to highlight the parts of your protein sequence which you want modified, pick the corresponding modification from the list of available modifications and hit `>>` to apply it.

<table>
<thead>
<tr>
<th>Tip</th>
</tr>
</thead>
<tbody>
<tr>
<td>Modifications can only include the elements listed in the above example since isotope patterns are only defined for these. Modifications with an elemental composition (more precisely: isotope distribution) that strongly differs from the isotope pattern of a peptide containing only standard amino acids may not be correctly identified by our NITPICK feature detection algorithm. If your modified peptide made it through feature detection, full modeling of the isotope pattern is carried out in subsequent steps such that accuracy is no different from unmodified peptides.</td>
</tr>
</tbody>
</table>
2.3 Filters

Background Protein Filter

Hexicon 2 can only analyze peptides corresponding to one protein at a time (referred to as protein of interest). If your experiment contains other proteins (background proteins) in nonnegligible amounts, there is a good chance of falsely assigning a sequence from the protein of interest to a peptide derived from a background protein. In order to avoid this, Hexicon 2 allows you to define the sequences of all background proteins in the filters tab. Import one plain text file for each background protein in your experimental mixture. Peptides that match a background sequence better than the protein of interest will be removed from the analysis after being used for internal mass calibration and false assignment estimation.

In some cases, such filtering may be too strict and peptides of interest may be falsely assigned to background sequences. If this is the case, you can load a reference map in mzXML format, containing only the protein of interest measured under identical LC-MS conditions. Peptides matching a background sequence but found in the reference map will be rescued from the filter. Note: This feature has been disabled in the current release (12 JUN 2014) as it provided no measurable performance gain in our benchmarking studies.

2.4 Parameters

Hexicon 2 provides a number of customizable parameters which can be used to tune the sensitivity of the analysis and to adjust it to your experimental conditions.

Peptide / LC Settings

NITPICK feature detection is applied in a divide-and-conquer scheme: feature detection is carried out in each scan independently and detected features with similar mass in subsequent scans are merged. The fields Minimum / Maximum # Scans let you define in how many subsequent scans a peptide has has to be detected by NITPICK in order to be carried into further analysis. The maximum scan number is rarely hit but if you have persistent contaminations consistently detected by NITPICK, this will remove them. The Maximum Gap Time setting allows you to set a tolerance window in which a peptide need not be detected and still be merged with a previous peptide signal. It is advisable to increase this setting when you notice fragmentation of your peptides, i.e., you get a large number of peptides with the same charge that are detected in more than one contiguous region of the LC-MS map (another reason for this to happen is too low mass precision tolerance in the FWHM setting).

Hexicon 2 will try to assign peptide sequences to any detected feature regardless of MS² identification. For this purpose, exhaustive in-silico digestion is used to procure all peptides within a given size range. Making the Peptide Length Range unnecessarily large will adversely affect runtime and sequence assignment specificity.
Tip

A good way to determine a suitable peptide length range for your experiment is to have a look at the peptides identified by MS² or to do a discovery run with Hexicon 2 using a large range (e.g. 5/45) and then to narrow down the range based on the results.

**MS Settings**

Since we are dealing with line spectra, there is no good way for Hexicon 2 to derive peak width from input data. A good estimate of the scan Resolution (FWHM@400) is therefore required for Hexicon 2 to estimate the attainable mass precision. A value of 40000 FWHM roughly translates to a precision of 6 ppm. Using this setting, features from subsequent scans with a relative mass difference $\Delta m/m$ of more than 6 ppm would be treated as two separate entities.

The **Calibration Error** parameter represents the accuracy of your measurement, i.e., how much measured mass deviates from actual mass and how much two measurements of the same peptide are allowed to differ in different maps. It is safe to set a fairly large value here since it is only used as worst-case value and empirically narrowed down during the analysis. If you feel that sequence assignment or chromatogram alignment perform poorly, try altering this parameter in 10 % intervals up or down.

