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1. Overview
Welcome to PEAKS Studio 11.

1.1 How to Use This Manual
This chapter provides an overview of PEAKS Studio 11’s features typical data analysis work flow in PEAKS Studio. It is strongly recommended for new users of PEAKS Studio to read this chapter to get a big picture of what PEAKS Studio provides and how to use PEAKS Studio. For users who are familiar with the PEAKS Studio product line from previous releases, it will be a similar user experience and this chapter may be used as reference.

1.2 What is PEAKS Studio 11?
PEAKS Studio 11 is the first release of PEAKS Studio on the new platform. PEAKS Studio 11 is an all-in-one software solution for data dependent acquisition (DDA) and data independent acquisition (DIA) shotgun proteomic mass spectrometry data analysis. It is a vendor-neutral platform capable of reading in raw mass spectrometry data and public data formats. Intuitive result visualization and validation tools are provided at every stage of analysis and results can be exported. In addition to the various DDA and DIA identification and quantification workflows with ion mobility support provided in previous releases, PEAKS Studio 11 highlights the new Glycan Search and Deep learning algorithm. Our newest release comes with new features and bugfixes. The following section provides an overview of the major additions in PEAKS Studio 11, including the new Quality Control function and enhanced PEAKS PTM Search with Signature Ion labelling. Additional feature requests were implemented to enhance user experience.

1.3 What’s New?
PEAKS Studio 11 introduces the enhanced Quality Control function for DDA/DIA Identification and Quantification results. PEAKS PTM Search has been improved with new Signature Ion detection and modified peptide view with confident PTMs.

Here are some of the highlights:

- Quality Control function
  Optional QC step added to both the DDA and DIA Identification and LFQ workflows. Analyze data quality to ensure data, identification, and quantification consistency across the sample set by evaluating key attributes for each sample.

- Confident PTM improvement and Signature ion detection
  Detect expected and potential signature ions in modified peptides. Report signature ions and display signature ions in spectrum chart. Highlight confident PTMs and compare modified and unmodified peptides.

- Mixed Mode support
  Added ‘MIX’ fragmentation mode support for Thermo mixed data which has two fragmentation modes.

- TMT QC
Added support for replicate channels in reporter ion quantification. Set CV filters for grouped replicate channels to improve labelled quantification.

- Local RT Normalization

Enhanced normalize to TIC and normalize to standard proteins with local RT normalization. Use TIC of nearby RT range as the normalization factor to improve ratio calculation.

- Modified Peptide View

Added a new view in Identification results to compare modified and unmodified peptides and display the spectrum chart in a mirror plot.

- Export Spectral Library from other results.

In addition to DDA Database search result, added support to generate PEAKS spectral libraries from these results: Denovo, Deepnovo, Peptide Search, DIA Denovo, DIA Database Search

- MGF, MSe, and Bruker .tsf loading

Added support for loading more data types.

- DeepNovo workflow

Enhanced DeepNovo search replaces De Novo if GPU is available.

There are also additional smaller features we believe that users will benefit from:

- Peptide View Filter adds length and -10lgP filter.

Provides a Length filter for quickly filtering for peptides of certain length and a -10lgP filter to improve user experience when looking for higher confidence peptides.

- Library Viewer support for OpenMS and Spectronaut spectral libraries.

Added some support for OpenMS and Spectronaut libraries to Library Viewer. Open and view libraries, convert them to PEAKS Studio format, and configure them for use in library search.

- Data mzXML Exporting

Added mzXML exporting for raw data. Combine with pepxml export to load into Skyline.

- Neutral Loss

Added support for specifying neutral loss when creating custom PTMs. Displays neutral loss in spectrum chart and ion match table.

- Skip Data Refine and Mass Correction

Provided the option to skip data refine for preprocessed data. Provided the option to not perform mass correction.

- Rename Sample

Added a way to rename samples that have been loaded or used in analyses already.
• TMT16plex / TMT18plex support

Added built-in TMT16plex and TMT18plex reporter ion quantification methods.

1.4 Installation and Activation

1.4.1 System Requirements

PEAKS 11 is recommended to be installed on 64-bit Windows operating system with Windows 10 or later. PEAKS 11 installer will take ~2.5GB of disk space and PEAKS 11 program files will use ~5GB disk space. The program files folder will take up more disk space over time as protein databases and glycan databases are configured and saved. The amount of disk space required for this purpose depends on the size of the user’s datasets. Protein databases may require up to double the file size of the configured .fasta file. It is recommended to allocate a large disk space for the purpose of running projects, and ensure the location of the project has a lot of free space to handle the size of the project.

The two main factors affecting PEAKS 11’s performance are CPU and RAM. The minimum, recommended and ideal requirements are as follows:

- Minimum: Quad-core processors and 32 GB of RAM.
- Recommended: 16 threads processors and 64GB of RAM with compatible GPU (described below).
- Ideal: Intel Core i7/i9/Xeon or AMD Ryzen 7/9/threadripper processors with total 32 threads or more, and 64-128 GB of RAM with compatible GPU.

For running DIA Database Search and DeepNovo, it is recommended that the machine is equipped with at least 64GB of free memory and a NVIDIA CUDA compute capability >= 4 GPU with at least 8GB of dedicated memory.

1.4.2 Installation on a Windows operating system

Double-click on the PEAKS Studio 11.exe installer.

Go through the steps of the Setup Wizard, selecting a directory for the application to be installed in. By default, it will select path C:\PEAKSStudio11

If you already have PEAKS Studio 11 installed and would like to update it, you can select the current installation path and overwrite that installation. When prompted, select “Yes to All” which will overwrite only the necessary files in the updated version. The configured databases and custom modifications, methods, etc. will not be replaced. However, the license key needs to be applied again and it will trigger a new performance configuration. Updated PEAKS Studio 11 has optimized the performance configuration further so it should have an improvement in runtime using the default configuration that gets applied.
1.4.3 Activation

Once set up, double-click on PEAKSStudio.exe in the installation folder.

You will be prompted for a license key. For purchasing a license or requesting a trial license key, please contact sales@bioinfor.com
After a successful activation, the application will perform an automatic performance configuration. Double-click on PEAKS.exe again to open the program and you will be greeted with the main interface.

1.4.4 Manual Activation

In case there is a problem in applying the license key conventionally, such as not being connected to the internet, or not being able to reach the license server, we have the option to activate PEAKS Studio manually.

Select the option in the bottom right and follow the steps outlined.

1. **Gather Information**: Input your name, email address, and license key to generate a license request file.
2. **Save License Request File:** Select a folder to save the license.request.

3. **Update License Request File:** On a computer with internet connection (it can be any computer, doesn’t have to be the computer with PEAKS Studio installed), go to the webpage [http://www.bioinfor.com/lcs20/index.jsp](http://www.bioinfor.com/lcs20/index.jsp)
   Click on Next and Choose File to select the license.request file. If this license.request file is from a different computer, use a USB key to move it between computers.
   Type in the 6 digit verification and then Register.

4. **Import License:** You should receive an email from support@bioinfor.com
   Back in Manual Activation, copy the license content in the email, or import the license file.

5. **Activation Complete:** Click Finish. Performance Configuration will automatically apply, and PEAKS Studio will need to be restarted.

### 1.4.5 License Key descriptions

To view the current license, click on the ![Information](image) icon under the Help menu located at the top of the interface. This opens the About BSI PEAKS Studio dialog where the current license details are listed.

![About BSI PEAKS Studio](image)

Each license key has a specified start / expire date. Licenses will have activation limits so it is intended to only be applied on the machines designated for PEAKS Studio 11. In the Module section, the following optional modules that are included in the license are listed.

**PEAKS Platform:** This is the base license without any add-on modules. Both DDA and DIA workflows are available.

**PEAKS IM module:** Enables running DDA and DIA analysis with ion mobility data, including timsTOF 1/k0 and FAIMS CV with results displayed in a 4D view.
PEAKS Q module: Enables running quantification workflows including Precursor ion quantification (SILAC/iTRAQ), Reporter ion quantification (TMT/iTRAQ), Label-free quantification for DDA and Label-free quantification for DIA.

PEAKS Glycan module: Enables running Glycan Search workflows.
1.5 Supported Data Formats

PEAKS 11 currently supports vendors with the following data formats:

1) Thermo - .raw, FAIMS, DIA
2) Bruker – timsTOF, tdf/tsf/baf/yep/fid, diaPASEF
3) Waters - MSe, .raw
4) SCIEX - .wiff, .wiff2 (requires license key), DIA, ZenoTOF
5) Agilent - .d
6) Shimadzu - .lcd
7) .mzml and .mzXML
8) .mgf

1.6 Quick Walkthrough

This chapter will go over the basic functions in creating projects and running analyses, as well as provide an overview of the different options available for deleting samples and analysis, modifying and running analysis, and other navigation options.

1.6.1 Start Page and Sample Project

Upon opening PEAKS 11 interface for the first time, you are presented with the Start Page seen below.
The Start Page lists recent news and highlights related to PEAKS Software line. Clicking on any of the titles will open the web browser showing interesting reads about recent developments.

In the top left “Recent Projects” section, there is an option to open the Sample Project. If you do not see the option, click on “Clear Project History”. The Sample Project is meant for users to have a quick view of what PEAKS 11 has to offer. This project cannot be modified or deleted in any way.

Once open, click on the arrow next to Analysis 1 and double-click on the DB Search result node to open the sample result. This is the Database Search result which you can learn more about in later sections.

### 1.6.2 Setting up a new project

This section is created for those not familiar with previous PEAKS Studio or PEAKS Online platforms. If you have experience with PEAKS products, the workflow settings and user interface (UI) for PEAKS Studio 11 will be straightforward.

To get started with a new project, in the toolbar, click on + to create a new project.
The Project Wizard opens at the Project Creation step. Here, you first add the raw mass spectrometric data files into the project. Add in a demo file from orbitrap instrument if you have one available. The data files selected need to be added into samples, and we provide an option to quickly configure a large number of samples by name using the button. Adding samples by **Delimiter** is a great feature for quickly grouping and naming samples for large experiments.

![Add sample(s) by Delimiter](image)

After selecting supported data files and adding them into samples, make sure to set the correct Instrument, Fragmentation, and Acquisition mode. Use the "**Copy to Whole Project**" button to copy the settings from the selected sample to all other samples.

Next, you may click “Finish” to load the data files only or you may proceed with setting up of your analysis by clicking on “Workflow Selection >”.

For raw data acquired in Data Dependent Acquisition (DDA) mode, five Workflow options are available:

- DeepNovo / De Novo
- PEAKS DB Search (optional PEAKS PTM Finder and SPIDER)
- PEAKS Q (LFQ, SILAC, TMT/iTRAQ)
- Glycan Search
- DeepNovo Peptidome
For raw data acquired in Data Independent Acquisition (DIA) mode, two Workflow options are available:

- DIA Identification (Spectral Library Search, Database Search, De Novo)
- DIA Quantification (Spectral Library Search, Database Search, LFQ)

Finally, you may create a new workflow or use a saved workflow from previous analysis, where saved parameters are automatically loaded into your new analysis. Click on the “Select Your Workflow” option to see a list of saved workflows. Any workflows you save later will appear here.
For the purpose of this demo, select the DDA PEAKS DB Workflow indicated below.

Click on “Data Refine” to begin setting up the PEAKS DB workflow.

The first step in PEAKS DB workflow is Data Refine.

**Associate feature with Chimera:** If enabled, PEAKS will assume that a tandem scan may contain two or more peptides, which will allow PEAKS to identify co-eluting peptides found within the acquisition window of the scan. It does this by searching for peptides that match the monoisotopic mass of any peptide feature in the acquisition window.

**Mass Correction:** The precursor m/z values given by some instruments are often not of the monoisotopic ion. This creates problems in downstream analysis. By examining the isotope shapes in the corresponding MS scans, this function can accurately adjust the precursor m/z to be equal to the monoisotopic ion m/z.

**Preprocessed Data - Skip Data Refinement:** For data that has already been preprocessed, such as those with centroiding, mass correction, and charge deconvolution. It is not recommended to check this option unless you are sure the data has been preprocessed.

Once selected, in the bottom right corner, click on “DB Search” to continue to the next step.
The DB Search setup contains parameters such as mass tolerance, enzyme and digest mode, missed cleavages, modifications (fixed or variable), database information, as well as advanced search options such as PTM Finder (search for additional modifications) and SPIDER (search for sequence variations).

Existing configured protein database may be selected, or a new database can be added and configured by clicking on “New” option. Note that for the Uniprot database, the taxonomy files (taxdmp and speclist) are already preloaded into the databases folder in the installation folder.

Once finished, click on QC to proceed to the next step.
The Quality Control function can be enabled to get important statistics about data quality as well as comparing the identification and quantification results with the selected control sample. It is optional and will run swiftly so it is encouraged to familiarize yourself with this function.

Click on Report to proceed to the next step in the workflow.
Report Filter is the final step in the workflow setup. Here, pre-selected scoring for PSM/peptide, protein, and De novo tag will determine the reported results. Important to note is that changing these filters and re-running the same analysis will take less time given that these filters are applied to the final output.

At this point, you may also choose to save your workflow parameters, which will allow you to re-select this workflow in the future. Lastly, the analysis can be given a name, or it can be renamed later in the project view.

Clicking on Finish will start loading the data files and running the analysis.

### 1.6.3 Monitoring analysis and task cancellation

Click on the **Progress** tab on the bottom left (below the project tree) to see progress status of projects or analyses being run. The statuses are as follows:

- **RUNNING**: The tasks in this step are running and in progress.
- **PENDING**: The tasks in this step are queued.
- **DONE**: All tasks in this step are complete.
- **FAILED**: An issue has prevented the task to complete.
- **CANCELLED**: The analysis was cancelled while in progress.
The progress bar is also colour-coded. Green indicates completed tasks in the step, blue indicates tasks in progress, orange indicates pending tasks in the step, red is an encountered failure during a task, and light blue is analysis cancellation.

In progress and pending tasks can be cancelled by right-clicking on the analysis.

A bar at the bottom left will also indicate if PEAKS Studio is currently busy running tasks.

Clicking on the Console tab will provide more details on the current tasks being run and may give more insight on how far the analysis progressed.
In the build directory there is a logs folder which also logs this running info. A tasks folder saves all the logs for each individual task ran.

When encountering issues, the user is encouraged to submit the logs folder to the PEAKS support team so that we can better assist you with the problem.

Clicking on the Properties tab will show details on each of the selected result nodes. For example, selecting a Glycan Search result node will display the parameters used in this setup in the Properties tab.

In addition, selecting either the Project node or Analysis node in the project tree will display the time that the project or analysis was created.
1.6.4 Project Tree Hierarchy

In the top left of the interface is the Project View. Project View will display any opened projects that are complete or in progress. The typical project tree has the following nodes:

1. **Project node (top level):** Displays the project name. It can collapse/expand the project. Right clicking will allow user to close the project or run another analysis in the project.

2. **Analysis node (second level):** Each analysis run will have its analysis node under the project. Each analysis can collapse/expand. Right clicking on the analysis node provides the option to change the name of the Analysis. There is also an option to delete the analysis from the project.

3. **Data Refine node (third level):** Expand to see the samples included in the analysis. Expand each sample to see the fractions or data files added to the sample. Double-click to open the data refine view. For more information on the Data Refine view, see 5. Data Refinement.
4. Result node (third level): The main result node. Double-click to open the result node. Later sections will cover details of the result view.

5. Export node (third level): The Export result node provides several options for exporting. This is particularly useful for exporting the results tables into a comma separated text file (csv). There are several other export options available such as mgf export, html export, and spectral library generation. Double-click to open the Export node and see the options available. For more details on each of the exports, refer to the Export sections in each of the result node chapters.

6. Sample node (second level): Displays the samples in the project. Click on the dropdown arrow to expand and reveal the data files in that sample. Right-click on the sample to delete it from the project. This can only be done if no analysis is using that sample and the project is not currently running an analysis.

7. Data File node (third level): Similar to the Data Refine Node but with unrefined (raw) data view. Double-click to open any of the raw data files.

1.6.5 Analysis and Sample renaming and deletion

Once the project analysis run is completed, the result nodes will appear in the project view (top left). Here, completed analyses can be renamed or deleted by right-clicking on them.

Samples can also be deleted by right-clicking on the Sample under ‘Data’ dropdown. Important to note that the Data can only be deleted if it is currently not a part of any analysis, and the project, that the sample is loaded into, is not running any analysis at that moment. Additionally, all analyses containing the sample to be deleted, must be deleted first. This is to prevent unwanted issues related to removal of the data from the project.

Samples can be renamed by right-clicking on the Sample while they are not being used in an analysis in progress. Once renamed, any results that are opened need to be reopened for the new name to apply.
1.6.6 Save Workflow from Analysis

Right-click on any analysis and select the option to save the parameters set in a saved workflow. Assign the workflow a name so that it can be used in future analyses without having to set parameters.

1.6.7 Running additional analyses in existing project and modifying analysis

In the Projects tab, click on to run a new analysis in the current project. Within Data Selection, In Project tab will allow selecting pre-loaded samples.

Another useful feature for running additional analyses is “Modify Analysis” function. Clicking on will prepare a new analysis with the same parameters as the existing one, which makes it a fast and convenient way to modify only specific parameters or report new filters without having to go through each of the settings again.

If the analysis is run with the same settings as the existing analysis in the project, the project should finish instantaneously, creating a duplicate of that analysis.

1.6.8 Project opening and closing

While the Projects tab is selected, the projects currently opened are at the top. Select the tab to switch between each project. Clicking ‘X’ on the tab will close the project.
Projects can also be closed by right-clicking on the project and selecting “Close Project” in the dropdown. It is recommended to close unused projects to preserve memory.

To open an existing project, click on the icon. You can also open from the list of recently opened projects by clicking on . Ten latest projects are listed here. Recent projects can also be opened from the Start Page Recent Projects lists.

1.7 Import PEAKS Online Project

PEAKS Studio 11 also supports importing archived PEAKS Online projects. Click on to select an archived Online project to view in PEAKS Studio 11.

Any result nodes that are available in both PEAKS Online and PEAKS Studio can be imported successfully into PEAKS Studio. Certain figures and tables will not be displayed since they are not available in the PEAKS Online project. Due to different project structures, running an analysis on an imported PEAKS Online project is prevented.
2. Configuration

Click on the button to open the configuration dialog. We provide a number of configuration options which will be explored in the following sections.

2.1 General Configuration - Default Directories Preferences

Default directories can be set in Configuration > General. Each of these directories are used as follows:

- **Default Input File Directory**: This is the folder that is selected initially when adding input data files. Choose the location of your source files for easy access when creating projects or analyses.

- **Default Project Directory**: This folder specifies the default location that projects will be created in. When opening the Project Wizard, this will be the path set in ‘Project Location’.

- **Default Export Directory**: This is the folder that is selected by default for all exports of PEAKS Studio. This includes exports of .csv and .html files from the Export node as well as any image exports provided.
Clicking on ‘Save’ will save the selected directories.
Clicking on ‘Reset’ will set the directories to default. A pop-up will prompt the user to confirm this action.

2.2 Enzyme Configuration

All of the built-in enzymes are listed here. A built-in enzyme cannot be modified or deleted.

Custom Enzymes can be created and updated. Provide the name of the new enzyme in the “Enzyme Name” field and specify how the custom enzyme will cleave the protein between two amino acids to create peptides in the “Enzyme Detail” panel.

The letter, X, denotes any amino acid at this position, while amino acids within the { and } brackets indicate any amino acid except for the one in the brackets. Choose where the cleavage sites are by selecting after or not after and before or not before to specify the range. Add multiple amino acids to indicate that cleavage happens before or after any of the stated amino acids. For example, after RK means after R or K not after R and K.
Click the Add/Update button to save the changes. The new enzyme will now appear in the “Enzyme List”, where it can be accessed later. To delete a customized enzyme, select the appropriate enzyme and click the Delete button.

2.3 Instrument Configuration

Create a new instrument.

Click the New button and provide a name for the instrument in the “Instrument Details” panel.

Use the “Ion Source” drop-down menu to select the ion source that was used: MALDI/SELDI or ESI (nano-spray). This will help the PEAKS Data Refine tool to determine the charge of the ions.

The names of the built-in instruments are provided in the “Instrument List”. Select an instrument to view the detailed instrument information in the “Instrument Detail” panel below.

The details of a built-in instrument cannot be deleted or edited.
Use the “MS Precursor Scan” drop-down menu to select the type of MS scan that was performed. This selection will tell the PEAKS Data Refine tool whether the survey scan is of sufficient resolution to determine the charge and the monoisotopic peak during examination of the survey scan.