The **Noise Quantile** defines the peak intensity quantile that is safe to be considered as noise. It is used together with the SNR parameters (Advanced Settings) to pre-filter the spectrum. If your data is heavily pre-filtered and no noise is present, set a very low value. Low values for this parameter increase sensitivity, false discovery and mostly runtime.

**m/z Range** and **Charge State Range** are self-explanatory. Having a charge state range that specifically matches your dataset will greatly accelerate feature detection and reduce false positive rates. We strongly recommend using **Positive Ionization** for all experiments. Negative ionization is incompatible with most current HDX protocols and was never tested in Hexicon.

**Export Settings**

There are several filters that you can apply for data export into the CSV file. The YAML file for processing using hxviewer will not be filtered. Depending on your ionization settings, most peptides will produce more than one charge state which can be detected by Hexicon 2. Finding multiple charge states with similar deuteration increases the confidence in data extraction. It may therefore be useful to discard results which have **only one charge state**, however it is not recommended. For a quick discovery analysis, you may want to discard peptides with ambiguous sequence assignment. Such assignments occur when multiple theoretical peptide sequences match the extracted mass within the attainable mass precision. Hexicon 2 provides several means to resolve such sequence conflicts and to find the most likely correct assignment, hence discarding ambiguous assignments up front is discouraged. As there are currently no means to inspect individual replicates, we recommend discarding features with inconsistent replicates. This filter will remove a peptide if the deuteration centroid standard deviation exceeds a certain percentage of its absolute value. This centroid deviation cutoff is set to 20 % by default. If you get a very small number of results in the csv file but not in the YAML file, it
might be worth checking your filters and looking for a bad replicate.

Hexicon lets you **Separately Export deuteration distributions** or a list of ambiguous peptides. The deuteration distributions will be exported into a separate directory with one csv file per feature. It will contain deuteration state abundance distributions for each replicate. The list of ambiguous peptides is a useful shortcut to defining peptides which you may want to have re-sequenced by MS$^2$. Both the deuteration subdirectory and the list of ambiguous peptides will be saved in the same directory as the main result list. This is also where the YAML file will be saved.

**Advanced Settings**

The first section deals with feature detection parameters. A running median will be computed for each scan to define a background noise level. Each peak is then smoothed using a **Gaussian Filter** and compared against the background noise. The width of this filter determines whether you emphasize local peaks or broad clusters to define which peaks lie above the noise level. The two SNR parameters determine the intensity ratio of any *raw (unsmoothed)* peak over the background and its *smoothed* counterpart over the background required to pass the filter. Only if both smoothed and raw peak over background intensities exceed the set threshold values, the peak is not treated as noise. These two SNR parameters are the main determinants of feature detection sensitivity in Hexicon 2.

The runtime of NITPICK increases quadratically with model size, hence Hexicon 2 attempts to fit models with independent (i.e., non-overlapping) isotope patterns separately. Busy spectra are often highly overlapping and no split points can be found. In such cases, the **Model Splitter** parameter determines the number of models after which Hexicon 2 forces the model set to split in order to keep runtime low. We do our best to stitch split models together in a sensitive way but the fits are subobtimal at such breakpoints. If you notice that feature detection is choking, try reducing this setting. Better results are obtained with higher numbers here but significant slowdown of feature extraction was observed at about 200 models.

The chromatogram alignment implemented in Hexicon 2 estimates two alignment tolerance terms based on the data and thereby already extends the search space for deuterated peptides generously. If you feel, however, that alignment fails, you may try to manually **extend the alignment window** for deuterated peptides.

<table>
<thead>
<tr>
<th>Tip</th>
</tr>
</thead>
<tbody>
<tr>
<td>If you suspect that the chromatogram alignment is not doing a good job, try plotting reference retention times versus retention times of deuterated peptides (both listed in the csv file). Large scatter will usually indicate alignment problems.</td>
</tr>
</tbody>
</table>

**Saving Parameters**

Parameters are not automatically saved and cannot be retrieved once Hexicon 2 has started processing data! Please save your parameter settings file for reproducibility and convenience. We
recommend using one directory per Hexicon 2 analysis run where you can also store the settings used to generated the results contained therein.