Use the “MS/MS Product Scan” drop-down menu to select the type of MS/MS scan that was performed. This selection will help PEAKS decide which internal parameters (for weighing fragments and amount of noise) to use during PEAKS auto de novo sequencing and a PEAKS DB search. Select LIT/FT if alternating high-res/low-res modes were used. This will allow the algorithm to determine the mass analyzer from the scan header.

Specify the values for “Parent mass error tolerance” and “Fragment mass error tolerance” in Daltons or ppm.

Click the Add/Update button to save the changes. The new instrument will appear in the “Instrument List” where it can be accessed when creating a new project file. To delete a customized instrument, select the appropriate instrument from the “Instrument List” and click the Delete button.

2.4 PTM Configuration

<table>
<thead>
<tr>
<th>General</th>
<th>Recent</th>
<th>Common</th>
<th>Uncommon</th>
<th>Glycosylation</th>
<th>Artificial</th>
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<td>(X@N)</td>
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<td>(X@N)</td>
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<tr>
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<td>(C)</td>
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</tbody>
</table>
Create a new PTM. To create a new PTM, click on the New button in the bottom right-hand corner of the PTM Configuration window to display the “New PTM” dialogue. Provide the necessary information about the new PTM.

- **PTM Name:** This name will appear in the PTM list for future use after it is saved.
- **PTM abbreviation:** The PTM name expressed in a shortened form.
- **Mass (Monoisotopic):** The mass that the residue gains or loses as a result of the PTM.
- **Residues that can be modified:** Enter residues that can be modified anywhere, or residues that can only be modified if they are at the N- or C-terminus. User will also need to specify if these residues can be modified on the peptide-level or protein-level.
- **Formula:** The chemical formula of the PTM. This should correspond to the mass listed above.
- **Rule:** This field is used for the user’s reference only. Users can use the field to enter a comment about the PTM.
- **Neutral Loss:** Specify potential neutral loss for this modification.
- **Signature Ions:** Specify expected signature ions for this modification.
Click the OK button to save the changes. The new PTM will now appear in the “Customized” PTM list, where it can be accessed later. To delete a customized PTM, select the appropriate PTM from the list and click the Delete button.

The “Recent” tab lists all the PTMs recently used in analyses.

2.5 Algorithm Configuration

The Algorithm Configuration is used to tune certain algorithm parameters that are used in the workflow. At present, PTM Search related algorithm parameters can be set.

Confident PTM

Peptide Tag Length ≥ X : The number of X consecutive ions at the modification site in the peptide that have ion intensity above the value Y set in the following algorithm parameter. If the PTM takes place at the N-term or C-term, then there needs to be a number of consecutive ions + 1 (default would be 3 + 1).

Peptide Tag Ion Intensity ≥ Y : Defines the minimum ion intensity Y that is used in considering whether the PTM is confident.

For example, ANGTTLVGVMPAGAK is the peptide of interest, which has Deamidation (N) at position 2. We look at X=3 consecutive amino acids in the peptide, so it would be ANG, or NGT. Whether it is b-ion or y-ion, we...
check \([b2, b3, b4]\) or \([b1, b2, b3]\) or \([y13, y14, y15]\) or \([y14, y15, y16]\) and their ion intensity. If those ion intensity exceeds \(Y=2\%\), then this modification will be reported as a confident PTM.

**Potential Signature Ions**

Peak ion intensity \(\geq Y\): The ion intensity for a peak must exceed \(Y\) for it to be considered as a potential signature ion. This is used in conjunction with the Use top \(X\) peaks in the spectra parameter to only select \(X\) peaks with ion intensity \(\geq Y\). If a signature ion does not meet these criteria anywhere, it will not display in the result.

Use top \(X\) peaks in the spectra: Only the top \(X\) peaks (in intensity) in the spectra are considered when searching for potential signature ions. This is used in conjunction with the Peak ion intensity parameter in determining how many peaks are considered when searching for potential signature ions.

Peak existence in \(\geq X\) \% of modified PSMs: For the list of Modified PSMs, if the number of PSMs with this signature ion is greater than \(X\) \% of the total, then the potential signature ion will be reported. Potential signature ions that do not pass this threshold will still be in the all potential signature ion export which is useful in determining how to tune this parameter for future runs.

\[\text{e.g. There are 5000 PSMs with ADP-Ribose, but only 1400 detected signature Ion ADP_232. Since this is 28\% of the total, and that is below the default 30\%, potential signature ion ADP-Ribose_232 is not reported.}\]

Difference between modified and unmodified PSMs \(\geq X\) \%: Both the list of Modified PSMs with the modification of interest that has the detected signature ion as well as the list of Unmodified PSMs that has the detected signature ion are recorded. A \% of the Modified PSMs will have the potential signature ion detected, and a \% of Unmodified PSMs will have the potential signature ion detected. If the \% of Modified PSMs is greater than \(X\) \% compared to the \% of Unmodified PSMs, then this signature ion is reported as a potential signature ion. Potential signature ions that do not pass this threshold will still be in the all potential signature ion export which is useful in determining how to tune this parameter for future runs.

\[\text{e.g. There are 5000 PSMS with ADP-Ribose, and 10000 without ADP-Ribose.}\]

\[\text{Of those 5000 ADP-Ribose PSMs, 3000 detected signature ion ADP_250. This comes to 60\% of the total.}\]

\[\text{Of those 10000 Unmodified PSMs, 2000 detected signature ion ADP-250. This comes to 20\% of the total.}\]

\[\text{Since 60\% - 20\% \geq default 30\%, then the potential signature ion ADP-Ribose_250 is reported.}\]

Modified PSM count \(\geq X\): There needs to be at least \(X\) Modified PSMs in the result with this potential signature ion for it to be reported in the result.
PEAKS Studio comes with a number of predefined labelled quantification methods for TMT and iTRAQ quantification. Custom label Q methods can be defined by users.

**Built-in Label Q Methods.** The built-in label Q methods within PEAKS Studio come standard with all installations and cannot be modified. To modify a built-in label Q method click the “Duplicate” button which will create a new label Q method (it will not modify the original method) with the same settings as the built-in method, which can then be modified.

**Create a new Label Q Method.** Click the New button to create a new label Q method.
PTM: The modifications used as labels for this label Q method. Click the Set PTM button to open the PTM options and set the modifications for this label Q method.

**Label Options:** The labels used by this label Q method. Each label must include:

- **Name:** (e.g.: iTRAQ-114). This will be the name associated with that channel in the labelled Q parameter page and on labelled Q result pages.
- **Reporter Ion Mass:** (e.g.: 114.11 for iTRAQ-114). The mass for this channel's reporter ion.

Click OK to save this method into the configured reporter ion quantification list.

**Edit a custom Label Q Method.** To edit a label Q method, click the “Edit” button to edit any fields.

**Delete a custom Label Q Method.** The user can delete the custom label Q Methods, but not built-in label Q methods. Click on the “Delete” button to remove the custom method.
2.7 SILAC Q Method Configuration

PEAKS Studio comes with a number of predefined SILAC quantification methods. Custom SILAC Q methods can be defined by users.

**Built-in SILAC Q Methods.** The built-in SILAC Q methods within PEAKS Studio come standard with all installations and cannot be modified. To modify a built-in SILAC Q method click the "Duplicate" button in the SILAC Q method's actions column. This will create a new SILAC Q method (it will not modify the original method) with the same settings as the built-in method, which can then be modified.

**Create a new SILAC Q Method.** Click the “New” to create a new SILAC Q method. Fill in the settings below and click “OK” to save this custom method.

Quantification Method Detail Name: This is the name that will appear in the SILAC Q methods list for future use after it is saved. If selecting an existing name, it will replace that method in the list. Names with built-in methods cannot be used and will not replace built-in methods.

Click on Add Row to add additional labels and set the Label Options as follows:
• Name: (e.g.: Light). This will be the name associated with condition in the SILAC Q parameter page and on SILAC Q result pages.

• Modi: The modifications used as a condition for this SILAC Q method. Click the button to open the PTM Options dialog in the Artificial tab and set the modifications for this SILAC Q method.

• Modi Detail: Shows the selected modification's monoisotopic mass and residue site.

**Delete a custom SILAC Q Method.** The user can delete the custom SILAC Q Methods, but not built-in SILAC Q methods. To delete a SILAC Q method find it in the Quantification Method List and click on the “Delete” button.

**Edit a custom SILAC Q Method.** To edit a SILAC Q method find it in the list and click the “Edit” button. All fields are editable, and if the method name is changed, it will save a separate custom SILAC Q method and keep the original as well.

### 2.8 Database Configuration

**Database List**

- uniprot_sprot_June-2-2020.fasta
- Swissprot-Human-20329.yeast.7904._v201803.fasta
- Reviewed_Human_2022-09-19.fasta

**Database Detail**

**FASTA format data...**

**UniProtKB (Swiss-Prot, TrEMBL, ...)**

**Basic Option**

- **Database name:** uniprot_sprot_June-2-2020.fasta
- **Database path:** D:\1-Databases\uniprot_sprot_June-2-2020.fasta

**Advanced Option**

- Rule to parse accessionId from FASTA title:
  ```
  ^>(\S+)(?i)\d+$
  ```
- Rule to parse description from title:
  ```
  ^\S+\1\$\n  ```

563062 sequences
The “Database List” on the top shows all the configured databases in the system. Select a database from the list to view the detailed information about the database in the “Database Details” panel.

**Create a new database.** Click the New button to create a new sequence database entry or select a database from the “Database List”. The database can be configured in the “Database Details” area below. Click the Add/Update button to save any changes.

**Note:** Databases can also be configured while creating a project in the Project Wizard.

**Moving/Updating a database.** To move a database to another directory, the location must be updated in PEAKS. Select the database and specify the new location using the Browse button next to the “Database path” field. Then, click Add/Update to save the new settings. If the database location is invalid, an error pop-up will be shown and the change will not be saved.

**Delete a previously configured database.** To delete a database file, select the database to be deleted from the “Database List” and click the Delete button at the bottom.

**Note:** A copy of the fasta database is copied into the installation folder `db` when configured. This is the main reason PEAKS Studio installation directory will occupy more disk space over time as each database file configured can be roughly two times the size of the original database.

**Note:** NCBI Database configuration can take a very long time depending on the size of the database. Please prepare for PEAKS Studio to be busy during that time and it is recommended to configure NCBI Databases when not actively using PEAKS Studio for other purposes.
2.9 Taxonomy Configuration

The PEAKS Studio build bundles a taxdmp and speclist in the `databases` folder of the installed directory. Clicking on the “Browse” button next to the text field will allow user to specify the path to a different taxdmp folder or speclist.txt file. For NCBI databases, set `gi_taxid_prot.dmp` file for the NCBI GI taxid prot and set `prot.accession2taxid.gz` file for NCBI accession to taxid.
## 2.10 Spectral Library Configuration

The “Library List” shows currently configured spectral libraries that can be used in PEAKS Studio DIA library search. Click on “Browse” Library path to select a valid library file and give it a name in Library name. Clicking “Add/Update” will configure this library with the name set, which will replace an entry if that name already exists.

Important to note that certain library files compatible with PEAKS Studio 11 can be configured. They are as follows:

- **PEAKS Studio generated spectral library**, which is exported from the Database Search result. See Section 6.9.4 PEAKS DB Search result - Exporting - Spectral Library generation for how to export this library. Select the `.tsv` file as the Library path, and if there is a `.info` file in the same path, it will also be read and additional information about the library will be shown.
- **PEAKS Online generated spectral library**, which is downloadable from PEAKS Online. Similar to PEAKS Studio libraries, select the `.tsb` file as the Library path, and if there is a `.info` file in the same path, it will also be read and more information about the library will be shown.
- **PEAKS Xpro generated spectral library**, which is exported from the Database Search result from the previous PEAKS Studio release. However, before being able to configure this library, it will have to be
converted into a compatible library format using the Library Viewer tool (see 12.2 Spectral Library Viewer).

Click on “New” to clear the Library path and Library name. Clicking on “View” will open the Library Viewer tool. More details on this will be covered in 12.2 Spectral Library Viewer.

Click on “Delete” to remove that library from configured library list.

2.11 Glycan Database Configuration

Much like protein databases in section 2.7 Database Configuration, Glycan databases can also be configured. PEAKS Studio 11 supports GDB and GlycoCT formats. Select a database by clicking on “Browse” next to Database path and give it a name. Click “Add/Update” to configure the selected glycan database. If the name is already in the glycan database list, it will replace that configured database. If the name is shared with a built-in database, it cannot be configured as built-in databases cannot be modified or removed.

Click on “New” to clear the Database name and Database path for a new entry.
Click on “Delete” to remove the configured glycan database. Built-in databases cannot be removed.

2.12 Workflow Configuration - Saved Workflows

Saved workflows are managed in the Workflow Configuration. To create a new saved workflow, check the Save Workflow option in the “Report” step of any workflow in the Project Wizard.

Alternatively, right-click on an analysis and select the Save Workflow option in the dropdown menu.

Select a workflow from the “Workflow list” to view more details on the parameters set for that workflow which will be automatically filled in the project wizard. A search box is provided for easy filtering of saved workflow names. Click on “Clear” to reset the search box.

Clicking the “Delete” button will delete any selected workflows in the Workflow list. A confirmation dialog will appear to ensure this is not done by mistake.
2.13 Performance Configuration

The Performance Configuration is accessed by clicking on the button under Configurations tab. It will list your computer resources including total physical memory and number of processors. Allocated resources for PEAKS Studio is automatically calculated based on your computer’s specifications and can be modified at any time. Be sure to leave some processors and memory free to allow other programs to function as normal while PEAKS Studio is open. The Detach Service JVM checkbox is selected by default. It allocates 2 of the total threads and 8GB of the total allocated memory specifically for the User Interface updates. Unchecking this may have performance issues when using PEAKS Studio, and it is only recommended to be disabled on computers with low thread count (8 or less) and less memory (32GB or less).

If the computer has a compatible GPU, it will be listed and the option to “Use GPU” can be selected. As a reminder, a compatible GPU needs to have a CUDA compute capability version >= 4, which can be discerned from NVIDIA’s website [https://developer.nvidia.com/cuda-gpus](https://developer.nvidia.com/cuda-gpus)

Ensure that the GPU has at least 8GB dedicated memory, and that the drivers are up to date.

Click “Apply” to save your preferred settings, and restart PEAKS Studio for the changes to be applied.
3. Data View & Data Refine View

The data view & data refine view gives a visualization of the mass spectrometric data loaded into the project. It allows for the inspection of LC-MS/MS details in an intuitive way for quality control and troubleshooting issues.

In the Data Refinement View, the information of a typical DDA LC-MS/MS data file is organized into 3 sections:

- **MS** - This section shows the TIC (total ion chromatogram) plot and all the MS scans. For each MS scan, the corresponding MS/MS scans are also displayed.

- **MS/MS** - This section lists all the MS/MS scans. For each MS/MS scan, the corresponding features are listed below, and associated number of features is displayed next to the MS/MS scan information header.

- **LC/MS** - This section provides a bird's eye view of the whole LC-MS dataset with detected features and MS/MS scans marked on refined data.

After opening a data file by double-clicking the data node from the Project View pane, the choice of different views can be made by choosing different tabs in the upper-left corner of the data view window. Hover mouse over an opened Data tab to see a tooltip of which data file it is and from which analysis.

3.1 Data Refinement - MS View

The MS View contains the TIC and all of the scans. The total ion chromatogram (TIC) is displayed on the left of the MS view. The navigation buttons are highlighted in the figure. To collapse the TIC chart, click the left navigation button. Clicking on a specific position in TIC will display the corresponding survey (MS) scan. To navigate the MS scans, use the up and down navigation buttons. The MS scans can also be navigated through using the up and
down arrows of the keyboard. The tandem (MS2) scans associated with the selected MS scan are shown in the panel below, where each can be viewed by selecting the different tabs identified by their "m/z" information.

The survey scans and tandem scans provide a few convenient ways to zoom and navigate the spectrum:

- **Zoom to an m/z region** - Click the desired start m/z and drag horizontally to the desired end m/z, release the mouse button.
- **Zoom in/out smoothly** - Place the mouse pointer at a particular m/z value (right below the x-axis line), scroll the mouse wheel button.
- **Increase/decrease the peak intensity** - Place the mouse pointer on the spectrum, scroll the mouse wheel button.
- **See the whole spectrum** - Double-click on the spectrum or click the 1:1 button.

### 3.1.1 Data Refinement - MS View - timsTOF Instruments

timsTOF Instruments have a different MS View that includes Ion Mobility information.

The total ion chromatogram (TIC) is displayed on the left of the MS view. A red bar on TIC curve points the current selected retention time. The top panel at right side shows the MS1 data at the selected retention time as an MS1 Frame - a heat map in m/z-1/k0 dimension.

The precursors associated with the selected MS1 Frame are listed at the bottom panel - one tab per precursor. The MS2 data associated with a precursor are pooled to form a combined MS2 spectrum. Each precursor tab shows the corresponding combined MS2 spectrum.
3.1.2 Data Refinement - MS View - DIA Acquisition

Data Refinement is also available for DIA data. The MS tab is similar to DDA but without any associated MS2 displayed for each MS1 scan. Select both timsTOF instrument and DIA acquisition to run diaPASEF data, which will have a similar MS View.

3.2 Data Refinement - MS/MS View

The MS/MS View shows the list of tandem scans on the left. If data refinement has been performed on the data file, each tandem scan in the list will indicate the total number of associated features at the end of the spectrum header [ex. "695.8360 (RT=26.95 Scan=1348) 2" from the image below indicates that there are 2 peptide features associated with the selected tandem scan]. In addition, when data refinement has been performed, a tandem scan header can be expanded and the associated features and their details (e.g. m/z, RT, charge) are listed below. When a tandem scan is selected from the list, the following information is provided to the right of the list: a summary of the spectral results, the MS/MS spectrum, the zoomed in LC/MS view, the Ion Match Table, and its Survey Scan. Similarly, when an individual peptide feature is instead selected from the list, the result for that feature is displayed. Zoom options are the same as described in the “MS View” section.

- **Summary of Spectral Results:** A table with a summary of the spectral results is displayed here, including the associated feature information, de novo sequence (if de novo sequencing has been performed), and database ID sequence (if a PEAKS search has been performed).

- **MS/MS Spectrum:** A view of the tandem scan will be displayed here. If de novo sequencing and/or a PEAKS search has been performed for the data file, the scan will be annotated. By clicking the sequence results from the Spectral Result table, the corresponding annotation will be displayed. Users will also have the ability to turn on the alignment and error map for the spectrum view when de novo sequencing and/or a PEAKS search is performed.

A search function is provided to search for specific Scan numbers (or precursor ID for timsTOF data files).

- **LC-MS Snapshot:** The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The MS/MS scan is indicated by a red framed square and the span of the yellow line designates the isolation window. If data refinement has been completed, purple circles will also be displayed on the heat map, and selecting an individual feature from the list on the left will highlight the feature in purple.

- **Parent Scan:** The “Survey” tab displays the precursor scan with the best signal within the isolation window. The buttons that appear in this section are identical to those explained in Section 5.2, “MS View”.

- **Ion Match Table:** An "Ion Match" tab will appear when one of the sequence results from the Spectral Results table is selected. The “Ion Match” tab in the bottom pane displays a table with the possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated fragment mass, where mass values displayed in color indicate fragment ions present in the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in “Spectrum Annotation Settings”.

- **Glycan Ion Match Table:** For glycosylated peptides identified using Glycan Search, a Glycan Ion Match table is provided showing the Theoretical m/z and Observed m/z of each Glycan Moiety matched ions.
3.2.1 Data Refinement - MS/MS View - timsTOF Instruments

MS/MS View for data acquired using a tamsTOF instrument includes Ion Mobility information.

The MS/MS View shows the list of precursors on the left panel. The right panel shows the details associated with the selected precursor. The combined MS2 spectrum is shown in the right middle panel. The MS1 Frame, to whom...
the precursor belongs, is displayed at the bottom-right panel zoomed in to show the precursor and the associated feature.

3.3 Data Refinement - LC/MS View
The LC/MS view shows the distribution of LC-MS signals/features as a heat map. Placing the cursor on the heat map will show the m/z value and the retention time of that point.

The LC/MS view provides various convenient ways to zoom and navigate the LC-MS features in the data:

- **Move to a specific heat map area**: Click on the heat map, hold and drag the cursor until the desired area of the heat map is reached and then release the mouse button.

- **Zoom to a specific heat map area**: Right-click on the heat map, hold and drag the cursor to select the desired area of the heat map and then release the mouse button.

- **Move to a specific heat map area from thumbnail**: Current view range of the heat map is shown as a rectangle in the Thumbnail. Click on the thumbnail view range rectangle, hold and drag the cursor until the desired area of the heat map is reached and then release the mouse button.

- **Jump to a specific heat map area**: Provide approximate m/z and RT values separated by a space in the Locate: m/z RT box and then hit the Enter key to zoom in to an area centered at the specified m/z and RT value.

- **Zoom in/out from thumbnail**: Place the cursor on the Thumbnail and then scroll the mouse wheel button.

- **Zoom in/out smoothly**: Place the cursor at a particular (m/z value, RT) position and then scroll the mouse wheel button.
• **Move to a specific RT range:** Click the cursor on the TIC or the RT axis, and then hold and drag the cursor until the desired range is reached.

• **Zoom to a specific RT range:** Place the cursor on the TIC or the RT axis, and then scroll the mouse wheel button.

• **Move to a specific m/z range:** When the m/z axis is zoomed in, click the mouse on the m/z axis, and then hold and drag the cursor until the desired range is reached.

• **Zoom to a specific m/z range:** Place the cursor on m/z axis and then scroll the mouse wheel button.

• **Change contrast of the heat map:** Place the cursor on the heat map, press and hold the Ctrl key, and then scroll the mouse wheel button.

• **Change peak height in the 3D intensity mode:** Place the cursor on the heat map, press and hold the Ctrl key, and then scroll the mouse wheel button.

• **View the whole heat map:** Click the 1:1 button in the control.

• **Move back/forward to previously specified heat map area:** Use the Back and Forward buttons in the control to navigate the previously specified heat map areas.

### 3.3.1 Additional LC/MS Controls

The heat map view offers various controls to study the LC-MS data features. The following sections describe the control options and their uses in detail.

#### 3.3.1.1 Noise Level

Select the appropriate threshold for noise filtering by peak and/or feature intensities. Once selected, the heat map view will reflect the changes dynamically.

#### 3.3.1.2 Show/Hide Features

To view the detected LC/MS features, mark the Features checkbox in the control. The features detected during the data refinement steps will be highlighted by small, red, diamond-shaped points on the heat map. Placing the cursor over such a feature point will show the boundary of the feature using a red polygon and display a pop-up window that includes the following information: the monoisotopic m/z, the mean RT, the charge state of the feature, and the total area under the XIC curves of the feature.
Note: Multiple features can be selected and represented by different colors by holding Ctrl and selecting the feature points.

3.3.1.3 Locate M/Z and Retention Time

Type an m/z and an RT value, such as "618.88 50.5" in the "Locate" text box to pinpoint a specific (m/z, RT) position in the LC/MS view.

3.3.1.4 Show/Hide MS/MS Spectra

Select the MS/MS Spectra checkbox to indicate where a tandem scan has been taken. The location will be marked with blue-outlined squares on the heat map. Place the cursor over a square to view the precursor m/z, the RT, the precursor charge state, and the scan number of that spectrum. Click on the square to view the acquisition window (indicated by the yellow line), and the spectrum in a pop-up window. Once selected, the square will become red-outlined.
3.3.1.5 Show/Hide Markers

Select the checkboxes to show the markers of where tandem scans are associated with different types of peptides.

- MS/MS Spectra will display all tandem scans. These are marked by empty blue markers.
- De novo tags will display all tandem scans associated with peptides found in De Novo search. These are shown as empty orange markers.
- De novo only tags are associated with the de novo only table in the DB Search result. These are shown as solid orange markers.
- Identified peptides are markers from Database search results such as DB Search, PTM Search, and SPIDER. These are shown as solid blue markers.
- Quantified peptides are markers associated peptides from TMT and SILAC quantification. These are shown as solid green markers.
- Glycopeptides are markers from Glycan Search glycopeptides. The location will be marked with a purple marker for N-linked glycopeptides and yellow for O-linked glycopeptides.

Place the cursor over a square to view more details about that spectrum in the tooltip. Click on the square to view the acquisition window (indicated by the yellow line), and the spectrum in a pop-up window. Once selected, the square will become red-outlined. Note that each of these markers can be jumped to from each of the respective result tables.
For FAIMS data, each marker will have different shades of colour to differentiate between CV values.
3.3.1.6 Intensity View

In addition to the heat map view, PEAKS also provides a 3D intensity view for visual inspection of the LC/MS data.

To switch between the heat map view and the intensity view, click the button in the top-right LC/MS control options. The intensity view has the same set of controls to easily zoom and navigate. Click the button to switch back to the 2D view.
3.3.1.7 Navigation History

The left and arrow buttons in the control panel provides the option to switch back and forth to previous views such as zoom levels, m/z or RT ranges, 3D view, and different noise level filters.

3.3.1.8 Export LC/MS

Click the icon on the LC/MS control options to export the LC/MS heat map or intensity view in PNG format.

3.3.2 Data Refinement - LC/MS View - timsTOF Instruments

The LC/MS view of timsTOF Instruments have an extra IMMS button to view the IM-MS dimension. Clicking on it will show a confirmation pop-up to process data reduction and index creation. This will take some time depending on the size of the data. Once complete, the IMMS pop-up will be shown with 1/k0 on the y-axis, m/z on the x-axis, and the RT range is based on the range in the LC/MS view. Adjust the RT in the LC/MS to adjust the RT range in the IMMS view.

The set of controls and functions in the in IMMS view is the same as in the LC/MS view.
3.3.3 Data Refinement - LC/MS View - DIA Acquisition

The Data Refinement LC/MS view is also provided for DIA data. It only has Features markers and no tandem scan markers. The controls are the same as conventional LC/MS view controls.

3.4 Data View

In the Project View, double-click on a data file under a Data node to view the raw data visualisation. This view is similar to the Data Refine view except it is from the raw data. The MS View, MS/MS View, and LC/MS View are all similar to what is shown in the Data Refine result node.
3.5 Project Wizard Data Refine

In the following chapters, each workflow type will be explored. Data Refinement is a mandatory step for every analysis. For DDA workflows, a Data Refine Option “Associate feature with Chimera [DDA]” is provided.

**Data Refine Options**

- [ ] Associate feature with Chimera [DDA]

The Data Refinement step has a single option “Associate feature with Chimera [DDA]”. If checked, PEAKS will consider a tandem scan can contain two peptides and PEAKS will search for chimera spectra to enable correct inference of co-eluted isobaric peptides.

3.6 Data Exports

When any DDA analysis is completed, an Export node appears with options for Data Exports. In Project View, double-click on Export to open the Export node. The loaded Data can be exported as Raw MS2 scans or Refined MS2 scans in MGF format. Raw scans can also be exported in mzXML format.
4. Denovo peptide sequencing

De novo sequencing derives a peptide sequence directly from the tandem mass spectrum without the need of a sequence database. It is the preferred method for identifying novel peptides and the study of unsequenced organisms.

PEAKS Studio provides a reliable and comprehensive solution for automated peptide de novo sequencing. It features:

- **Accurate de novo sequencing**
  
The PEAKS Studio de novo sequencing algorithm and scoring functions are constantly tuned for each instrument type to ensure optimal accuracy.

- **Easy-to-use user interface**
  
PEAKS Studio generates a comprehensive result summary and provides interactive views of annotated spectra.

- **Support of all major instrument types**
  
PEAKS Studio supports de novo sequencing of spectra generated by various mass analyzers (Ex. Orbitrap, Ion Trap, Time-of-Flight, Quadrupole) and instrument types (Ex. Orbitrap Eclipse with FAIMS, timsTOF, ZenoTOF) in common fragmentation modes (Ex. CID, HCD, ETD, ETHCD).

- **Automated result validation**
  
PEAKS Studio assigns a local confidence score to each amino acid in a de novo sequence. The local confidence score ranges from 0 to 99%, indicating how confident the algorithm considers a particular amino acid residue to be the correct assignment. Moreover, the peptide sequence is evaluated by an ALC (Average of Local Confidence) score, which is the average of the local confidence scores of all the amino acids in the sequence.

**Note:** For details about the PEAKS de novo sequencing algorithm, please refer to the initial publication:


Other PEAKS Studio tools, including PEAKS DB for database search, PEAKS PTM for unspecified PTM search, and SPIDER for homology search, depend on the de novo sequencing result to achieve superior sensitivity and accuracy.

4.1 De Novo Analysis Workflow and Parameter Settings

To run a De novo Analysis, Select the De Novo option in Workflow Selection

After deciding the Data Refine option, proceed to the next step to see the Denovo parameters.
The following parameter settings are provided:

**Precursor Mass Error Tolerance**: The maximum mass difference between the de novo peptide and the monoisotopic mass of the precursor ion.

**Precursor Mass Tolerance unit**: PEAKS de novo generates peptides within the precursor mass error tolerance. Precursor mass error tolerance can be specified in either Daltons or ppm.

**Fragment Mass Error Tolerance**: PEAKS de novo uses this value when scoring de novo peptide sequences. PEAKS Studio considers a fragment ion to be matched if the calculated m/z is within the specified error tolerance. Units are Daltons.

Both precursor and fragment error tolerance parameters need to be set consistently with the mass accuracy of the instrument.

**Enzyme Settings**: Select the enzyme used for protein digestion from the enzyme drop-down menu. PEAKS de novo respects the enzyme specificity at both ends when generating peptides.

Note: When the selected data node is a project of multiple samples, "Specified by each sample" allows samples to be analyzed separately, using their respective specified enzymes during project creation.
Note: "None" is a special enzyme allowing non-specific cleavage at both ends of the peptide. It is the recommended option when no digestion enzyme was used or when the digestion enzyme exhibits a high degree of non-specificity.

**PTM Settings:** Click “Set PTM” to open the **PTM Options** dialog and specify the fixed PTMs and variable PTMs expected in the experiment. Highlight PTMs and click on the button to place them as either Fixed or Variable PTM. From there, highlighted PTMs can be removed by clicking the “Remove” button. Clicking “Remove All” will reset the PTM options. Highlighting and clicking “Switch Type” will move the Fixed PTM in into the Variable PTM list and vice-versa. Click on “New” to create a new custom PTM. While a PTM is selected, click on “View” to see more details.

Note: Multiple PTMs can be highlighted by holding down the CTRL key or SHIFT key on the keyboard.

Note: For a specific residue, only one fixed modification is allowed.

Once the PTMs are selected, the maximum number of variable PTMs per peptide can be specified and a value of less than 4 is recommended.

Note: The use of variable modifications increases the size of the computational search space for the de novo sequencing algorithm. It is recommended not to select too many variable modifications in PEAKS de novo. See PEAKS PTM for alternate search options.
Click on "Report" to proceed to the final step. The Denovo result does not have any Report filters that need to be set. From here, there is the option to specify an Analysis Name (which can be later renamed) and/or decide to save the Workflow to quickly re-use in future analyses.

4.2 Understanding PEAKS De Novo Sequencing Results

When de novo sequencing is done, double-click on the De Novo result node to open the Denovo result. The result contains three tabs:

- **Summary**: This shows an outline of the PEAKS de novo search results with key statistics. The overall quality of the experiment can be examined and the filters for de novo ALC can be adjusted at the top of the page.
- **De novo**: This shows a list of peptide sequences that pass the ALC threshold detected by de novo sequencing.
- **Feature**: This shows a list of features detected in the refined data. Features associated denovo peptides are easily identified here.

The following sections will explore each of these three tabs in the Denovo result.

4.3 Denovo result - Summary tab

In the summary tab, a PEAKS Studio de novo sequencing result can be filtered using Average Local Confidence (ALC) score. Low quality de novo sequences can be filtered out by specifying a minimal threshold of the ALC score. The purpose of filtering is to remove poor sequences in which residues are incorrectly identified. By default, the ALC threshold is set to 50%. After changing the threshold, click to apply the new filter. The result in the Denovo result view will be updated accordingly.

Click on to open the Notes dialog to save useful information regarding this Analysis. The Notes will appear in Section 1 of the Summary page at the top.
Section 2 Result Statistics gives key statistics in Table 1 on the number of MS2 scans and # PSMs total as well as with different ALC thresholds for each sample. The #PSM / #MS2 provides a de novo sequencing identification rate metric for each sample.

Table 2 simply shows what is the current Denovo ALC filter set. This is placed here so that when the Summary page is exported, the viewer will know what the ALC threshold was at the time.

Section 3 Other Information shows the parameters that were set when the analysis was ran.

4.4 De novo result - De novo View

The "De Novo View" displays the de novo sequencing results in greater detail, as shown in the next figure. The table on the top displays de novo sequences, while the bottom section provides additional information about the peptide-spectrum match.

4.4.1 De novo View - De novo table

The following list describes the contents in each of the columns of the Denovo table:

- **Scan**: The scan number and data file it belongs to. The scan number is a unique index for tandem mass spectra in the data.
- **Peptide**: The amino acid sequence of the peptide as determined by de novo sequencing. A modified residue is followed by a pair of parentheses enclosing the mass of that modification.
- **ALC(%)**: The average local confidence. ALC is calculated as the total of the residue local confidence scores in the peptide divided by the peptide length. The higher the score the more confident the denovo candidate.
- **Length**: The number of amino acids in the sequence backbone.
- **m/z**: The precursor mass-to-charge ratio.
• **z:** The precursor charge.