2.5 Other Customizations

Protease Specificity Scores

Hexicon 2 assigns cleavage scores based on cleavage probabilities empirically determined by Hamuro and colleagues. Table 2 of the corresponding publication was adapted (W-W cleavage was not defined originally because the sequence WW had not been observed in the study) and reformatted to match a shape which can be read by Hexicon 2: In a whitespace-separated text file, $N$ fields of the first line define the one-letter amino acid alphabet $A$ (typically 20 letters). It follows a $N \times N$ square matrix of nonnegative real-valued cleavage probabilities $[0,1]$ where the value at position $i, j$ ($i$ denoting the row, $j$ the column) stands for cleavage between $A[j]$ (in $P1$) and $A[i]$ (in $P1'$). Note the unconventional column-first notation which was kept for compliance with Hamuro et al..

Consider the following example file defining an arbitrary four-letter alphabet:

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>C</th>
<th>D</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.06</td>
<td>0.18</td>
<td>0.07</td>
<td>0.2</td>
</tr>
<tr>
<td>2</td>
<td>0.15</td>
<td>0.8</td>
<td>0.1</td>
<td>0.3</td>
</tr>
<tr>
<td>3</td>
<td>0.2</td>
<td>0.31</td>
<td>0.01</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>0.21</td>
<td>0.83</td>
<td>0.03</td>
<td>0.08</td>
</tr>
</tbody>
</table>

Examples for cleavage scores would be $AAC_A\cdot DEACED_A$ (C/D): 0.31 or $AACDEACED_A\cdot A$ (D/A): 0.07. Internal peptides result from two cleavage events, hence a compound cleavage score must be found. Hexicon 2 assigns an internal peptide cleavage score

$$s = \sqrt{s_1 s_2}$$

with $s_1$ and $s_2$ being the scores of the N- and C-terminal individual cleavage sites, respectively. Accordingly, the subsequence $DEACED$ from the above example (denoted $C\cdot DEACED\cdot A$ using MASCOT convention) would receive a score of 0.15.

Specificities are read from the plain text file "pepsintable.txt" which must be in the search path of Hexicon 2. In the Windows release, this would be the same directory as the hexicon.exe binary.

2.6 Running Hexicon 2

Hexicon 2 will produce a log file that contains much useful information about the state of data processing. The log file is shown in the Run tab of Hexicon 2 and saved in the directory of run output. Should Hexicon 2 quit unexpectedly without an error message, please have a look at the log file which will contain debug output.

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1Hamuro Y. et al., Rapid Commun Mass Spectrom. 2008; 22(7):1041-6
3 Hexicon 2 Result Browser

Hexicon 2 writes its results in YAML format which can be read by HXViewer, a graphical tool for postprocessing and visualization of Hexicon 2 results.

3.1 User Interface

HXViewer has a split graphical user interface (GUI) which contains file and result management in the left column and a multiple document interface (MDI) area for various tasks related to visualization as well as interactive manipulation (Figure 3). Selecting the graphical representation of a peptide in any of the views will select and highlight the peptide in the tree view on the left. Modifications to either representation of the data will update the underlying model and affect any other data displays.

![Figure 3: Hexicon 2 Result Browser User Interface](image)

3.2 File Handling

File Loading

You can load one or more YAML files containing Hexicon 2 results by drag&drop from your filesystem browser into the tree view on the left, by clicking on the green + symbol above that area or by using the menu item **File > Open**. Since YAML is a text-based format, loading of data may take a while and happens in the background while you can start naming your datasets or
work on other, previously loaded data. Completely loaded datasets will appear in the dropdown menu on the left from where you can select the *active dataset* to be displayed in the tree view.