• **RT:** The retention time (elution time) of the spectrum as acquired by mass spectrometer.
• **Area:** The area under the curve of the peptide feature found at the same m/z and retention time as the MS/MS scan. This can be used as an indicator of the abundance of the peptide.
• **Mass:** The monoisotopic mass of the peptide.
• **ppm:** The precursor mass error, calculated as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$.
• **PTM:** The types and the numbers of modifications present in the peptide shown in color-coded icons.
• **Mode:** The fragmentation mode in which de novo sequencing is performed by the algorithm.

Click on any of the column headers to sort the column.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

In the table, Amino acids in de novo sequences are color-coded according to their local confidence scores. Red represents a very high confidence (greater than 90%); purple represents a high confidence (80 to 90%); blue represents a medium confidence (60 to 80%); and black represents low confidence (less than 60%). Hover the cursor over the sequence to view local confidence scores of each individual amino acid.

### 4.4.2 De novo View - De novo table controls
The menu above the Denovo table provides options to scroll through the table of de novo peptides, filter for mass tags, switch between samples, search for specific entries, and export the table.

#### 4.4.2.1 De novo table controls - Sample Selection
Result for different samples can be viewed by selecting the sample using the "Show Results for Specified Sample" drop-down menu.

#### 4.4.2.2 De novo table controls - Pagination
1000 rows are visible at any one time in the Denovo table. Use the top left Pagination options to switch between result records.

#### 4.4.2.3 De novo table controls - Mass Tag Filter
Click on $\pm$ 0% to use the mass tag confidence slider filter which will replace amino acids in the sequence with their mass tag values if their local confidence does not meet the threshold.

#### 4.4.2.4 De novo table controls - Search function
Use the search function to search for scan # or precursor ID, m/z, RT, or sequence contains. Search results can be navigated between using the Up or Down arrows.
Options for searching include:

- **Scan =** Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction. For timsTOF, there is the option to search for Precursor ID instead.

- **m/z ≈:** Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/− 1 of the last significant digit.
  
  For example, searching for m/z = 580.8 will find all m/z between [580.7, 580.9]

- **RT ≈:** Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/− 1 of the last significant digit set.
  
  For example, searching for RT = 64 will find all m/z between [63, 65]

- **Seq contains:** Search for the sequence in the table. If specifying a sequence without glycans or PTMs, the results will include that sequence and any sequences with glycans or PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered.

  Additionally, since the glycan peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”

4.4.2.5 De novo table controls - Table exporting

Click on the button to export the contents of the current table. This considers sorting applied to the table and is consistent with the view. There is the option to export the De novo table and/or the De novo all candidates table in CSV format.

4.4.3 De novo View - De novo All Candidates

Click on the button to view the All de novo candidates for that spectrum. By default, the top 5 candidates are reported. The highest ALC scoring candidate is the one displayed in the De novo table.

4.4.4 PSM selection

Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.
All other components in the De novo View interface will update according to the currently selected PSM.

For timsTOF instruments, Precursor ID is used rather than Scan number.

4.4.4.1 Show spectrum in Data View

Click on to open the Data Refine MS/MS tab (if not already opened) and jump to the corresponding selected Scan.

4.4.4.2 Show spectrum in LC/MS View

Click on to open the Data Refine LC/MS tab (if not already opened) and jump to the corresponding selected Scan, highlighting the De novo marker and the feature if present.

4.4.4.3 Show Raw Spectrum View

Click on to show the raw spectrum chart of the currently selected Scan.

4.4.5 Annotated Spectrum Chart

For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

4.4.5.1 Annotated Spectrum Chart - Chart Navigation

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.
**4.4.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings**

Click on the button to show the spectrum annotation settings.

The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the "Ion Match" table, as well. It is possible to annotate the spectrum with various ion types for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.
To reset the defaults, click on the “Reset default” button.

**Show Decimal Places:** Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation:** Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated:** Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation:** Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info:** Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity:** Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

### 4.4.5.3 Annotated Spectrum Chart - Additional Chart Controls

Click on ☐ or ☒ to toggle between absolute and percentage intensity values in the Y axis.

Click on ☐ to reset the zoom level of the annotated spectrum chart to the default.

Click on ☐ to zoom into the X axis by a factor of 2.

Click on ☐ to zoom into the Y axis by a factor of 2.

The "alignment" check box ☑ allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box ☑ is used to show/hide the error map underneath the chart.

### 4.4.6 Ion Match Table

The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings" (see 4.4.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings).
Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

4.4.7 Survey Chart
The “Survey” tab displays the precursor ion spectrum. The buttons that appear in this section are identical to those explained above in 4.4.5.3 Annotated Spectrum Chart - Additional Chart Controls.

4.4.8 LC/MS Snapshot
The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The span of the orange line in the LC-MS Snapshot designates the isolation window.

Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a m/z RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.
4.4.9 Precursor Profile

The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with the retention time along the x-axis and the intensity along the y-axis.

4.5. De novo result - Feature View

The "Feature View" displays all the feature information in greater detail, as shown in the next figure. The table on the top displays features, while the bottom section provides additional information about the feature, precursor profile, and if available peptide-spectrum match.

4.5.1 Feature View - Feature table

The following list describes the contents in each of the columns of the Feature table in De novo result:

- **Fraction**: The fraction from which the feature is detected.
- **Feature ID**: An arbitrary number provided by PEAKS during data refinement for each identifiable feature.
- **m/z**: The precursor mass-to-charge ratio.
- **z**: The precursor charge.
- **RT**: The retention time (elution time) for the spectrum as recorded in the data.
- **RT Begin**: The beginning retention time (elution time) for the feature as recorded in the data.
- **RT End**: The ending retention time (elution time) for the feature as recorded in the data.
- **Area**: The area under the curve of the peptide feature found at the same m/z and retention time as the MS/MS scan. This can be used as an indicator of the abundance of the peptide.
- **Denovo Peptide**: The amino acid sequence of the peptide associated with the feature, as determined by de novo sequencing. A modified residue is followed by a pair of parentheses enclosing the mass of that modification. If multiple MS2 are associated with the feature, all de novo peptide results are listed.
- **Denovo Score (%)**: The denovo score represents the quality of the peptide spectrum match. Factors that affect this score includes ALC score and RT difference.
- **ppm**: the mass error in ppm of the de novo peptide with the highest ALC score

Click on any of the column headers to sort the column.
To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to hover mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

In the table, Amino acids in Denovo Peptide are color-coded according to their local confidence scores. Red represents a very high confidence (greater than 90%); purple represents a high confidence (80 to 90%); blue represents a medium confidence (60 to 80%); and black represents low confidence (less than 60%). Hover the cursor over the sequence to view local confidence scores of each individual amino acid.

### 4.5.2 Feature View - Feature table controls

The menu above the Denovo table provides options to scroll through the table of de novo peptides, filter for mass tags, switch between samples, search for specific entries, and export the table.

#### 4.5.2.1 Feature table controls - Sample Selection

Result for different samples can be viewed by selecting the sample using the "Show Results for Specified Sample" drop-down menu.

#### 4.5.2.2 Feature table controls - Pagination

1000 rows are visible at any one time in the Denovo table. Use the top left Pagination options to switch between result records.

#### 4.5.2.3 Feature table controls - Feature View Filter

Click on [Feature View Filter] to open the Feature View Filter dialog.
There are three options for filter provided:

- With Identification: When checked, features that are associated with a De novo peptide will be displayed in the table. Uncheck this to see all features detected in the samples.
- Feature Area ≥: Filters out all features with area below this threshold.
- Feature Quality ≥: Filters out all features with quality below this threshold. In PEAKS Studio 11, all features in a certain dataset may have quality score of 0 as there is no scoring associated with the features. It is recommended to not use this filter in that case.

Click on the **Reset** button to restore the default filters.

Click on **OK** to apply these filters to the Feature table.

Click on **Cancel** to discard changes and close the Feature View Filter dialog.

### 4.5.2.4 Feature table controls - Search function

Use the search function to search for sequence or m/z. Search results can be navigated between using the Up or Down arrows.

![Search function](image)

Options for searching include:

- m/z ≈: Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit.
  
  For example, searching for m/z ≈ 580.8 will find all m/z between [580.7, 580.9]

- Seq contains: Search for the sequence in the table. If specifying a sequence without glycans or PTMs, the results will include the bare sequence as well as sequences containing glycans or PTMs with the same peptide backbone. If specifying a sequence and a mass value within a bracket, then only sequence that has a substring with an exact match will be considered.

  Additionally, since the glycan peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”

### 4.4.2.5 Feature table controls - Table exporting

Click on the **Export** button to export the contents of the current table. This considers sorting applied to the table and is consistent with the view. There is the option to export the De novo table and/or the De novo all candidates table in CSV format.

### 4.4.2.6 Feature table - PSM visualization

The rest of the components of the Feature View are the same as the De novo View Chapter.

Those sections are:

- 4.4.4 PSM selection
• 4.4.5 Annotated Spectrum Chart
• 4.4.6 Ion Match Table
• 4.4.7 Survey Chart
• 4.4.8 LC/MS Snapshot
• 4.4.9 Precursor Profile

4.6. De novo result - Exporting

In Project View, double-click on Export to open the Export node.

There are several options for exporting the De novo results:

• De novo Summary CSV: Exports the statistics tables from the De novo Summary page into a CSV format.
• De novo CSV: Exports the De novo table. This export does not consider any sorting and will export a separate file for each sample.
• De novo CSV with all Candidates: Exports the De novo all candidates. This export does not consider any sorting and will export a separate file for each sample.
• De novo Features CSV: This export does not consider any Feature view filters or sorting, but will consider the De Novo ALC (%) filter in the Summary tab. Each sample will be in a separate export.
• Spectral Library TSV: PEAKS De Novo results can be exported as a spectral library that can be used by the Library Search engine.

For more information on how to configure a PEAKS generated spectral library, refer to Section 2.9 Spectral Library Configuration

4.7. De novo result - DeepNovo

In the PEAKS 11 upgrade release, a standalone DeepNovo workflow is enabled if the computer has a GPU selected. To run DeepNovo, open Performance Configuration and check the GPU Resource and restart PEAKS Studio. To run standard De Novo, uncheck the GPU resource.
5. Deepnovo Peptidome

5.1 Deepnovo Peptidome Analysis Workflow and Parameter Settings

To run a Deepnovo Peptidome Analysis, Select DeepNovo Peptidome in Workflow Selection

After deciding the Data Refine option, proceed to the next step to see the Deepnovo Peptidome Search parameters.

The following parameter settings are provided:

**Precursor Mass Error Tolerance:** The maximum mass difference between the peptide and the monoisotopic mass of the precursor ion.

**Precursor Mass Tolerance unit:** PEAKS generates peptides within the precursor mass error tolerance. Precursor mass error tolerance can be specified in either Daltons or ppm.
**Fragment Mass Error Tolerance:** PEAKS uses this value when scoring peptide sequences. PEAKS Studio considers a fragment ion to be matched if the calculated m/z is within the specified error tolerance. Units are Daltons.

Both precursor and fragment error tolerance parameters need to be consistent with the mass accuracy of the mass analyzers used for MS and MSMS acquisitions respectively.

**Enzyme Settings:** Select the enzyme used for protein digestion from the enzyme drop-down menu. For Deepnovo Peptide Search, only None and Trypsin are provided as Enzyme options.

Note: "None" is a special enzyme allowing non-specific cleavage at both ends of the peptide. It is the recommended option when no digestion enzyme was used or when the digestion enzyme exhibits a high degree of non-specificity.

**PTM Settings:** Click “Set PTM” to open the PTM Options dialog and specify the fixed PTMs and a few common variable PTMs expected in the experiment. Highlight PTMs and click on the button to place them as either Fixed or Variable PTM. From there, highlighted PTMs can be removed by clicking the “Remove” button. Clicking “Remove All” will reset the PTM options. Highlighting and clicking “Switch Type” will move the Fixed PTM in into the Variable PTM list and vice-versa. Click on “New” to create a new custom PTM. While a PTM is selected, click on “View” to see more details.

Note: Multiple PTMs can be highlighted by holding down the CTRL key or SHIFT key on the keyboard.

Note: For a specific residue, only one fixed modification is allowed.

Once the PTMs are selected, the maximum number of variable PTMs per peptide can be specified and a value of less than 4 is recommended.

Note: The use of variable modifications increases the size of the computational search space for the de novo sequencing algorithm. It is recommended not to select too many variable modifications in PEAKS de novo. See PEAKS PTM for alternate search options.

**Target Database:** Select a protein sequence database for the search. Select one from the list of databases that have been configured in PEAKS and set the taxonomy by clicking on , if applicable. To configure a new database, click on next to the Target Database dropdown to open the Database Configuration dialog.

**Contaminant Database:** To search for contaminants, select a contaminant database from the list of databases that have been configured in PEAKS. Keep the selection as N/A if no contaminant database is required for the search.

Click on “Report” to proceed to the final step. Report filters can be set here to filter the final result.
These filters are provided:

**PSM/Peptide:** Select one of the options to filter the PSMs or Peptides.

**Deep Novo:** Set a Deep Novo ALC(%) filter to filter the Deep novo peptides. This number can be adjusted in the Deep novo Summary tab to filter the Deep novo table.

Finally, there is an option to specify the Analysis Name (which can also be renamed post-analysis) and/or an option to save the Workflow for a timely re-use in future analyses.

### 5.2 Understanding PEAKS Deep Novo Results

When Deep Novo sequencing is done, double-click on the result node to open the Deep novo result. The result contains two tabs:

- **Summary:** This tab shows an outline of the PEAKS deep novo search results with key statistics and chart visualizations. The overall quality of the experiment can be examined and the filters for deep novo ALC can be adjusted at the top of the page.
• Deep Novo: This tab shows a list of peptide sequences detected by de novo sequencing, which passed the set ALC score threshold.

The following sections will explore each of these two tabs in the Deep Novo result.

5.3 Deep Novo result - Summary tab

In the summary tab, PEAKS Studio de novo sequencing result can be filtered using Average Local Confidence (ALC) score. Low quality de novo sequences can be filtered out by specifying a minimal threshold of the ALC score. The purpose of filtering is to remove low confidence sequences in which residues are likely incorrectly identified. By default, the ALC threshold is set to 50%. When changing this threshold, click [Apply] to apply the new filter. The result in the Denovo result view will be updated accordingly.

1. Notes

2. Result Statistics

Table 1. Statistics of data and result

<table>
<thead>
<tr>
<th>Sample Name</th>
<th># Precursors</th>
<th># PSMs</th>
<th># PSMs (ab &gt; 3x)</th>
<th># PSMs (ab &gt; 5x)</th>
<th># PSMs (ab &gt; 10x)</th>
<th># PSMs (ab &gt; 20x)</th>
<th># PSMs (ab &gt; 30x)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AB</td>
<td>28143</td>
<td>18117</td>
<td>11645</td>
<td>6021</td>
<td>3019</td>
<td>149.0%</td>
<td></td>
</tr>
<tr>
<td>Sample 2</td>
<td>19492</td>
<td>10310</td>
<td>5798</td>
<td>1810</td>
<td>144.0%</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Click on [Notes] to open the Notes dialog to save useful information regarding this Analysis. The Notes will appear in Section 1 of the summary page at the top.

Section 2 Result Statistics gives key statistics in Table 1 by providing the total number of MS2 scans and PSMs total as well as the number of MS2 scans and PSMs with different ALC thresholds for each sample. The #PSM / #MS2 provides a de novo sequencing identification rate metric for each sample. For timTOF data, Table 1 shows # Precursors instead.

5.3.1 Deep Novo result - Summary tab - ALC versus precursor mass error scatterplot

Figure 1. a) Scatterplot of peptide ALC score versus precursor mass error shows the distribution of precursor mass error according to their ALC score. Red indicates a higher concentration of peptides at that mass error and ALC score, and green indicates a lower concentration. A vertical dashed line shows the current Deep Novo ALC(%) filter applied.
5.3.2 Deep Novo result - Summary tab - Distribution of local residue confidence

Figure 1. b) Distribution of local residue confidence shows the overall distribution of local confidence scores of residues across all peptides.

5.3.3 Deep Novo result - Summary tab - ALC RT distribution of first fraction

Similar to Figure 1, Figure 2. a) ALC Filtered RT figures of first fraction shows the distribution of $\Delta$RT (min) according to their ALC score. Red indicates a higher concentration of peptides at that $\Delta$RT and ALC score, and green indicates a lower concentration. A vertical dashed line shows the current Deep Novo ALC(%) filter applied.

5.3.4 Deep Novo result - Summary tab - Peptide RT versus Predicted RT scatter plot

Figure 2. b) Scatterplot of peptide RT versus Predicted RT figure shows the scatter plot of predicted RT versus actual RT in minutes for the first fraction. A regression function is shown to fit the plot.
Table 2 simply shows what is the current Deep novo ALC filter set. This is placed here so that when the Summary page is exported, the viewer will know what the ALC threshold was at the time.

Section 3 Other Information shows the parameters that were set when the analysis was ran.

5.4 Deep Novo result - Deep Novo View

The "Deep Novo View" displays the deep novo sequencing results in greater detail, as shown in the next figure. The table on the top displays deep novo sequences, while the bottom section provides additional information about the peptide-spectrum match.
5.4.1 Deep Novo View - Deep Novo table

The following list describes the contents in each of the columns of the Denovo table:

- **Scan**: The scan number and data file it belongs to. The scan number is a unique index for tandem mass spectra in the data.
- **Peptide**: The amino acid sequence of the peptide as determined by de novo sequencing. A modified residue is followed by a pair of parentheses enclosing the mass of that modification.
- **ALC(%)**: The average local confidence. ALC is calculated as the total of the residue local confidence scores in the peptide divided by the peptide length. The higher the score the more confident the denovo candidate.
- **Precursor Id**: This column is only displayed for timsTOF data. A unique identifier is assigned for each precursor obtained from the raw data.
- **Length**: The number of amino acids in the sequence backbone.
- **m/z**: The precursor mass-to-charge ratio.
- **z**: The precursor charge.
- **RT**: The retention time (elution time) of the spectrum as recorded in the data.
- **Δ RT**: The difference in retention time from the data and the RT prediction in deep learning algorithm.
- **1/k0 Range**: This shows the ion mobility range associated with precursor.
- **Δ 1/k0**: The difference in 1/k0 from the data and the 1/k0 prediction in deep learning algorithm.
- **Area**: The area under the curve of the peptide feature found at the same m/z and retention time as the MS/MS scan. This can be used as an indicator of the abundance of the peptide.
- **Mass**: The monoisotopic mass of the peptide.
- **ppm**: The precursor mass error, calculated as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$.
- **PTM**: The types and the numbers of modifications present in the peptide shown in color-coded icons.
- **Mode**: The fragmentation mode in which de novo sequencing is performed by the algorithm.

Click on any of the column headers to sort the column.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

In the table, Amino acids in de novo sequences are color-coded according to their local confidence scores. Red represents a very high confidence (greater than 90%); purple represents a high confidence (80 to 90%); blue represents a medium confidence (60 to 80%); and black represents low confidence (less than 60%). Hover the cursor over the sequence to view local confidence scores of each individual amino acid.

5.4.2 Deep Novo View - Deep Novo table controls

The menu above the Deep Novo table provides options to scroll through the table of deep novo peptides, filter for mass tags, switch between samples, search for specific entries, and export the table.
5.4.2.1 Deep Novo table controls - Sample Selection
Result for different samples can be viewed by selecting the sample using the "Show Results for Specified Sample" drop-down menu.

Show Results for Specified Sample  Sample 1

5.4.2.2 Deep Novo table controls - Pagination
1000 rows are visible at any one time in the Deep Novo table. Use the top left Pagination options to switch between result records.

5.4.2.3 Deep Novo table controls - Mass Tag Filter
Click on [ ] 0% to use the mass tag confidence slider filter which will replace amino acids in the sequence with their mass tag values if their local confidence does not meet the threshold.

5.4.2.4 Deep Novo table controls - Search function
Use the search function to search for scan # or precursor ID, m/z, RT, or sequence contains. Search results can be navigated between using the Up or Down arrows.

seq contains: [ ] TIP

Options for searching include:

- **Scan =**: Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction. For timsTOF, there is the option to search for Precursor ID instead.

- **m/z ≈**: Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit.
  
  For example, searching for m/z ≈ 580.8 will find all m/z between [580.7, 580.9]

- **RT ≈**: Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/- 1 of the last significant digit set.
  
  For example, searching for RT ≈ 64 will find all m/z between [63, 65]

- **Seq contains**: Search for the sequence in the table. If specifying a sequence without glycans or PTMs, the results will include that sequence and any sequences with glycans or PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered.
  
  Additionally, since the glycan peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”

5.4.2.5 Deep Novo table controls - Table exporting
Click on the [ ] button to export the contents of the current table. This considers sorting applied to the table and is consistent with the view. There is the option to export the Deep Novo table and/or the Deep novo all candidates table in CSV format.
5.4.3 Deep Novo View - Deep Novo All Candidates

Click on the button to view the All deep novo candidates for that spectrum. By default, the top 5 candidates are reported. The highest ALC scoring candidate is the one displayed in the De novo table.

5.4.4 PSM selection

Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.

All other components in the De novo View interface will update according to the currently selected PSM.

For timsTOF instruments, Precursor ID is used rather than Scan number.

5.4.4.1 Show spectrum in Data View

Click on to open the Data Refine MS/MS tab (if not already opened) and jump to the corresponding selected Scan.

5.4.4.2 Show spectrum in LC/MS View

Click on to open the Data Refine LC/MS tab (if not already opened) and jump to the corresponding selected Scan, highlighting the De novo marker and the feature if present.

5.4.4.3 Show Raw Spectrum View

Click on to show the raw spectrum chart of the currently selected Scan.

5.4.5 Annotated Spectrum Chart

Click on button to switch from the Mirror plot view and annotated spectrum chart view.
For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

5.4.5.1 Annotated Spectrum Chart - Chart Navigation

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.

Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.

Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.

See the whole spectrum: Double click in the spectrum or click the 1:1 button.

5.4.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings

Click on the button to show the spectrum annotation settings.
The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ion types for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places:** Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation:** Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated:** Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation:** Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info:** Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity:** Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

### 5.4.5.3 Annotated Spectrum Chart - Additional Chart Controls

Click on to toggle between absolute and percentage intensity values in the Y axis.

Click on to reset the zoom level of the annotated spectrum chart to the default.

Click on to zoom into the X axis by a factor of 2.

Click on to zoom into the Y axis by a factor of 2.
The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

5.4.5.4 Annotated Spectrum Chart - Mirror Plot

Deep Novo results have a Mirror Plot view. Click on the button to switch to the Mirror Plot.

This view shows the ions from the query spectrum compared to the predicted spectrum.

Hold down CTRL when scrolling the mouse wheel to zoom only on the top half or bottom half of the mirror plot.

5.4.6 Ion Match Table

The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated fragment mass, where mass values displayed in color indicate fragment ions present in the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings" (see 4.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings).
Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

### 5.4.6.1 Ion Match Table - Predicted matching table

While in mirror plot view, the Ion Match table updates to display the ion matches between fragment query ions and predicted ions.

<table>
<thead>
<tr>
<th>#</th>
<th>Label</th>
<th>Predicted M/Z</th>
<th>Predicted Intensity (%)</th>
<th>Query M/Z</th>
<th>Query Intensity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>b2</td>
<td>259.15</td>
<td>6.67</td>
<td>258.14</td>
<td>4.06E(6.80%)</td>
</tr>
<tr>
<td>2</td>
<td>y2</td>
<td>310.18</td>
<td>48.38</td>
<td>310.17</td>
<td>2.02E(6.19%)</td>
</tr>
<tr>
<td>3</td>
<td>b3</td>
<td>345.16</td>
<td>5.21</td>
<td>345.17</td>
<td>5.24E(7.59%)</td>
</tr>
<tr>
<td>4</td>
<td>y3</td>
<td>409.24</td>
<td>26.13</td>
<td>409.24</td>
<td>1.51E(7.17%)</td>
</tr>
<tr>
<td>5</td>
<td>b4</td>
<td>508.24</td>
<td>7.93</td>
<td>508.24</td>
<td>3.23E(4.66%)</td>
</tr>
<tr>
<td>6</td>
<td>y4</td>
<td>572.81</td>
<td>8.81</td>
<td>572.86</td>
<td>3.88E(9.51%)</td>