**File Unloading**

Unload an opened dataset to free the memory it occupies by clicking on the red – button while it is active. This will only succeed if no other views (graphical displays) of that dataset are open. **Note:** Changes you made to the dataset will be lost unless you explicitly save it!

**File Saving and Export**

Your work is not saved unless you explicitly do so. Since HXViewer can only read YAML files, you should save your progress. We recommend saving to a new YAML file using the **File > Save as** menu item such that the original Hexicon 2 output is not lost. Peptides marked as disabled will be saved as such and can be restored using HXViewer. **Important:** Only the active dataset (in the dropdown menu on the left - not the graphics view) will be saved. Make sure you save work in all your datasets before closing HXViewer.

If you wish to create a csv file similar to the output originally generated by Hexicon 2, you can use the menu item **File > Export to csv**. The dialog will ask you whether you wish to create a separate directory for deuteration files. Please note that answering **Yes** will create a deuteration directory in the same location as the exported csv file. If this happens to be the directory of the original Hexicon 2 output, it will be overwritten.

**3.3 Dataset Navigation**

**Selecting the Active Dataset**

Use the dropdown menu on the left to choose the active dataset. It will be displayed in the tree view below and all filters or visualizations will be applied to this dataset. You can change the active dataset while there are still graphical displays of it, however interactive manipulation is only possible to the active dataset. **It is not** required to save your work before changing the active dataset.

**The Tree View**

The tree view on the left side of HXViewer presents rich information about the results of a Hexicon 2 run. It groups the results by peptide and lists all peptides that were retrieved at least once in each time point (i.e., have complete time series). Right-click the title bar to change how peptides are sorted. The default is by position in the protein sequence. Expanding a peptide will show all found ions (charge states) which you can view in further detail by expanding. Ions displayed in **red font** have multiple sequence assignments which you can view by expanding their **other assignment(s)** field. Right-click or double-click on the other assignment to jump to it in the tree view. Ions displayed in **boldface** are confirmed by MS2 sequencing. Peptides that contain at least one ambiguous or **confirmed** sequence are displayed in **red** or **bold**, respectively. Double click on an ion to open a graphical display showing its deuterium incorporation over
time or on a peptide to show that display for all related ions. Other peptides or ions can be added to the plot by dragging tree items and dropping them over an active plot. This can be done across datasets.

Right-click on an ion to disable it or on a peptide to disable all related ions. Disabled items will be removed from any statistics or graphical displays, they will not be listed as sequence conflicts and not be exported to CSV. When saving the active dataset to a YAML file, disabled state is preserved. Right-click on a disabled entity to re-enable it.

**MDI Tabs**

Graphical displays opened by a dataset will be grouped in one tab of the MDI area in the right part of the viewport. While this helps you keep your work structured, closing a tab provides a convenient way of destroying all views on a dataset and thereby making it ready for unloading (unless there are open multi-dataset views open in other tabs).

### 3.4 Filtering and Processing

HXViewer provides two filter operations that we routine use as first-pass quality control: **Filter by intensity** and **Resolve sequence conflicts**. If you plan on applying both filters, we recommend first applying the intensity filter and then proceeding to conflict resolution. Filters will operate on the active dataset only. The intensity filter looks for ions with large deviation in intensity extracted from different maps. This is usually an indicator for either poor extraction in some datasets or for mis-alignment, i.e., assignment of a deuterated signal to a noncognate reference ion. It has two passes: The filter will remove all features which exceed a relative intensity standard deviation larger than the first cutoff set regardless of total intensity in the first pass. The second pass removes all ions with a given standard deviation and an absolute intensity lower than a specified quantile. This filter is of limited utility if you know that due to experimental conditions, your samples show large variation in intensity.

The conflict resolution filter inspects all ions with multiple sequence assignments and attempts to determine, solely based on sequence, if any of the suggested sequences is more likely to be correct than the others. It allows the user to specify residues after which cleavage is not allowed to occur. Furthermore, cleavage scores computed by Hexicon 2 can be compared. If one sequence is significantly more likely to be produced by the protease than another, it will be accepted as correct sequence. Check the box next to **Extend MS/MS to related ions** if you want MS² confirmation of an ion also to be applied to related ions and thus all related sequences to be preferred over any conflicting sequences.

The **Set all peptides** menu contains two elements: **Enable** and **Disable**. This allows to (re-)set your peptide selection quickly, e.g., if you want to consider only a small number of manually selected peptides. Please note that hxviewer does not have an undo function, hence you should save your dataset to preserve the state of any manual processing you may have done prior to enabling or disabling all peptides.
3.5 Graphical Representations

Coverage Statistics

Activate the dataset of interest and choose the menu option Visualization > Coverage statistics to view a summary of general statistics about that dataset (Figure 4). Note that the recovery statistics in this window do not match up with what is printed in the Hexicon 2 log file since recoveries in the log file do not check for presence of a unique sequence for a particular ion. The histogram in the upper part of the view shows the sequence assignment mass error. Since Hexicon 2 performs internal mass-recalibration, the histogram should be zero-centered and its width should not exceed the attainable mass precision of your mass spectrometer. The bar plot in the bottom part of the view shows the number of features per backbone amide. It takes into account that the N-terminal amide back-exchanges rapidly under quench conditions and cannot be used to extract HDX information. This figure is interactive and supports dragging as well as mouse wheel zooming. Right-click on the figure to open the graphics export dialog (cf. section 3.6).

![Sequence Coverage](image)

**Figure 4:** HXViewer Result Statistics View

Peptide Map View

The peptide map view can either display deuteration of all peptides in a single dataset (dynamics mode, Figure 5) or compare deuteration differences between datasets (difference mode). Open
the peptide map view through the menu item **Visualization > Peptide map**. Choosing only a reference dataset will open the dynamics view which shows deuterium incorporation of each peptide color-coded from blue (low) through green to red (high). The scale begins at zero and the **Dynamic Range** parameter determines the upper limit of the scale. You can adjust the number of colors to obtain a more coarse or fine scale of color codings.

Choosing two different datasets will open a deuteration difference view which shows the deuter-

![Figure 5: HXViewer Peptide Map View](image)

ation difference of the target dataset with respect to the reference on a blue-gray-red color scale running from the negative end of the dynamic range through zero to the positive end of the dynamic range. Deuteration differences between two peptides take into account the standard deviation of deuterium incorporation into the two displayed peptides, hence peptides with large measurement uncertainty will appear less significantly blue or red than accurately measured peptides.

Each box in the plot represents one peptide which may contain multiple charge states that are combined to one compound deuteration value. Each box is split horizontally into a number of segments which correspond to D₂O incubation time points. Therefore, the different colors in a box, bottom-up, indicate deuteration values or differences at each of the incubation time points. A white diamond in a peptide box indicates MS² confirmation of at least one ion related to that peptide.

The peptide map view is interactive and supports drag and mousewheel zoom operations. Left-clicking a peptide will select the corresponding item in the tree view, right-clicking allows you to disable a peptide or to view its time series. If you are in difference mode, you will have the option to disable either the reference, the target or both peptides. In this case, the time series will show both peptides’ deuterium incorporation over incubation time.

Click the **Secondary Structure** icon to load a comma-separated secondary structure annotation which can be displayed above the protein sequence. Hexicon2 reads the following

Click the Export icon to open the graphics export dialog (cf. section 3.6).

Time Series View

The time series view shows deuterium incorporation into one or more peptides over D₂O incubation time. In order to open a time series view, double-click on an ion in the tree view. Double clicking on a peptide will display all ions associated with the sequence. Each time a new item is added to the time series view, you will be asked to specify a color. When a peptide with multiple charge states is loaded, each charge state will be plotted as an individual line, the median deuteration will be plotted as separate bold line and the deuteration range spanned by all related ions will be shaded.