</tr>
<tr>
<td>7</td>
<td>b5</td>
<td>595.27</td>
<td>12.40</td>
<td>595.27</td>
<td>7.17E(10.35%)</td>
</tr>
<tr>
<td>8</td>
<td>y5</td>
<td>671.38</td>
<td>14.42</td>
<td>671.37</td>
<td>9.82E(14.19%)</td>
</tr>
<tr>
<td>9</td>
<td>b6</td>
<td>694.34</td>
<td>55.14</td>
<td>694.33</td>
<td>2.81E(46.55%)</td>
</tr>
<tr>
<td>10</td>
<td>y6</td>
<td>758.41</td>
<td>41.06</td>
<td>758.40</td>
<td>2.88E(15.50%)</td>
</tr>
<tr>
<td>11</td>
<td>b7</td>
<td>857.40</td>
<td>63.27</td>
<td>857.40</td>
<td>4.37E(8.05%)</td>
</tr>
</tbody>
</table>

Predicted intensity and Query intensity are reported relative to the highest intensity ion. All columns can be rearranged and sorted.

### 5.4.7 Survey Chart

The “Survey” tab displays the precursor ion spectrum. The buttons that appear in this section are identical to those explained above in 4.4.5.3 Annotated Spectrum Chart - Additional Chart Controls.

![Survey Chart](image)

### 5.4.8 LC/MS Snapshot

The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The span of the orange line in the LC-MS Snapshot designates the isolation window.
Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a m/z RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

5.4.9 Precursor Profile

The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with the retention time along the x-axis and the intensity along the y-axis. The first two isotopes are also drawn as M+1 and M+2 respectively.

5.5. Deep Novo result - Exporting

In Project View, double-click on to open the Export node.
There are several options for exporting the Deep Novo results:

- **DeepNovo Summary HTML**: Exports the Summary page including all figures in HTML format. The content will be the same as shown in PEAKS Studio.
- **DeepNovo CSV**: Exports the Deep Novo table. This export does not consider any sorting and will export a separate file for each sample.
- **DeepNovo CSV with all Candidates**: Exports the Deep Novo all candidates. This export does not consider any sorting and will export a separate file for each sample.
- **Spectral Library TSV**: PEAKS DeepNovo results can be exported as a spectral library that can be used by the Library Search engine.
  
  For more information on how to configure a PEAKS generated spectral library, refer to Section 2.9 Spectral Library Configuration.

### 5.6 Peptide Search result

Double-click on the result node to open the result. The analysis results for peptide search tools are presented in two tabs:

- **Summary**: This tab shows an outline of the PEAKS Studio peptide search results with key statistics. The overall quality of the experiment can be examined and the filters for peptide and protein identifications can be adjusted. This page shows after the search is done.
- **Peptide**: This tab shows a list of peptide identifications. This view also provides spectrum annotation and other detailed information for peptide precursor spectrum matches.

The following sections will explore each of these two tabs in the Identification result.

### 5.6.1 Peptide Search result - Summary tab

The Summary view reports key statistics as an overview of the result. In the report, several statistical charts can be examined to assess result quality.
1. Notes

2. Result Statistics

Figure 1. False discovery rate (FDR) curve. X axis is the number of peptides being kept. Y axis is the corresponding FDR.

Click on [Notes] to open the Notes dialog to save useful information regarding this Analysis. The Notes will appear in Section 1 of the summary page at the top.

The Summary view reports key statistics as an overview of the result. In the report, several statistical charts can be examined to assess result quality.

5.6.1 Peptide Search result - Figure 1. False Discovery Rate (FDR) curve.

Figure 1 shows the FDR curve with respect to the number of PSMs being kept after filtration by the peptide -10lgP score. By lowering the score threshold, more PSMs are kept in the filtered result. Conversely, the FDR increases because more false positives are kept. In this figure, the vertical line indicates the current score threshold. The number of PSMs and the corresponding FDR at the current score threshold are shown in the top-left corner.

PEAKS Search estimates FDR using the "decoy-fusion" approach. Decoy-fusion is an enhanced target-decoy approach that makes more conservative FDR estimations. For details of the "decoy-fusion" approach, please refer to the publication, "PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification", Mol Cell Proteomics, 2011 Dec 20.

It is commonly recommended to set FDR at <1% by adjusting the score threshold. If a rapid growth of the FDR around the 1% FDR threshold is noticed, some PSMs may be sacrificed to significantly reduce the FDR.

**Note:** When counting the number of PSMs, PEAKS Studio keeps one peptide per spectrum at most. Thus, the number of PSMs is actually the number of spectra with assigned peptides.

**Note:** During Analysis creation, if Peptide FDR (%) filter was used in Report step, the Summary page FDR curve in the results will show the Peptide FDR curve and corresponding number of Identified peptides.

**Note:** Decoy matches are not counted in the number of PSMs. Unless otherwise specified, decoy matches are also excluded from the other statistical numbers shown in the Summary view.
5.6.1.2 Peptide Search result - Table 1. Statistics of data
Statistics of Data tabulates MS1 and MS2 information, as well as identification in the peptide and protein level. This table in the Summary view provides a condensed statistical information for the analysis.

Table 1. Statistics of data.

<table>
<thead>
<tr>
<th></th>
<th>#MS Run</th>
<th>#Frames</th>
<th>#Precursors</th>
<th>#PSMs</th>
<th>#Peptides</th>
<th>#Sequences</th>
<th>#PSM/#Precursors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>9</td>
<td>6432</td>
<td>5252</td>
<td>124115</td>
<td>47300</td>
<td>47799</td>
<td>40088</td>
</tr>
<tr>
<td>Sample 1</td>
<td>3</td>
<td>2005</td>
<td>1799</td>
<td>41228</td>
<td>15724</td>
<td>21806</td>
<td>18964</td>
</tr>
<tr>
<td>Sample 2</td>
<td>3</td>
<td>2097</td>
<td>1771</td>
<td>40286</td>
<td>16393</td>
<td>22174</td>
<td>19140</td>
</tr>
<tr>
<td>Sample 3</td>
<td>3</td>
<td>2330</td>
<td>1682</td>
<td>42601</td>
<td>15183</td>
<td>21797</td>
<td>19049</td>
</tr>
</tbody>
</table>

#MS Run: The number of data files in each sample

#MS1: Total # of MS1 spectra for each sample. For timsTOF data this shows the # MS1 frames.

#MS/MS: Total # of MS2 spectra for each sample. This does not include chimeric scans. For timsTOF data this shows the # MS2 frames.

#Precursors: The number of precursors in the result. Only displayed for timsTOF data.

#PSMs: The number of peptide-spectrum matches.

#Peptides: The unique number of peptide sequences with modifications not including I/L differentiation.

#Sequences: The unique number of peptide sequences not including modifications and I/L differentiation.

#PSM/MS2: The identification rate calculated by dividing #PSM by the #MS2. For timsTOF data, it is calculated by dividing #PSM by the #Precursors.

5.6.1.3 Peptide Search result - Figure 2. PSM score distribution
Figure 2 in the Summary View show the precursor mass errors of PSMs in filtered results. It shows the distribution of the precursor mass errors in a histogram with and without software calibration. The precursor mass error is calculated in ppm as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$. Calibration is for display purposes only and PEAKS corrects m/z value for each PSM to correct systematic biases and instrument measurement drift. For a well-calibrated instrument, precursor mass errors should center at 0 ppm across the range of m/z.

**Figure 2. Distribution of precursor mass error of filtered PSM in ppm.**
5.6.1.4 Peptide Search result - Result filtration parameters

Tables 2 show the statistical numbers of the data and results based on report filters set.

<table>
<thead>
<tr>
<th>Table 2. Statistics of filtered results.</th>
</tr>
</thead>
<tbody>
<tr>
<td>DB Peptides</td>
</tr>
<tr>
<td>Homolog Peptides</td>
</tr>
<tr>
<td>DeepNovo Peptides</td>
</tr>
<tr>
<td>FDR (Peptide-Spectrum Matches)</td>
</tr>
<tr>
<td>FDR (Peptide Sequences)</td>
</tr>
<tr>
<td>-10LgP Cutoff (Peptide-Spectrum Matches)</td>
</tr>
</tbody>
</table>

5.6.1.5 Peptide Search result - RT Figures

RT figures with outliers removed are displayed for the first data file in the analysis.

Figure 3(a) shows the ΔRT distribution with a histogram. Peptides found by Database Search, DeepNovo, and Homolog are separated into colour-coded bars. Figure 3(b) shows the alignment of predicted RT and real RT in minutes. The peptides found are split into different colour-coded points in the scatterplot and a regression function is drawn to indicate the correlation.

Figure 3. (a) Histogram of peptide ΔRT. (b) Scatterplot of peptide RT versus Predicted RT.

RT figures for 20180924_50ngHeLa_1.0.25.1_Hystar5_0SR1_S2-A1_1_2042.d with outliers removed

For timsTOF data, a Regression plot of 1/k0 is also provided in Figure 4.
Figure 4. Regression plot of peptide 1/k0 versus Predicted 1/k0.

5.6.1.6 Peptide Search result - Boxplot
Figure 5 shows a boxplot of the median ΔRT across all MS runs.

5.6.1.7 Peptide Search result - Other Information
Table 3 summarizes the parameters defined by the user to analyze the data. This information includes error mass tolerances, database(s) used, etc.
3. Other Information

Table 3. Search parameters:
- Search Engine Name: PEAKS
- Parent Mass Error Tolerance: 20.0 ppm
- Fragment Mass Error Tolerance: 0.05 Da
- Enzyme: Trypsin
- Fixed Modifications:
  - Carbamidomethylation (+57.02)
- Variable Modifications:
  - Deamidation (NQ) (+0.98)
  - Oxidation (M) (+15.99)
- Max Variable PTM Per Peptide: 2
- Database: uniprot_sprot_June-2-2020.fasta
- Taxon: Homo sapiens (Human)
- Searched Entries: 20376
- Deep Novo ALC: 50.0%
- Deep Novo Protein Association Tag Sharing: 5

5.6.2 Peptide Search result - Peptide Tab

The Peptide Table shows the filtered native peptide identification results. Each row in the table is a native peptide identification represented by its highest-scoring PSM. When there are more than 1000 peptides, the table is split into multiple pages.

Note: If there are multiple PSMs matched to a peptide, the table displays the information for the top-scoring PSM. Peptides with the same sequence but different modifications are considered different and shown in different rows.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.
The following list describes the contents in each of the default columns:

- **Peptide**: The amino acid sequence of the peptide, as determined by the PEAKS search workflow. A modified residue is followed by a pair of parentheses enclosing the modification.
- **-10lgP**: The peptide -10lgP score. The score indicates the scoring significance of a peptide-precursor spectrum match.
- **Mass**: The monoisotopic mass of the peptide.
- **Length**: The number of amino acids in the sequence backbone.
- **ppm**: The precursor mass error, calculated as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$.
- **m/z**: The precursor mass-to-charge ratio.
- **RT**: The retention time (elution time) of the spectrum as recorded in the data.
- **ΔRT**: The difference between predicted RT and RT of peptide.
- **1/k0**: The 1/k0 range for timsTOF data only.
- **Δ1/k0**: The difference between average predicted 1/k0 and 1/k0 of peptide for timsTOF data.
- **Area per Sample**: The combined area for all features in that sample associated with this peptide.
- **Scan / Precursor Id**: The scan number of the spectrum that matches the peptide sequence with the highest -10lgP. For timsTOF, this is the Precursor Id instead.
• **#Feature:** For single sample analysis, shows number of features assigned to the peptide sequence from identification.

• **Accession:** The accessions of proteins that contain this peptide.

• **PTM:** The types and the numbers of modifications present in the peptide shown in color-coded icons.

• **AScore:** Localization score assigned to modifications on the peptide. It is the \(-10\) log of a p-value. In this case, the p-value is the probability that the modification occurs at the reported position compared to other possible positions. A \(-10\log P\) of 20 is equal to a p-value of 0.01. Scores are listed in the format amino acid, amino acid residue position, modification, AScore. For example, serine at amino acid 3 is phosphorylated with an AScore of 54.01 will be reported as: S3 P 54.01.

• **Found By:** The name of the PEAKS search workflow that identifies the peptide; this can be either Database Search, Homolog, or DeepNovo.

5.6.3 Peptide Table controls
The menu above the peptide table provides options to scroll through the table of peptides, filter the table, show optional columns and search for specific entries.

5.6.3.1 Peptide Table controls - Pagination
For easy viewing and to improve performance, the peptide table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.

In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

5.6.3.2 Peptide Table controls - View Filter
Click on the button to open the Peptide View Filter which is used to filter the Peptide table. The following are the filter options provided:
• Peptide sequence contains: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains for further filtering if necessary.

• Peptide score (-10lgP): Filters out all peptides with -10lgP scores below this threshold.

• Peptide length: Filters for peptides of length within this min/max threshold.

• Peptide sample area: Filters based on total peptide area found in each sample. Sample areas are calculated using the total of all peptide features from all spectra that identified the native peptide. If the peptide is found in multiple samples, only one sample must pass this filter for the peptide to be included.

• PTM contains: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text they will all be included in the table.

• Show FAIMS CVs: For FAIMS data, only keep Peptides that have PSMs with CV values selected.

• Found By: A dropdown that allows selection of only keeping peptides found by Database Search, DeepNovo, or Homolog.

• CAA (%): Filters for peptides with minimum CAA (%) threshold.

• MS2 Correlation: Filters for peptides with minimum MS2 Correlation threshold.
• ΔRT: The filter for the range of RT difference.

• Δ1/k0: The filter for the range of 1/k0 difference. Only shown for timsTOF data.

Note: In all text filters, multiple entries can be separated by a semi-colon.

Click on the Reset button to reset all filters to their default setting.

Click on OK to apply the filters to the Peptide table.

Click on Cancel to discard the changes in the View Filter.

### 5.6.3.3 Peptide Table controls - Optional Columns

Click on the button to open the optional columns dropdown. By default the sample area columns are displayed. Sample Feature count columns can also be shown. These sample feature counts are always from identification, whether or not there are feature vectors present. For projects with many samples, it might be useful to hide these optional columns and reduce the width of the peptide table.

### 5.6.3.4 Peptide Table controls - Search function

On the top right of menu, there is a search function with 4 options. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- **Scan =**: Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction. For timsTOF data, this search function is for Precursor Id rather than Scan.

- **m/z ≈**: Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit. For example, searching for m/z ≈ 560.21 will find all m/z between [560.20, 560.22]

- **RT ≈**: Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/- 1 of the last significant digit set. For example, searching for RT ≈ 55.5 will find all m/z between [55.4, 55.6]
- **Seq contains**: Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered. Additionally, since the peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVK”, “IVLK”, and “IVIK”

### 5.6.4 Feature selection breadcrumb

There is a sample and feature selection breadcrumb which allows you to switch between features associated with the peptide. Click on the Sample in the breadcrumb to reveal other samples and the features in that sample related to the current peptide selection. From here you can select other samples, fractions, or features.

By clicking on the fraction level, all fractions in this sample are revealed as well as the features in each fraction associated with this peptide. From here you can select other fractions and features to view.

Lastly, clicking on the feature level will show all features in the currently selected fraction. Features with Feature Id = 0 are known as fake features and are constructed to enhance identification.

### 5.6.5 Peptide View - PSM selection

Underneath the feature breadcrumb will shown the currently selected PSM. Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.

All other components in the Peptide tab interface will update according to the currently selected PSM.

For timsTOF data, Precursor Id is used rather than Scan.
5.6.5.1 Show spectrum in Data View
Click on to open the Data Refine MS/MS tab (if not already opened) and jump to the corresponding selected Scan.

5.6.5.2 Show spectrum in LC/MS View
Click on to open the Data Refine LC/MS tab (if not already opened) and jump to the corresponding selected Scan, highlighting the peptide MS/MS marker and the feature if present.

5.6.5.3 Show Raw Spectrum View
Click on to show the raw spectrum chart of the currently selected Scan.

5.6.6 Peptide View - Annotated Spectrum Chart
Click on button to switch from the Mirror plot view and annotated spectrum chart view.

For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

5.6.6.1 Annotated Spectrum Chart - Chart Navigation
Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.
Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.

Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.

See the whole spectrum: Double click in the spectrum or click the 1:1 button.

5.6.6.2 Annotated Spectrum Chart - Spectrum Annotation Settings

Click on the button to show the spectrum annotation settings.

The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places:** Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation:** Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated:** Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation:** Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info:** Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.
**Intensity**: Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

### 5.6.6.3 Annotated Spectrum Chart - Additional Chart Controls
Click on ∧ or ∨ to toggle between absolute and percentage intensity values in the Y axis.

Click on ▼ to reset the zoom level of the annotated spectrum chart to the default.

Click on × to zoom into the X axis by a factor of 2.

Click on ÷ to zoom into the Y axis by a factor of 2.

The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

### 5.6.6.4 Annotated Spectrum Chart - Mirror Plot
Spectral Library results have a Mirror Plot view. Click on the button to switch to the Mirror Plot.

This view shows the ions from the query spectrum compared to the library spectrum.

Hold down CTRL when scrolling the mouse wheel to zoom only on the top half or bottom half of the mirror plot.

### 5.6.6.5 Peptide View - Ion Match Table
The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings".
Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

5.6.7 Peptide View - LC/MS Snapshot

The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The peptide feature is highlighted in purple and the selected PSM marker is blue. The span of the orange line in the LC-MS Snapshot designates the isolation window.

Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a m/z RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

5.6.8 Peptide View - Precursor Profile

The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with
the retention time along the x-axis and the intensity along the y-axis. The first two isotopes are displayed in the chart as well as M+1 and M+2.

5.7 Peptide Search result - Exporting
In Project View, double-click on to open the Export node.

5.7.1 Peptide Search result - Exporting - comma separated values (CSV) format
There are several options for exporting the PEAKS Identification results in CSV format:

- **Peptide CSV**: All of the identified peptides and details will be saved to peptides.csv.
- **PSM CSV**: The peptide-spectrum matches (PSMs) will be exported to peptideDn.psms.csv. Peptides differentiated with only I/L isoform are represented by separate entries. As a result, the number of entries in this file might be bigger than the number of PSMs mentioned in the Summary view.

5.7.2 Peptide Search result - Exporting - HTML format
**Export in HTML**: This will generate single or multiple HTML reports in the specified location.

The following exporting options are available:

- **Summary view**: The “Summary” view page will be saved as a summary.html file in HTML format in the specified location.
- **Export peptide list in html**: creates a separate HTML-containing peptide identifications.

5.7.3 Peptide Search result - Exporting - Spectral Library generation

Spectral Library TSV: PEAKS peptidome identification results can be exported as a spectral library that can be used by the Library Search engine. Select “Use original RT” to keep original RT values in the library. Select “Map to iRT”
to save iRT values in the library. The map to iRT function maps experimental retention times for peptides to an indexed RT (iRT) to account for chromatographic differences between runs (e.g. different gradient lengths).

For more information on how to configure a PEAKS generated spectral library, refer to Section 2.9 Spectral Library Configuration
6. Peptide, PTM, and Mutation Identification (PEAKS DB, PEAKS PTM, and SPIDER)

PEAKS Studio provides a complete set of database search tools for in-depth protein analysis. With PEAKS DB, PEAKS PTM, and SPIDER, proteins in the sample can be identified with high sensitivity and accuracy. It is also possible to find potential sites for modification and mutation. PEAKS Studio supports multiple enzyme digestion by which almost full sequence coverage can be achieved for single protein studies. PEAKS Studio database search tools have built-in result validation using an enhanced target-decoy approach. The false discovery rate (FDR) is estimated to ensure that only valid results are reported. PEAKS DB is a database search tool assisted by PEAKS de novo sequencing to achieve high sensitivity and accuracy.

**Note:** For more details, refer to the paper: "PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification", Mol Cell Proteomics, 2011, Dec 20.

PEAKS PTM is a dedicated search tool for peptides with unspecified modifications and mutations. It can search with unlimited number of modifications, allowing searches with all modifications in the Unimod database.

**Note:** For more details, refer to the paper: "PeaksPTM: Mass Spectrometry Based Identification of Peptides with Unspecified Modifications", Journal of Proteomics Research, 2011, 10(7): 2930-2936.

SPIDER is a dedicated search tool for finding novel peptides that are homologous to peptides in a given protein database.

**Note:** For more details, refer to the paper: "SPIDER: Software for Protein Identification from Sequence Tags Containing De Novo Sequencing Error", J Bioinform Comput Biol, 2005, Jun;3(3):697-716.

6.1 Database Search Workflow and Parameter Settings

PEAKS DB, PEAKS PTM, and SPIDER can be launched together as a work flow in a PEAKS Studio analysis. Users can also do PEAKS DB only, PEAKS DB plus PEAKS PTM, or PEAKS DB plus SPIDER.

To run a Database Search workflow, select PEAKS DB (In-depth de novo assisted search) in the Project Wizard.

After deciding the Data Refine option, proceed to the next step to see the DB Search parameters.
The following parameter settings are provided:

**Precursor Mass Error Tolerance**: The maximum mass difference between the peptide and the monoisotopic mass of the precursor ion.

**Precursor Mass Tolerance unit**: PEAKS generates peptides within the precursor mass error tolerance. Precursor mass error tolerance can be specified in either Daltons or ppm.

**Fragment Mass Error Tolerance**: PEAKS uses this value when scoring peptide sequences. PEAKS Studio considers a fragment ion to be matched if the calculated m/z is within the specified error tolerance. Units are Daltons.

Both precursor and fragment error tolerance parameters need to be set consistently with the mass accuracy of the instrument.

**Enzyme Settings**: PEAKS Studio digests the protein database in silico to generate peptide candidates. It is necessary to specify the enzyme for protein digestion from the Enzyme drop-down menu.
**Note:** When the selected dataset is digested with different enzymes, "Specified by each sample" allows samples to be analyzed separately using their respective enzymes specified during project creation.

**Note:** "None" is a special enzyme allowing non-specific cleavage at both ends of the peptide. It is the recommended option when no digestion enzyme was used or when the digestion enzyme exhibits a high degree of non-specificity.

**Missed Cleavages:** This specifies the number of missed cleavage sites allowed in a peptide.

**Digest Mode:** This specifies the type of in silico cleavages allowed at a peptide termini based on the method used for protein digestion. If "Specific" is selected, the specificity of the selected enzyme is strictly enforced at both termini of a peptide. "Semispecific" allows one terminus to disobey the enzyme specificity rules.

**Note:** If the enzyme is specified as "None", then no matter which mode is selected for "Digest Mode", the "None" enzyme digest rule will be applied.

**PTM Settings:** Click “Set PTM” to open the PTM Options dialog and specify the fixed PTMs and a few common variable PTMs expected in the experiment. Highlight PTMs and click on the ➡️ button to place them as either Fixed or Variable PTM. From there, highlighted PTMs can be removed by clicking the “Remove” button. Clicking “Remove All” will reset the PTM options. Highlighting and clicking “Switch Type” will move the Fixed PTM in into the Variable PTM list and vice-versa. Click on “New” to create a new custom PTM. While a PTM is selected, click on “View” to see more details.

**Note:** Multiple PTMs can be highlighted by holding down the CTRL key or SHIFT key on the keyboard.

**Note:** For a specific residue, only one fixed modification is allowed.

Once the PTMs are selected, the maximum number of variable PTMs per peptide can be specified and a value of less than 4 is recommended.

**Note:** The use of variable modifications increases the size of the computational search space for the de novo sequencing algorithm. It is recommended not to select too many variable modifications in PEAKS de novo. See PEAKS PTM for alternate search options.

**Target Database:** Select a protein sequence database for the search. Select one from the list of databases that have been configured in PEAKS and set the taxonomy by clicking on ➜️, if applicable. To configure a new database, click on ➜️ next to the Target Database dropdown to open the Database Configuration dialog.

**Contaminant Database:** To search for contaminants, select a contaminant database from the list of databases that have been configured in PEAKS. Keep the selection as N/A if no contaminant database is required for the search.

**Peptide Length:** The length for a peptide can be set when creating an analysis, where the default peptide length range is between 6 and 45.
**Deep Learning Boost:** Check this option to use PEAKS enhanced deeplearning algorithm to better identify PSMs in the DB Search process.

**Find Unspecified PTMs with PEAKS PTM:** Select this option to enable a PEAKS PTM search after a PEAKS DB search finishes. PEAKS PTM analyzes spectra with good de novo sequences that remain unidentified by PEAKS DB. The default setting for PEAKS PTM is to search with all the built-in modifications in the "Common" and "Uncommon" lists, which include all of the natural modifications in the Unimod database.

Alternatively, PEAKS PTM can search with a list of preferred modifications. Once "Search with preferred modifications" has been selected, click the Additional Modifications button to select desired PTMs for the PTM search. Importantly, PEAKS PTM allows an unlimited number of variable modifications to be searched, however, limiting the number can improve search accuracy.

Check “Find Potential Signature Ions” to add potential signature ion detection and report the potential signature ion statistics in the summary page. Potential signature ions are ions that meet the signature ion detection algorithm set in section 2.5 Algorithm Configuration. Expected signature ions, which are defined as signature ions in PTMs that were selected in the analysis, will always be displayed. Signature ions can be specified when creating custom Modifications. See 2.4 PTM Configuration for configuring custom PTMs.

By default, PEAKS PTM considers the same number of maximum number of variable PTMs per peptide as set in the DB search parameters. A threshold on the de novo ALC score can be specified so that only the spectra with good de novo sequences are analyzed by PEAKS PTM. In the PEAKS PTM result node, search results from PEAKS DB and PEAKS PTM are combined and shown.

**Find More Mutations with SPIDER:** Select this option to enable a SPIDER search. SPIDER performs homology searches on spectra with good de novo sequences (i.e. ALC>15%) that remain unidentified by either PEAKS DB and/or PEAKS PTM. SPIDER identifies novel peptides that are homologous to peptides in the searched protein database. SPIDER is good for cross-species searches and for finding point mutations of the protein. In the SPIDER result page, search results from PEAKS DB, PEAKS PTM (if applicable), and SPIDER are combined and shown.

In PEAKS Studio 11, the next optional step is the newly introduced QC function. See section 6.12 Identification Quality Control (QC) for further explanation on how this function is utilized.
When QC is checked, a QC result node is added in the analysis. Select a control sample and acceptance tolerance, where the tolerance will mark a sample as pass or fail depending on the relation to the control sample. These two cannot be adjusted in the QC result node, and a new analysis would have to be run. The Data and Identification QC attributes are the attributes of interest. If a sample is outside the acceptance tolerance in any of these attributes with respect to the control sample, it will be marked as a failed sample. These attributes can be changed in the QC Summary filter, so they can be adjusted without needing to run another analysis should you want to change the selections. Lastly the Standard Peptides list is used to specify peptides and their charge in the format shown (e.g. LLDMGET, 2). These standard peptides will have their RT and m/z mapped for each sample that reported the peptide and charge. At this time, only peptides without modifications can be used (so adding in a modification to the sequence will not work).

Click on “Report” to proceed to the final step. Report filters can be set here which will filter the final result.
These filters are provided:

**PSM/Peptide:** this section is a mandatory section to filter out some results under the user-defined filter. Any changes to the filter will create a new analysis with old parameters and update the results and statistics.

**PSM -10lgP:** The PEAKS peptide score (-10lgP) is calculated for every peptide-spectrum match (PSM) reported by PEAKS DB, PEAKS PTM, and SPIDER. The score is derived from the p-value that indicates the statistical significance of the peptide-spectrum match. A peptide may be matched to many spectra, resulting in multiple PSMs. In that case, the peptide's score is calculated as the maximum among all PSM scores. For details of the scoring algorithm, please refer to the publication, "PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification ", Mol Cell Proteomics, 2011 Dec 20. A minimum requirement can be set and all identifications must pass this filter.

**PSM FDR (%):** The PSM filter can also be set to use the false discovery rate. FDR is estimated using a "decoy-fusion" approach.

**Peptide FDR (%):** The Peptides can also be set to use the false discovery rate. Peptides are scored based on their top PSM and using a "decoy-fusion" approach, the result is reported at that FDR.

**Note:** Decoy fusion is an enhanced target-decoy method for result validation with FDR. Decoy fusion appends a decoy sequence to each protein as the "negative control" for the search. See BSI's web tutorial (http://www.bioinfor.com/fdr-tutorial/) for more details.
**De Novo ALC (%):** This filter controls the minimum ALC score required for display in the de novo only result. A default of 50% is set.

**Denovo Only Tag Sharing:** The number set here corresponds to the length of de novo only tags to be shown in the result Protein coverage. For de novo only peptides that are not fully matched, they must share a tag length with at least minimum of this parameter with the protein residue to be displayed.

**Denovo Only Fully Matched:** Checking this option will show the de novo only peptides in the Protein coverage regardless of what tag sharing parameter is set if the entire de novo only sequence matches the protein residue.

In the Report step, there also is the option to specify an Analysis Name (which can be later renamed) and/or decide to save the Workflow to quickly re-use in future analyses.

### 6.2 Understanding PEAKS DB Search Results

After a PEAKS Search is complete, all searches selected during the analysis creation will generate result nodes in Project View. One of the result nodes is the de novo sequencing result, which includes de novo peptide sequences listed, spectrum annotation and other detailed information. The other result nodes are identification results from the database search tools: PEAKS DB, PEAKS PTM, and SPIDER. If more than one database search tool is enabled, the results from previous search tools are automatically merged.

Double-click on the result node to open their respective results. The analysis results for database search tools are presented in five tabs:

- **Summary:** This shows an outline of the PEAKS Studio database search results with key statistics. The overall quality of the experiment can be examined and the filters for peptide and protein identifications can be adjusted. This page shows after the search is done.
- **Proteins:** This shows a list of protein identifications. This view also visualizes protein sequence coverage and helps with protein characterization.
- **Peptide:** This shows a list of peptide identifications. This view also provides spectrum annotation and other detailed information for peptide precursor spectrum matches.
- **Modified Peptide:** This shows a list of peptide identifications with modifications and their respective unmodified form. A mirror plot is provided in the spectrum chart for further comparison.
- **De novo only:** This shows a list of quality peptide sequences detected by de novo sequencing that remain unidentified by the PEAKS Studio database search.
- **Feature:** This shows a list of features with and without peptide identifications.

The following sections will explore each of these six tabs in the Identification result.

#### 6.3 PEAKS DB Search result - Summary tab

The Summary view reports key statistics as an overview of the result. In the report, several statistical charts can be examined to assess result quality. At the top of each summary result page, protein filters can be set. Identifications that fall below the filters will not be visible in the result pages or exports.

**Proteins**

- **-10logP:** 25.19
- **FDR:** 1.0
- **% and ≥ 1:** unique peptides

**PTM Confidence**

- **Score:** 15.0
- **Ion Intensity:** 2.0
**Proteins -10lgP:** The PEAKS protein score (-10lgP) is calculated as the weighted sum of the -10lgP scores of the protein's supporting peptides. After removing any redundant peptides, the supporting peptides are sorted by -10lgP scores in descending order, and the k-th ranked peptide contributes to the weighted sum with a weight of 1/k. A default threshold of 15 is set.

**Proteins FDR:** The Proteins can also be filtered in accordance to the false-discovery rate of the protein set.

**Proteins Unique Peptides:** The minimum number of unique supporting peptides for a protein identification. A unique peptide is defined as a peptide that can be mapped to only one protein group.

**PTM Confidence Ascore and Ion Intensity:** These filters set the default Ascore and ion intensity filters in the Protein coverage pane. They are also used to determine which modifications are considered confident in the modified peptides table.

After changing the filters, click **Apply** to apply the new filter and update the results accordingly.

1. Notes
2. Result Statistics

![FDR curve](image1.png)  
**Figure 1.** False discovery rate (FDR) curve. X axis is the number of peptides being kept. Y axis is the corresponding FDR.

![PSM score distribution](image2.png)  
**Figure 2.** PSM score distribution. (a) Distribution of PEAKS peptide score. (b) Scatterplot of PEAKS peptide score versus precursor mass error.

Click on **Notes** to open the Notes dialog to save useful information regarding this Analysis. The Notes will appear in Section 1 of the summary page at the top.

The Summary view reports key statistics as an overview of the result. In the report, several statistical charts can be examined to assess result quality.
6.3.1 PEAKS DB Search result - Figure 1. False Discovery Rate (FDR) curve.

Figure 1 shows the FDR curve with respect to the number of PSMs being kept after filtration by the peptide -10lgP score. By lowering the score threshold, more PSMs are kept in the filtered result. Conversely, the FDR increases because more false positives are kept. In this figure, the vertical line indicates the current score threshold. The number of PSMs and the corresponding FDR at the current score threshold are shown in the top-left corner. PEAKS Search estimates FDR using the "decoy-fusion" approach. Decoy-fusion is an enhanced target-decoy approach that makes more conservative FDR estimations. For details of the "decoy-fusion" approach, please refer to the publication, "PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification", Mol Cell Proteomics, 2011 Dec 20.

It is commonly recommended to set FDR at <1% by adjusting the score threshold. If a rapid growth of the FDR around the 1% FDR threshold is noticed, some PSMs may be sacrificed to significantly reduce the FDR.

Note: When counting the number of PSMs, PEAKS Studio keeps one peptide per spectrum at most. Thus, the number of PSMs is actually the number of spectra with assigned peptides.

Note: During Analysis creation, if Peptide FDR (%) filter was used in Report step, the Summary page FDR curve in the results will show the Peptide FDR curve and corresponding number of Identified peptides.

Note: Decoy matches are not counted in the number of PSMs. Unless otherwise specified, decoy matches are also excluded from the other statistical numbers shown in the Summary view.

6.3.2 PEAKS DB Search result - Figure 2. PSM score distribution

Figures 2(a) and 2(b) in the Summary view show the peptide score distribution of PSMs. When FDR estimation is enabled in the PEAKS Search parameters, the target and decoy matches are respectively shown in two different colors.

Figure 2(a) is a stacked histogram showing the number of target and decoy matches in each score interval. There should be a similar number of target and decoy matches with low scores, but very few decoy matches with high scores. The vertical line indicates the current score threshold.

Figure 2(b) is a scatter plot showing the peptide score versus the precursor mass error in ppm for all PSMs. This figure is most useful for high resolution instruments. The PSMs with high scores should be centered near the mass error of 0, and the PSMs with low scores are usually scattered to larger mass error. The vertical dotted line indicates the current score threshold.
6.3.3 PEAKS DB Search result - Table 1. Statistics of data

Statistics of Data tabulates MS1 and MS2 information, as well as identification in the peptide and protein level. This table in the Summary view provides a condensed statistical information for the analysis.

<table>
<thead>
<tr>
<th>Sample</th>
<th>#MS1</th>
<th>#MS/MS</th>
<th>#Chimera</th>
<th>#Features</th>
<th>Identified #PSMs</th>
<th>Identified #Scans</th>
<th>Identified #Features</th>
<th>#Peptides</th>
<th>#Sequences</th>
<th>#Proteins*</th>
</tr>
</thead>
<tbody>
<tr>
<td>SMA_1</td>
<td>3906</td>
<td>19343</td>
<td>17093</td>
<td>64673</td>
<td>9711</td>
<td>9006</td>
<td>7417</td>
<td>6735</td>
<td>6251</td>
<td>891</td>
</tr>
<tr>
<td>SMA_2</td>
<td>3799</td>
<td>19317</td>
<td>16581</td>
<td>63508</td>
<td>9500</td>
<td>8847</td>
<td>7289</td>
<td>6614</td>
<td>6121</td>
<td>876</td>
</tr>
<tr>
<td>SMA_3</td>
<td>3915</td>
<td>19445</td>
<td>16358</td>
<td>63352</td>
<td>9621</td>
<td>8970</td>
<td>7310</td>
<td>6662</td>
<td>6161</td>
<td>877</td>
</tr>
<tr>
<td>WT_1</td>
<td>3742</td>
<td>19403</td>
<td>16850</td>
<td>68969</td>
<td>9225</td>
<td>8587</td>
<td>7185</td>
<td>6354</td>
<td>5946</td>
<td>898</td>
</tr>
<tr>
<td>WT_2</td>
<td>3950</td>
<td>19935</td>
<td>16746</td>
<td>67043</td>
<td>9392</td>
<td>8753</td>
<td>7204</td>
<td>6410</td>
<td>6005</td>
<td>923</td>
</tr>
<tr>
<td>WT_3</td>
<td>3792</td>
<td>19811</td>
<td>17522</td>
<td>70194</td>
<td>9396</td>
<td>8766</td>
<td>7265</td>
<td>6457</td>
<td>6055</td>
<td>933</td>
</tr>
</tbody>
</table>

* * proteins with significant peptides are used in counts.

** features are identified by DB search only.

#MS1: Total # of MS1 spectra for each sample. For timsTOF data this shows the # MS1 frames.

#MS/MS: Total # of MS2 spectra for each sample. This does not include chimeric scans. For timsTOF data this shows the # MS2 frames.

#Chimera: The number of chimeric tandem scans in the sample.

#Features: Total # of MS1 features detected from each sample.

Identified #PSMs: The number of peptide-spectrum matches.

Identified #Scans: The number of tandem scans that have a peptide-spectrum associated with it.

Identified #Features: The number of features that have a PSM.

#Peptides: The unique number of peptide sequences with modifications not including I/L differentiation.

#Sequences: The unique number of peptide sequences not including modifications and I/L differentiation.

#Proteins Groups: PEAKS Studio groups proteins identified by a common set of peptides. This number shows the number of protein groups in the filtered result based on All proteins.

#All Proteins: Total number of proteins with significant peptides, according to current protein filters.

#Top Proteins: Total number of top proteins with significant peptides, according to current protein filters.

6.3.4 PEAKS DB Search result - Figure 3. Sample overlap for Proteins and Peptides

Figures 3(a), 3(b), and 3(c) overlaps between samples in the search for all proteins, top proteins, and peptides, respectively.

Note: Top proteins are supported by the most number of unique peptides in the protein group.

Note: Peptide sequences with I/L differences will only be counted once.

Sample overlaps are displayed as venn diagrams for 2-4 samples.
6.3.5 PEAKS DB Search result - Figures 4 and 5. Distribution of peptide feature detection

Figures 4(a), 4(b), and 5(a) in the Summary View show the distribution of peptide feature detection.

Figure 4(a) in the Summary View provides all detectable features distributed over the m/z range. This stacked histogram shows the number of detectable features in each m/z interval. In addition, the number of detectable features with database results are highlighted in orange and the number of detectable features with de novo results in addition to the database results are highlighted in light blue. In a typical dataset, it is often observed that a higher number of detectable features occur in the lower m/z range, which is reflected within the identification results.
Figure 4(b) in the Summary View provides all detectable features distributed over the total retention time. This stacked histogram shows the number of detectable features in each RT interval. In addition, the number of detectable features with database results are highlighted in orange and the number of detectable features with database and de novo results are highlighted in light blue. In a typical dataset, it is often observed that a higher number of detectable features occur in the middle of the RT window, which aligns well with the identification results.

![Stacked histogram of detectable features](b)

Figure 5(a) in the "Summary" view provides all detectable features distributed over the feature abundance measured as the area under the XIC curve. This stacked histogram shows the number of detectable features measured with a certain total abundance. In addition, the number of detectable features with database results are highlighted in orange and the number of detectable features with de novo results in addition to the database results are highlighted in light blue. In a typical dataset, it is often observed that more peptide features will be assigned a database ID when they exhibit a stronger LC-MS signal.

**Figure 5.** Distribution of identified peptide features. (a) Feature abundance distribution;

![Stacked histogram of feature abundance](a)

6.3.6 PEAKS DB Search result - Result filtration parameters

Tables 2-4 show the statistical numbers of the data and results.
Table 2. Result filtration parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peptide -10logP</td>
<td>≥ 16.22</td>
</tr>
<tr>
<td>Protein -10logP</td>
<td>≥ 15</td>
</tr>
<tr>
<td>Proteins unique peptides</td>
<td>≥ 1</td>
</tr>
<tr>
<td>De novo score(%)</td>
<td>≥ 50%</td>
</tr>
</tbody>
</table>

Table 3. Statistics of filtered result.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDR (Peptide-Spectrum Matches)</td>
<td>1.0%</td>
</tr>
<tr>
<td>FDR (Peptide Sequences)</td>
<td>1.9%</td>
</tr>
<tr>
<td>FDR (Protein Group)</td>
<td>2.0%</td>
</tr>
<tr>
<td>De Novo Only Spectra</td>
<td>19875</td>
</tr>
</tbody>
</table>

Table 4. PTM profile.

<table>
<thead>
<tr>
<th>Name</th>
<th>ΔMass</th>
<th>Position</th>
<th># peptides</th>
<th># proteins</th>
<th># modification sites</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carboxymethylation</td>
<td>57.02</td>
<td>C</td>
<td>1901</td>
<td>1135</td>
<td>1765 (1765)</td>
</tr>
<tr>
<td>ADP-Ribose</td>
<td>541.06</td>
<td>CDKETSNR</td>
<td>1712</td>
<td>826</td>
<td>328 (1679)</td>
</tr>
<tr>
<td>Oxidation (M)</td>
<td>15.99</td>
<td>M</td>
<td>429</td>
<td>315</td>
<td>258 (403)</td>
</tr>
<tr>
<td>Acetylation (N-term)</td>
<td>42.01</td>
<td>N-term</td>
<td>332</td>
<td>211</td>
<td>134 (221)</td>
</tr>
</tbody>
</table>

Table 2. Result filtration parameters: Table 2 summarizes the result filtration parameters set at the top of the Summary View page. This information provides a helpful reference for exported data.

Table 3. Statistics of filtered results: Table 3 presents the statistics based on the data analysis results after setting the desired filters. See below for further explanation:

- **FDR (Peptide-Spectrum Matches):** The total number of decoy database assignments to spectra relative to the total number of target database assignments to spectra represented as a percentage.
- **FDR (Peptide Sequences):** The total number of decoy database assignments to unique peptide sequences relative to the total number of target database assignments to unique peptide sequences represented as a percentage.
- **FDR (Protein Group):** The total number of decoy protein groups where the top hit in the protein group is a decoy database protein relative to the total number of target database protein groups where the top hit is from the target database.
- **De Novo Only Spectra:** Corresponds to the number of existing de novo sequencing results without a positive protein identification, given the filters set for the search results.

Table 4. PTM Profile: shows modifications identified in the filtered result and the number of peptides, proteins, and protein modification sites containing each modification.

6.3.7 PEAKS DB Search result - Experiment Control

Figures 6(a) and 6(b) in the Summary View show the precursor mass errors of PSMs in filtered results. Figure 6(a) shows the distribution of the precursor mass errors in a histogram with and without software calibration. Figure 6(b) shows precursor m/z versus precursor mass error in a scatter plot after software calibration. The precursor mass error is calculated in ppm as \(10^6 \times \frac{\text{precursor mass} - \text{peptide mass}}{\text{peptide mass}}\). Calibration is for display purposes only and PEAKS corrects m/z value for each PSM to correct systematic biases and instrument measurement drift. Figures 5(a) and 5(b) help examine whether or not the mass spectrometer is properly calibrated. For a well-calibrated instrument, precursor mass errors should center at 0 ppm across the range of m/z.
Figure 6. Precursor mass error of peptide-spectrum matches (PSM) in filtered result. (a) Distribution of precursor mass error in ppm; (b) Scatterplot of precursor m/z versus precursor mass error in ppm.

Table 5 shows the number of identified peptides in each sample with the number of missed cleavages, indicating the enzyme digestion efficiency.

**Table 5. Number of identified peptides in each sample by the number of missed cleavages.**

<table>
<thead>
<tr>
<th>Missed Cleavages</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4+</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>132</td>
<td>56</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Sample 2</td>
<td>113</td>
<td>51</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Sample 3</td>
<td>133</td>
<td>60</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

6.3.8 PEAKS DB Search result - Other Information

Table 6 summarizes the parameters defined by the user to analyze the data. This information includes error mass tolerances, missed cleavages, database(s) used, etc.

4. Other Information

**Table 6. Search parameters.**

- Search Engine Name: PEAKS
- Parent Mass Error Tolerance: 10.0 ppm
- Fragment Mass Error Tolerance: 0.02 Da
- Precursor Mass Search Type: monoisotopic
- Enzyme: Specified by each sample
- Max Missed Cleavages: 1
- Digest Mode: Semi-Specific
- Peptide Length Range: 6 - 45
- Fixed Modifications:
  - Carbamidomethylation (+57.02)
- Variable Modifications:
  - Deamidation (NQ) (+0.98)
  - Oxidation (M) (+15.99)
- Max Variable PTM Per Peptide: 2
- Database: uniprot_sprot_june-2-2020.fasta
- Taxon: Mammalia
- Searched Entries: 67217
- Deep Learning Boost: False
- FDR Estimation: Enabled
6.4 PEAKS DB Search result - Proteins View

The Protein View lists protein identifications that have been filtered by the current settings in Summary View. It also visualizes the protein sequence coverage for identified proteins. The view has four main components:

- **Protein table**: This lists protein identifications that have been filtered by the current settings in the Summary View and the “Protein View Filters”.
- **Coverage**: This visualizes the protein sequence coverage of the selected protein, showing the mapping of support peptides and de novo tags to the protein sequence.
- **Peptides**: This lists supporting peptides mapped to the selected protein.
- **De novo Tags**: This lists "de novo only" peptides mapped to the selected protein.

### 6.4.1 Proteins View - Protein Table controls

At the top of the Proteins View, there are the options to filter, show/hide columns, search, and export the protein table.

### 6.4.1.1 Protein Table controls - Protein View Filters

Click on the **Protein View Filter** button to open the Protein View Filter.
The following options are available for filtering the protein table:

**Show Top, All, or First proteins in each group:** Adjust the protein list based on Protein grouping, top is selected by default. Proteins are grouped based on parsimony.

- **Top:** Show only the top proteins in each group. These proteins are supported by the most unique peptides in the group. Proteins in the group that share a subset of the unique peptides that support the top protein (subproteins) will not be displayed.
- **All:** Show all proteins in each group. Proteins are grouped together if they are supported by the same set or a sub-set of the top protein in the group. If a protein is supported by a peptide not supporting the top protein it will be added to a new group.

**Protein accession/name contains:** Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

**Protein description contains:** Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

**Note:** In these above two filters, multiple entries can be separated by a semi-colon.

**Protein sample area:** Filter proteins based on total protein area found in each sample. Sample areas are calculated using the total of all peptide features from unique supporting peptides. If a protein is identified and has areas from multiple samples, only one of the samples must pass this filter to be included.

**PTM contains:** Filter protein results by the presence of selected PTMs. Proteins that are supported by peptides containing modifications that match the selected PTMs will be included. If multiple PTMs match the modification name, they will all be included in the table. For SPIDER results, the mutations will also be listed in the PTM contains filter.

Click on the **Reset** button to remove all filters.

Click on **OK** to apply the Protein View Filter to the Protein table.

Click on **Cancel** to discard changes and close the dialog.
6.4.1.2 Protein Table controls - Protein count
At the top of the protein table shows the count of protein groups and number of proteins in the table based on the current Protein View Filters.

57 protein groups, total 142 proteins

6.4.1.3 Protein Table controls - Optional columns
Click on the button to see a list of sample-based optional columns.

Select the corresponding sample-based Coverage, Spectra count, or Area to add into the Protein Table view.

- **Coverage by Sample**: The protein coverage as a percentage will be displayed for the individual sample when selected.
- **#Spec by Sample**: The total number of spectra identified that support the given protein. Totals are given for each sample included in the search.
- **Area by Sample**: The total area of peptide features from unique supporting peptides in each sample will be displayed.

The exports will always include the contents of these optional columns even if they are hidden.

6.4.1.4 Protein Table controls - Protein Table Search Function
An easy-to-use search function is available for the protein table. The protein accession or description substring can be searched for, and all search results can be navigated through with the Up and Down arrows.

**Protein accession/name contains**: Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.
**Protein description contains:** Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

### 6.4.1.5 Protein Table controls - Protein Table Export

Click on the button to export the contents of the current table. This considers the Protein View Filters, the sorting applied to the table and is consistent with the view. There is the option to export the Protein table and/or the Supporting peptide table in CSV format.

### 6.4.2 Proteins View - Protein Table

Each row in the table represents a group of proteins that are supported by a common set of peptides. To expand the group, click the right arrow button at the left.

The columns in the Protein Table are:

- **Accession:** The accession number of the protein as seen in the FASTA database.
- **Cluster:** A unique identifier that classifies which protein group the protein belongs to.
- **Top:** Indicates whether this protein is a top protein.
- **-10lgP:** The protein confidence score.
- **Coverage (%):** The percentage of the protein sequence that is covered by supporting peptides. The coverage is visualized by a colored bar. Light blue blocks indicate the parts of the sequence covered by low-confidence peptides. Dark blue blocks indicate the parts covered by high-confidence peptides. This is the total coverage including results from all samples.
- **#Peptides:** The number of high-confidence supporting peptides.
- **#Unique:** The number of high-confidence supporting peptides that are mapped to only one protein group. Unique peptides with same sequence but different modifications are only counted once in this number.
- **PTM:** The identified modifications displayed with color-coded icons.
- **Avg. Mass:** The protein mass calculated using the average mass.
- **Description:** The protein's header information as seen in the FASTA database.
**Note:** For #Peptides and #Unique, two peptides with the same starting and ending positions in the protein are counted as one, regardless of their PTM forms. This is to follow the MCP (Molecular & Cellular Proteomics) guidelines.

### 6.4.3 Proteins View - Coverage Pane
The Protein Coverage view visually maps the supporting peptides and de novo tags to the protein selected in the Protein table. It also shows all identified sites with modifications or mutations to assist with protein characterization at the amino acid level. The "Protein Coverage" view is composed of three major components - Protein Sequence Coverage, Coverage Control Panel, and Protein Tools.

#### 6.4.3.1 Protein Sequence Coverage
This area visualizes the coverage of the protein sequence.

Regions in the protein sequence that are covered by supporting peptides are displayed in bold font with a grey background. Confident modifications and mutations identified in supporting peptides are displayed as icons above the protein sequence. Modifications are represented by colored icons with the first letter of its modification name. If a residue is modified by more than one modification in the same supporting peptide, "*" is used instead of a letter. Mutations are represented by framed white icons with a letter indicating the post-modification residue. Placing the cursor over a mutation icon shows the full name of the modification/mutation. The number above a mutation icon indicates the position in the protein sequence. The header information of the protein is shown on the top of the protein sequence.

The supporting peptides and matched de novo tags can be shown as colored bars under the protein sequence. Placing the cursor over a bar shows a tooltip with detailed information of the peptide. Left clicking on a bar shows the annotated spectrum from which the supporting peptide is identified.
Right clicking on a peptide bar shows additional options. “Show in spectrum” has the same behavior as left clicking on the peptide bar which is to open the annotated spectrum chart. “Show in table” has the function to jump to the supporting peptides table.

6.4.3.2 Coverage Control Pane
This panel controls what to display in the protein sequence coverage view.

**Sequence Display Mode:** The protein sequence can be displayed in either "outline" or "coverage" mode.

**Outline mode:** In outline mode, the protein sequence is displayed without supporting peptides and matched de novo tags. However, the sequence coverage at a specific position can be examined by left-clicking a residue.