You can add further peptides to the time series view by dragging them from the tree view into the plot area. HXViewer will try to construct a logical sequence from the peptides present in a time series view such that loading one peptide from a dataset will allow you to browse through all peptides of that dataset using either the scrollbar below the plot area or the mouse wheel while over the plot area. When corresponding peptides from two or more datasets with identical (reference) sequence are loaded, HXViewer will let you scroll through all peptides present in all datasets (intersection mode) or through all peptides present in any of the loaded datasets (union mode).

Click the distribution button to toggle the display of deuteration distribution estimates. If multiple peptides are loaded, each peptide’s distribution will be plotted in a separate row.

HXViewer will automatically estimate the required Y-axis range to accommodate all peptides in the current view. The field Required #D range lets you override this setting such that the Y-axis range will never be less than specified here. This may be useful if you want to get all plots in a series on the same scale.

There are three modes of interacting with the data plotted in the time series view: In Selection Mode, you can click on peptides and charge states to select them in the plot and in the tree view. Selected peptides or charge states will be highlighted in blue and you can hit the Delete key to disable the selected item. Right-clicking will open a context menu that lets you disable, select or remove the selected item. Please note the difference between disabling an item and removing it from the plot: a disabled item will still be associated with the time series view and therefore influences how the sequence of peptides is constructed. Removed items on the other hand remain active but are no longer associated with the plot. The latter corresponds to reversing the drag-and-drop operation from the tree view.

In Marker Mode, clicking on a graph value will display the corresponding deuteration value. This value is transiently plotted and replotting, e.g., by scrolling back and forth in the sequence or by disabling/activating a charge state, will remove the label.

Measurement Mode lets you measure deuteration difference between two different data points. Click on the first data point to initiate the measurement and click on a second point to complete it. You can abort a measurement by hitting Escape.
3.6 Graphics Export

Hexicon 2 has a standardized graphics export scheme. Export parameters can be set up in the export dialog (Figure 6). Export into a PNG file guarantees that the current view will be saved as-is. Vector graphics can be obtained by exporting into PDF. Graphics exceeding the display width can be saved as multi-page PDF files whereas HXViewer ensures correct breakpoints. There is a bug in PDF export through Qt (Qt bug 23142, unresolved in Qt 4.x as of 2013/11/30) which causes Adobe PDF readers to misinterpret the colorspace information of graphics containing transparency. This is invisible to physical printing and can be easily fixed by opening and re-saving the file in a vector graphics editor.

![Figure 6: HXViewer File Export Dialog](image)

B-Factor Export

Mapping of deuteration values or differences onto a crystal structure model is a popular way of visualizing HDX data. One commonly used way of encoding such information in a structure is to alter the B-Factors of a PDB file. PyMOL scripts like `data2bfactor` and `color_b` ([http://pldserver1.biochem.queensu.ca/~rlc/work/pymol/](http://pldserver1.biochem.queensu.ca/~rlc/work/pymol/), accessed 2013/12/04) can be used to modify and color-code B-Factors according to a text file. Use the menu item **Visualization > Create B-factor table** to generate a text file compatible with the `data2bfactor` script.

As with the peptide map (section 3.5), you can choose between one or two datasets to create B-Factors from, to get B-factors corresponding to dynamics or difference output, respectively. The task of integrating information from multiple peptides for each amide is resolved as follows: for any amide, if MS$^2$-confirmed peptides are available to cover this amide, use the longest of these. Otherwise, information from the longest MS-1 assigned peptide is used.
We are aware that there are multiple ways of mapping deuteration values from multiple overlapping peptides onto each amide of a structure. While the presented approach consistently yielded the most faithful representation of our experimental data, this may not apply to other datasets. We are planning to include other deuteration export policies in further releases. In the meantime you can take full control of what is exported with this "hack": Save your dataset first. Then disable all peptides using the Processing > Set all peptides > Disabled menu item (cf section 3.4). Open the peptide map view (section 3.5) and go through the peptide list (section 3.3) to activate only the peptides that you want exported. The peptide map will visually assist you identifying the correct peptides.