```
1  MPEEYHGE  EVETFAGAE  IAJLMSLIIN  TFYSNKEIFL  RELINSADA  LDKIRYESLT  DPKLDGSKE  LKIIDIIPNQ
```

**Coverage mode:** In coverage mode, all the supporting peptides and matched de novo tags are shown.

```
1  MPEEYHGE  EVETFAGAE  IAJLMSLIIN  TFYSNKEIFL  RELINSADA  LDKIRYESLT  DPKLDGSKE  LKIIDIIPNQ
```

**AAs per line and 10AA gap:** Specify the number of amino acids per line and whether to show a gap for every 10 amino acids in the protein sequence.

**Sequence Display Option:** Select whether to view the protein sequence as Fasta or a specific enzyme used in the project. When a particular enzyme is selected, the protein sequence will be highlighted to emphasize expected peptides generated by the selected enzyme's cleavage specificity.
Modification minimal ion intensity / Ascore: A modification or mutation is considered confident if there are fragment ions supporting the modified residue with the minimal peak intensity above the specified threshold. Only confident modifications and mutations are displayed as icons above the protein sequence.

![Modificationminimalionintensity/Ascore](image)

Modification table: The modification table shows the modifications identified in supporting peptides on the protein. For each modification, the number of supporting PSMs with this particular modification is shown. The checkbox on the left controls whether to indicate the modification in the protein coverage.

<table>
<thead>
<tr>
<th>AA</th>
<th>Modification</th>
<th>PSM</th>
<th>#</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>Carbamidomethylation</td>
<td>03</td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>Deamidation (NQ)</td>
<td>07</td>
<td></td>
</tr>
<tr>
<td>O</td>
<td>Oxidation (Met)</td>
<td>05</td>
<td></td>
</tr>
<tr>
<td>K</td>
<td>2-amino-3-oxo-butanoic</td>
<td>03</td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>Formylation</td>
<td>03</td>
<td></td>
</tr>
<tr>
<td>L</td>
<td>Acetylation (Protein)</td>
<td>02</td>
<td></td>
</tr>
<tr>
<td>S</td>
<td>Sodium adduct</td>
<td>02</td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>Amidation</td>
<td>01</td>
<td></td>
</tr>
</tbody>
</table>

6.4.3.3 Proteins View - Protein Tools

The Proteins View toolbar is at the upper-right corner of the protein sequence coverage tab and contains helpful protein tools.

Full screen: When selected, the protein coverage window expands in full screen mode. Click to return from full screen mode.

Tool box: The tool box contains the following tools:

- Copy protein sequence. This copies the protein sequence of the current selected protein into the system clipboard.
- Save protein coverage as image. Different scale options are provided to increase the image size.
- Coverage statistics. This provides useful statistics based on the current selected protein.
PTM Profiling: Calculates the difference in abundance between peptides with confidently identified PTMs versus unmodified peptides. Peptide feature areas are used for this comparison.

For the Identification results (DB/PTM/SPIDER), the PTM Profiling tool provides quantitative information of modified peptides compared with unmodified peptides for the modification sites of the protein across all MS samples.

The inputs for the profiling can be set using the drop-down lists at the top-left corner of the dialog.

- **Proteins**: The accession ID of the selected protein to be analyzed for PTM profiling is shown in the title of the dialog. The selected protein cannot be changed within the dialog. To run profiling on another protein, close the dialog, select the protein of interest from the protein table and run the profiling by clicking the button.

- **Modification**: The drop-down list contains the detected and selected confident PTMs for the protein. Select the type of modification to study a specific PTM or choose “All the PTMs” to visualize the profiling information of all PTM sites at together.

  **Note**: The fixed PTMs are not used for profiling.

- **Peptides**: Data source used for the analysis. “All” considers all the peptides for this protein. When “Fully digested” is selected, only the full enzyme digested peptides will be used to calculate the modification abundance.

PTM Profile Table
The PTM Profile Table shows the detailed quantitative information for the modification sites. Each row in the table shows the modified and unmodified area for each sample of the selected PTM at each detected modification site. The table can be sorted by any column. The following list describes the contents in each column:

- **(Check box):** Allow the selection of each table row. Only the marked rows are used for the PTM Profile Chart.
- **Protein Position:** The detected modification sites in the protein. The modified amino acid symbol together with the position in the protein sequence is shown.
- **PTM:** The type of PTM being profiled.
- **-10logP:** The best -10logP identification score for the corresponding modified peptides.
- **AScore:** The best PTM AScore associated with the corresponding PTM at the specified site of the modified peptides.
- **Ion intensity:** The best PTM ion intensity associated with the corresponding PTM at the specified site of the modified peptides.
- **Sample Modified:** Total area of the features associated with the corresponding modified peptides for each sample.
- **Sample Unmodified:** Total area of the features associated with the corresponding unmodified peptides for each sample.

When “Fully digested” is selected, there are two more columns shown in the PTM Profile Table:

- **Peptide:** The fully digested peptide sequence.
- **Peptide Position:** Position of the modified amino acid in the peptide sequence.

### PTM Profile Chart

The PTM Profile Chart is located at the bottom left corner in the dialog. It gives a visualization of the ratio between unmodified and modified amino peptides at each detected modification sites. The x-axis indicates the percentage (%) of modified or unmodified peptide feature areas relative to the total abundance whereas the y-axis indicates the position in the protein sequence where a modification was identified for each individual sample.

**Note:** Since the percentage is calculated over the sum of the modified and unmodified areas, the modified and unmodified portion always sum up to 100%.

**Note:** For samples with the specific PTM not detected, the corresponding bars are left as blank.
Spectrum Annotation View

The Spectrum Annotation View provides a graphical representation for the best modified peptide and the best unmodified peptide when “All” is selected from the “Peptides” drop-down list. The top spectrum corresponds to the best modified peptide, whereas the bottom spectrum shows the best unmodified peptide. When “Fully digested” is considered, the best modified peptide and the best unmodified peptide are displayed in a compare view for quick visualization and validation.
PTM Profile Export

The PTM profile can be exported as an image as well as in text format.

**Image export:** To save the profile chart, right click and choose Save to Image... menu item to save the displayed graph as a PNG file.

**Text format export:** Click the button at the top-right corner to export the calculated PTM Profile data to a text file in CSV format. The PTM Profile of the selected modification on the selected protein will be exported by default. In addition, profiles on all proteins for the selected modification may also be exported, if “Export ptm profile for all proteins” is checked. The exported file contains all necessary information to construct the PTM profile graph. The PTM profile data are grouped together by protein, then by modification type, and then by the modification site on the protein sequence. The headers of the CSV column are similar to the PTM Profile Table and also self-explanatory. The following items are explained for clarity:
• **Modified Peptide Sequence Window**: protein sequence containing 10 AA ahead of the modification site and 10 AA after the modification site with padding as necessary.

• **Sample Best AScore**: best AScore for each sample.

**Note**: When “Export PTM profile for all proteins” is not checked, only the profiling data that is marked in the PTM Profile Table will be exported.

**Peptide Mapping**

A peptide map is often referred to as a “fingerprint” of a protein. It is a powerful tool that provides a comprehensive understanding of the protein being analyzed. It gives a detailed picture of the ion signal of all peptides associated with the protein. In addition to generating the peptide map, PEAKS has integrated an annotated tool-tip which allows for easy identification of the peptide shown at any point on the map. If multiple data files are searched together in the analysis, then multiple peptide maps may be compared to one another. This allows users to easily determine the consistency of the protein between samples.

PEAKS provides two different approaches to analyze peptide mapping of a protein.
Annotated Chromatogram: The annotated chromatogram of the selected protein is displayed in profile mode. The intensity of each peptide feature associated to the protein is mapped to its retention time (RT). The annotated chromatogram has its own input selection, result display, and result exporting options.

- **Fraction Choices:** The data source used for peptide mapping. When the search result involves multiple fractions, a comparison of annotated chromatograms for the selected protein among (a maximum of three) fractions, can be done by selecting the second and third fraction from the corresponding drop down menu.
- **Minimal Ion Intensity:** A filter used to set an intensity threshold, where peptide features below this value will not be included in the display.
- **Type:** Configures the display to view the Total Ion Chromatogram (TIC) or Base Peak Intensity (BPI).
- **Display Mode:** PEAKS offers two modes to view multiple chromatograms for better comparison.
  - **Compare:** In compare mode, the protein chromatograms for the specified data samples will appear on opposite sides of the x-axis. This allows the user to directly compare the protein chromatograms between two samples.
  - **Side by Side:** In Side-by-Side mode, the chromatograms are displayed individually; each with their own set of x- and y-axes. This display mode allows for the comparison of up to three samples.
- **PEAKS DB:** Peptide features associated with peptides identified by database searching are highlighted in the annotated chromatogram.
- **De novo only:** This highlights the peptide features associated with de novo only peptides.
- **Annotation:** When selected, the retention time of major peptide features are displayed on the peptide map.
- **Tool Tip:** When selected, a pop-up will appear when the cursor hovers over any peptide within the peptide map. The pop-up contains the retention time, the intensity (TIC or base peak), and the peptide sequences found at that particular retention time.

**Annotated Chromatogram Chart controls:** This chart gives a visualization of the peptide map for the selected data. It provides a few convenient ways to zoom-in and navigate the chromatogram.
Zoom to an RT region - Click the desired starting retention time, drag horizontally to the desired final retention time, and then release the mouse button.

Zoom in/out smoothly - Place the cursor at a particular RT value (right below the x-axis line), scroll the mouse wheel button.

Increase/decrease the ion intensity - Place the cursor on the chromatogram.

See the whole chromatogram - Double-click on the chromatogram.

Pan to an RT region: Place the cursor on the chromatogram, right-click, and then hold and drag to pan to an RT region.

- Result Export: The annotated chromatogram chart can be exported as an image in PNG format. Click the button to save the displayed chart.

**Peptide Comparison:** Another option PEAKS offers its users to analyze the peptide map is in terms of a peptide comparison view. In this view, the total area of the features associated with a peptide is mapped to the mass of the peptide. The peptide comparison has its own input selection, result display, and result exporting options.
**Inputs:** The inputs for the peptide comparison can be set using the drop-down menus at the top-left corner of the dialogue.

- **Fraction Choices:** The data source used for peptide mapping. Peptide comparison allows a maximum of two fractions as the source. After making choices for the fraction selection, click the Apply button to update the chart.
- **Minimal Feature Abundance:** A filter used to set a feature area threshold, where peptide features below this value will not be included in the display.
- **Annotation:** When selected, the masses of the major peptides are displayed on the comparison chart.
- **Tool Tip:** When selected, a pop-up will appear when the cursor hovers over any peptide within the comparison chart. The pop-up contains the peptide mass, the total area for that peptide, and the peptide sequence.

- **Peptide Comparison Chart:** This chart gives a visualization of the peptide comparison for the selected data. The chart is displayed in Peak Mode, where each peptide is represented as a bar and the height of that bar represents the total area of the associated peptide features in the specified fraction. The chart provides similar convenient ways to zoom-in and navigate the comparison chart.

- **Result Export:** The peptide comparison of a protein can be exported as an image, as well as in text format.
  - **Image export:** Click the **Save to Image** button to save the displayed graph as a PNG file.
  - **Text format export:** Click the **Save to Text Format** button to export the peptide comparison data to a text file in CSV format. The exported file contains all of the necessary information to construct the peptide comparison chart. The data are grouped together by peptide sequence and mass. The information for the same peptide are combined together if they belong to the same group, as mentioned earlier. The headers of the CSV column are self-explanatory. The following items are explained for clarity:
    - **Unique:** This specifies that the peptide is unique (Y) or not unique (N).
    - **-10lgP or ALC:** The significance score of the peptide identified by database searching or by de novo sequencing for de novo only peptides.
    - **Mass:** The mass of the peptide.
    - **ppm:** The error tolerance of the peptide.
    - **Feature Abundance:** This list of areas of the features associated with the corresponding peptide. The format of the feature areas is (area [space] +charge [semi-colon])* For example, 500 +2; 800 +2; 700 +3
    - **Total Abundance:** The sum of the areas of the features associated with the corresponding peptide in the specified fraction.
    - **Ratio:** When the data sources contain two fractions, the ratio of the total areas of the corresponding peptide between two fractions is calculated.

**Note:** -10lgP, ALC(%), and ppm values are from the best PSM of the corresponding peptide.
6.4.3.4 Proteins View - Supporting Peptides

The "Peptides" tab displays a table of supporting peptides for the protein of interest. This table is similar to the peptide table in the Peptide view, except that "Accession" is excluded and the columns "(Checkbox)", "Unique", "Start", and "End" are included. Descriptions of extra columns are given in the following sections.

- **(Checkbox):** This controls whether a peptide is shown in the protein coverage as a blue bar.
- **Unique:** This shows whether the peptide is a unique supporting peptide to the protein group.
- **Start:** This shows the peptide's starting position in the protein.
- **End:** This shows the peptide's ending position (inclusive) in the protein.

The supporting peptides with peptide -10lgP score below the peptide filtering threshold are listed in the table as well. By default, they are shown in grey and are not displayed in the protein coverage.

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6.4.3.5 Proteins View - De novo Tags

The "Denovo Only Tags" tab shows a table of "de novo only" peptides with sequence tags matched to the protein.

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6.4.3.5 Proteins View - De novo Tags

The "Denovo Only Tags" tab shows a table of "de novo only" peptides with sequence tags matched to the protein.
6.5 PEAKS DB Search result - Peptide View

The "Peptide View" displays the resulting peptide identifications that have been filtered by the current settings in Summary View and "Peptide View Filters". The table on the top shows the details of each peptide identification. The bottom section provides additional information about the peptide-spectrum matches for the native peptide selected in the table.

6.5.1 Peptide Table

The Peptide Table shows the filtered native peptide identification results. Each row in the table is a native peptide identification represented by its highest-scoring PSM. When there are more than 1000 peptides, the table is split into multiple pages.

Note: If there are multiple PSMs matched to a peptide, the table displays the information for the top-scoring PSM. Peptides with the same sequence but different modifications are considered different and shown in different rows.

To copy the sequence, right click on a row (or multiple rows) in the table and select "Copy selected peptide sequence(s)". Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

The following list describes the contents in each of the default columns:
- **Peptide**: The amino acid sequence of the peptide, as determined by the PEAKS search workflow. A modified residue is followed by a pair of parentheses enclosing the modification.

- **-10lgP**: The peptide -10lgP score. The score indicates the scoring significance of a peptide-precursor spectrum match.

- **Mass**: The monoisotopic mass of the peptide.

- **Length**: The number of amino acids in the sequence backbone.

- **ppm**: The precursor mass error, calculated as \(10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}\).

- **m/z**: The precursor mass-to-charge ratio.

- **RT**: The retention time (elution time) of the spectrum as recorded in the data.

- **Area per Sample**: The combined area for all features in that sample associated with this peptide.

- **Scan**: The scan number of the spectrum that matches the peptide sequence with the highest -10lgP.

- **#Feature**: For single sample analysis, shows number of features assigned to the peptide sequence from identification. For multiple samples, if peptide has feature vectors, it will show number of features used (area > 0) in all feature vectors. If not, it shows the number of features from identification.

- **Accession**: The accessions of proteins that contain this peptide.

- **PTM**: The types and the numbers of modifications present in the peptide shown in color-coded icons.

- **AScore**: Localization score assigned to modifications on the peptide. It is the -10 log of a p-value. In this case, the p-value is the probability that the modification occurs at the reported position compared to other possible positions. A -10lgP of 20 is equal to a p-value of 0.01. Scores are listed in the format AminoAcid, peptide position, modification, Ascore. For example, serine at amino acid 3 is phosphorylated with an Ascore of 54.01 will be reported as: S3 P 54.01.

- **Found By**: The name of the PEAKS search workflow that identifies the peptide; this can be either PEAKS DB, PEAKS PTM, or SPIDER.

### 6.5.2 Peptide Table controls
The menu above the peptide table provides options to scroll through the table of peptides, filter the table, show optional columns and search for specific entries.

### 6.5.2.1 Peptide Table controls - Pagination
For easy viewing and to improve performance, the peptide table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.
In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

6.5.2.2 Peptide Table controls - View Filter

Click on the button to open the Peptide View Filter which is used to filter the Peptide table. The following are the filter options provided:

- **Peptide sequence contains**: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains for further filtering if necessary.

- **Peptide score (-10lgP)**: Filters out all peptides with -10lgP scores below this threshold.

- **Peptide length**: Filters for peptides of length within this min/max threshold.

- **Peptide sample area**: Filters based on total peptide area found in each sample. Sample areas are calculated using the total of all peptide features from all spectra that identified the native peptide. If the peptide is found in multiple samples, only one sample must pass this filter for the peptide to be included.

- **PTM contains**: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text they will all be included in the table.

  **Note**: In all filters, multiple entries can be separated by a semi-colon.

Click on the Reset button to reset all filters to their default setting.

Click on OK to apply the filters to the Peptide table.

Click on Cancel to discard the changes in the View Filter.

6.5.2.3 Peptide Table controls - Optional Columns

Click on the button to open the optional columns dropdown. By default the sample area columns are displayed. Sample Feature count columns can also be shown. These sample feature counts are always from identification, and does not consider whether there are feature vectors present. For projects with many samples, it might be useful to hide these optional columns and reduce the width of the peptide table.
6.5.2.4 Peptide Table controls - Search function

On the top right of menu, there is a search function with 4 options. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- **Scan =**: Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction.

- **m/z ≈**: Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit. For example, searching for m/z ≈ 560.21 will find all m/z between [560.20, 560.22]

- **RT ≈**: Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/- 1 of the last significant digit set. For example, searching for RT ≈ 55.5 will find all m/z between [55.4, 55.6]

- **Seq contains**: Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered.

Additionally, since the peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVIK” is the same as searching for “LVLK”, “IVLK”, and “IVIK”

6.5.3 Feature selection breadcrumb

There is a sample and feature selection breadcrumb which allows you to switch between features associated with the peptide. Click on the Sample in the breadcrumb to reveal other samples and the features in that sample related to the current peptide selection. From here you can select other samples, fractions, or features.

By clicking on the fraction level, all fractions in this sample are revealed as well as the features in each fraction associated with this peptide. From here you can select other fractions and features to view.

Lastly, clicking on the feature level will show all features in the currently selected fraction. Features with Feature Id = 0 are known as fake features and are constructed to enhance identification.
6.5.4 PSM selection
Underneath the feature breadcrumb will shown the currently selected PSM. Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.

All other components in the Peptide tab interface will update according to the currently selected PSM.

For timsTOF instruments, Precursor ID is used rather than Scan number.

6.5.4.1 Protein jump button
Click on to jump to a protein that contains the currently selected PSM. If there are more than 1 protein, a dropdown is revealed which allows the user to select which protein to jump to. If a protein is filtered out due to a protein summary filter or summary page protein filter, then the option to jump will be grayed out.

6.5.4.2 Show spectrum in Data View
Click on to open the Data Refine MS/MS tab (if not already opened) and jump to the corresponding selected Scan.

6.5.4.3 Show spectrum in LC/MS View
Click on to open the Data Refine LC/MS tab (if not already opened) and jump to the corresponding selected Scan, highlighting the peptide MS/MS marker and the feature if present.

6.5.4.4 Show Raw Spectrum View
Click on to show the raw spectrum chart of the currently selected Scan.

6.5.5 Annotated Spectrum Chart
For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

6.5.5.1 Annotated Spectrum Chart - Chart Navigation
Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.
Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.
Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.
See the whole spectrum: Double click in the spectrum or click the 1:1 button.

6.5.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings
Click on the button to show the spectrum annotation settings.

Click on the button to show the spectrum annotation settings.
The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places:** Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation:** Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated:** Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation:** Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info:** Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity:** Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

### 6.5.5.3 Annotated Spectrum Chart - Additional Chart Controls

Click on ☐ or ☑ to toggle between absolute and percentage intensity values in the Y axis.

Click on ☐ to reset the zoom level of the annotated spectrum chart to the default.

Click on ☐ to zoom into the X axis by a factor of 2.

Click on ☐ to zoom into the Y axis by a factor of 2.
The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

### 6.5.6 Ion Match Table

The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings".

<table>
<thead>
<tr>
<th>Ion Match</th>
<th>Survey</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td>b</td>
</tr>
<tr>
<td>10</td>
<td>1076.503</td>
</tr>
<tr>
<td>12</td>
<td>1344.626</td>
</tr>
<tr>
<td>13</td>
<td>1481.659</td>
</tr>
<tr>
<td>14</td>
<td>1596.682</td>
</tr>
<tr>
<td>15</td>
<td>1453.703</td>
</tr>
<tr>
<td>16</td>
<td>1781.733</td>
</tr>
<tr>
<td>17</td>
<td>1878.815</td>
</tr>
<tr>
<td>19</td>
<td>2122.880</td>
</tr>
</tbody>
</table>

Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

### 6.5.7 Survey Chart

The “Survey” tab displays the precursor ion spectrum. The buttons that appear in this section are identical to those explained above in Section 6.5.5.3 Annotated Spectrum Chart - Additional Chart Controls.

### 6.5.8 LC/MS Snapshot

The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The peptide feature is highlighted in purple and the selected PSM marker is blue. The span of the orange line in the LC-MS Snapshot designates the isolation window.
Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a $m/z$ $RT$ (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

6.5.9 Precursor Profile
The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with the retention time along the x-axis and the intensity along the y-axis.
6.6 PEAKS DB Search result - Modified Peptide View

The Modified Peptide View introduced in updated PEAKS Studio 11 is similar to the Peptide View except the modifications are colour-coded based on whether they are considered confident or unconfident modifications. Green indicates a confident modification while red indicates an unconfident modification. The criteria for determining confident modifications can be adjusted in the Summary page by changing the PTM Confidence Ascore and ion intensity filters. This view also pairs the modified and unmodified sequence together for easy viewing of whether there is an unmodified form of the peptide reported. Only those peptides that exist with sequence without the modification of interest will be displayed underneath the modified peptide for comparison purposes. All searching and filtering function will be done on modified peptides only, and those unmodified peptides will be filtered out alongside the modified peptide search criteria.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>-10lgPw</th>
<th>Mass</th>
<th>Length</th>
<th>ppm</th>
<th>m/z</th>
<th>z</th>
<th>RT</th>
<th>T/kb</th>
<th>Area</th>
<th>Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>SV(+2.01)TVIVNDTGSHEDMIMDAQMDYVGTR</td>
<td>108.54</td>
<td>3038.3386</td>
<td>26</td>
<td>3.9</td>
<td>1015.0701</td>
<td>3</td>
<td>73.30</td>
<td>1.1189-1.1411</td>
<td>1.92E4</td>
</tr>
<tr>
<td>2</td>
<td>MC(-4.61)EIVDK</td>
<td>140.96</td>
<td>1848.2522</td>
<td>26</td>
<td>1.0</td>
<td>889.5792</td>
<td>3</td>
<td>80.52</td>
<td>1.0592-1.0914</td>
<td>4.75E3</td>
</tr>
<tr>
<td>3</td>
<td>MC(-2.01)TVIVNDTGSHEDMIMDAQMDYVGTR</td>
<td>108.54</td>
<td>3038.3386</td>
<td>26</td>
<td>3.9</td>
<td>1015.0701</td>
<td>3</td>
<td>73.30</td>
<td>1.1189-1.1411</td>
<td>1.92E4</td>
</tr>
<tr>
<td>4</td>
<td>MC(-4.61)EIVDK</td>
<td>140.96</td>
<td>1848.2522</td>
<td>26</td>
<td>1.0</td>
<td>889.5792</td>
<td>3</td>
<td>80.52</td>
<td>1.0592-1.0914</td>
<td>4.75E3</td>
</tr>
<tr>
<td>5</td>
<td>MC(-4.61)EIVDK</td>
<td>140.96</td>
<td>1848.2522</td>
<td>26</td>
<td>1.0</td>
<td>889.5792</td>
<td>3</td>
<td>80.52</td>
<td>1.0592-1.0914</td>
<td>4.75E3</td>
</tr>
<tr>
<td>6</td>
<td>MC(-4.61)EIVDK</td>
<td>140.96</td>
<td>1848.2522</td>
<td>26</td>
<td>1.0</td>
<td>889.5792</td>
<td>3</td>
<td>80.52</td>
<td>1.0592-1.0914</td>
<td>4.75E3</td>
</tr>
<tr>
<td>7</td>
<td>MC(-4.61)EIVDK</td>
<td>140.96</td>
<td>1848.2522</td>
<td>26</td>
<td>1.0</td>
<td>889.5792</td>
<td>3</td>
<td>80.52</td>
<td>1.0592-1.0914</td>
<td>4.75E3</td>
</tr>
<tr>
<td>8</td>
<td>MC(-4.61)EIVDK</td>
<td>140.96</td>
<td>1848.2522</td>
<td>26</td>
<td>1.0</td>
<td>889.5792</td>
<td>3</td>
<td>80.52</td>
<td>1.0592-1.0914</td>
<td>4.75E3</td>
</tr>
<tr>
<td>9</td>
<td>HGTPGSLV (+15.99)DEAGTPGNSQQFFIC(-37.02)TAK</td>
<td>150.89</td>
<td>2065.3167</td>
<td>27</td>
<td>2.4</td>
<td>1157.9089</td>
<td>3</td>
<td>85.92</td>
<td>1.3796-1.3978</td>
<td>5.07E3</td>
</tr>
<tr>
<td>10</td>
<td>HGTPGSLV (+15.99)DEAGTPGNSQQFFIC(-37.02)TAK</td>
<td>150.89</td>
<td>2065.3167</td>
<td>27</td>
<td>2.4</td>
<td>1157.9089</td>
<td>3</td>
<td>85.92</td>
<td>1.3796-1.3978</td>
<td>5.07E3</td>
</tr>
<tr>
<td>11</td>
<td>HGTPGSLV (+15.99)DEAGTPGNSQQFFIC(-37.02)TAK</td>
<td>150.89</td>
<td>2065.3167</td>
<td>27</td>
<td>2.4</td>
<td>1157.9089</td>
<td>3</td>
<td>85.92</td>
<td>1.3796-1.3978</td>
<td>5.07E3</td>
</tr>
</tbody>
</table>

Click on to switch to a mirror plot view comparing the modified peptide and unmodified peptide.

6.7 PEAKS DB Search result - De novo only View

The "De novo only View" displays the filtered de novo only sequences. A de novo sequence is de novo only if it is from an MS/MS spectrum that is not confidently assigned to any database peptide. The table shows the filtered de novo only sequences by the current settings in the Summary View. De novo only sequences may suggest novel peptides, peptides with unknown modifications, or other interesting research subjects. The interface is identical to the Peptide View in a de novo sequencing result node. For detailed instructions, please refer to Section 4.4 De novo result - De novo View.
6.8 PEAKS DB Search result - Feature View

The "Feature View" displays all the feature information in greater detail, as shown in the next figure. The table on the top displays features, while the bottom section provides additional information about the feature, precursor profile, and if available peptide-spectrum match. For detailed instructions, please refer to Section 4.4 De novo result - De novo View.
6.9 PEAKS DB Search result - Exporting

In Project View, double-click on ![Export](image) to open the Export node.
There are several options for exporting the PEAKS Identification results in CSV format:

- **Proteins CSV**: The list of protein identifications, filtered by the protein filters in the Summary view and will be saved to proteins.csv.
- **Peptide CSV**: All of the identified peptides and details will be saved to peptides.csv.
- **Support peptides CSV**: A list of supporting peptides of each protein identification, filtered by the protein filters in the "Summary" view and will be exported to protein-peptides.csv. This usually contains more entries than the Peptide table as a peptide can be identified in several proteins and, therefore, can be reported multiple times in this file.
- **PSM CSV**: The peptide-spectrum matches (PSMs) will be exported to DB search psms.csv. Peptides differentiated with only I/L isoform are represented by separate entries. As a result, the number of entries in this file might be bigger than the number of PSMs mentioned in the Summary view.
- **De novo only CSV**: Exports the De novo only table. This export does not consider any sorting and will export a separate file for each sample.
- **DB Features CSV**: This export does not consider any Feature view filters or sorting. Each sample will be in a separate export.
6.9.2 PEAKS DB Search result - Exporting - HTML format

Export in HTML: This will generate single or multiple HTML reports in the specified location.

The following exporting options are available:

- **Summary view:** The “Summary” view page will be saved as a summary.html file in HTML format in the specified location.
- **Protein coverage:** The coverage pane will be saved for each protein.
- **Supporting peptides:** A list of supporting peptides will be saved for each protein.
- **Best unique PSM:** The best unique PSM will be saved for each protein.
- **Export peptide list in html:** creates a separate HTML-containing peptide identifications.

An individual protein will have its own HTML output file, where the corresponding protein coverage, the supporting peptides, and the best unique PSM are gathered.

6.9.3 PEAKS DB Search result - Exporting - pepXML and mzIdentML

**Peptides - pepXML:** A list of peptide-spectrum matches will be saved to pep.xml in pepXML format. The number of the “spectrum_query” elements will be the same as the number of PSMs mentioned in the "Summary" view, and the number of the “search_hit” elements will be the same as the number of PSMs counted from Peptide table (summing up #Spec column). Current version of pepXML export is 1.22.

**Peptides - mzidentml:** A list of peptide-spectrum matches will be saved to mzidentml.xml in mzIdentML format version 1.2.0. To validate the export, search for the following keywords and compare with the results in PEAKS Studio:

```xml
<PepXMLExport>
  <Peptide>
    <PeptideSequence>Equal to the number of peptides in the Peptide table</PeptideSequence>
    <DBSearch accession>Equal to the total protein count in the Protein table</DBSearch accession>
  </Peptide>
</PepXMLExport>
```

**</ProteinAmbiguityGroup>**: Equal to the number of protein groups
“Leading Protein:” Equal to the number of top proteins, disregarding protein filters

“Non-leading Protein:” Equal to the number of non-top proteins, disregarding protein filters

“protein group passes threshold” value = “true”: Equal to the number of protein groups based on current protein filters

passThreshold=”true”>: Equal to the number of proteins based on current protein filters

6.9.4 PEAKS DB Search result - Exporting - Spectral Library generation

Spectral Library TSV: PEAKS identification results can be exported as a spectral library that can be used by the Library Search engine. Check “Include De Novo only” and select an ALC threshold to include de novo only peptides in the library. Select “Use original RT” to keep original RT values in the library. Select “Map to iRT” to save iRT values in the library. The map to iRT function maps experimental retention times for peptides to an indexed RT (iRT) to account for chromatographic differences between runs (e.g. different gradient lengths).

For more information on how to configure a PEAKS generated spectral library, refer to Section 2.10 Spectral Library Configuration

6.10 PEAKS PTM Finder

PEAKS PTM Finder is an optional dedicated search tool for unspecified modification search performed after Database Search. Open the PEAKS PTM result by double-clicking on the PEAKS PTM node in the Project View.

PEAKS PTM uses the entire list of common and uncommon built-in PTMs in PEAKS Studio unless specified by the user (see section 6.1 Database Search Workflow and Parameter Settings)


In the Protein table it will be common to see Proteins with many PTMs associated with their supporting peptides.

<table>
<thead>
<tr>
<th>PTM</th>
<th>Avg. Mass</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>273427</td>
<td>Fatty acid synthase OS= Homo sapien</td>
</tr>
<tr>
<td>O</td>
<td>164939</td>
<td>Carbamoyl-phosphate synthase amm</td>
</tr>
<tr>
<td>T</td>
<td>83281</td>
<td>Heat shock protein HSP90 beta OS=</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PTM</th>
<th>Avg. Mass</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td>Dehydration (-18.01)</td>
</tr>
<tr>
<td>D</td>
<td></td>
<td>Deamidation (NQ) (+0.98)</td>
</tr>
<tr>
<td>C</td>
<td></td>
<td>Carbamidomethylation (+57.02)</td>
</tr>
<tr>
<td>C</td>
<td></td>
<td>Carboxymethyl (+56.01)</td>
</tr>
<tr>
<td>O</td>
<td></td>
<td>Oxidation (M) (+15.99)</td>
</tr>
<tr>
<td>P</td>
<td></td>
<td>Pyro-glu from E (-18.01)</td>
</tr>
<tr>
<td>R</td>
<td></td>
<td>Replacement of 2 protons by calcium (+37.95)</td>
</tr>
<tr>
<td>D</td>
<td></td>
<td>Dimethylation(KR) (+28.03)</td>
</tr>
<tr>
<td>P</td>
<td></td>
<td>Phosphorylation (STY) (+79.97)</td>
</tr>
<tr>
<td>M</td>
<td></td>
<td>Methyltion(others) (+14.02)</td>
</tr>
<tr>
<td>a</td>
<td></td>
<td>2-amino-3-oxo-butanolic acid (-2.02)</td>
</tr>
<tr>
<td>C</td>
<td></td>
<td>Carboxylation (E) (+43.99)</td>
</tr>
</tbody>
</table>
### 6.10.1 PEAKS PTM Finder - Signature Ions

In updated PEAKS Studio 11, PEAKS PTM was enhanced to include Signature Ion detection, displaying expected and potential signature ions. In the Summary page, there is an additional table and chart which provides statistics on how many potential or expected signature ions were found in the result.

#### Table 5. Detected Signature Ions.

<table>
<thead>
<tr>
<th>Modification</th>
<th>Signature Ion Name</th>
<th>Mass</th>
<th>#PSM with modification</th>
<th>#PSM with modification with signature ion</th>
<th>#PSM without modification</th>
<th>#PSM without modification with signature ion</th>
<th>Specificity (Expected/Potential)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADP-Ribose</td>
<td>Adenine</td>
<td>136.0600</td>
<td>5025</td>
<td></td>
<td>49744</td>
<td></td>
<td>Expected</td>
</tr>
<tr>
<td></td>
<td>Adenosine-H2O</td>
<td>250.0900</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>AMP</td>
<td>348.0500</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ADP</td>
<td>428.0200</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ResidueLossS</td>
<td>524.0500</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ResidueLossK</td>
<td>542.0600</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oxidation (HW)</td>
<td>Oxi_130.06</td>
<td>130.0600</td>
<td>373</td>
<td>247</td>
<td>54396</td>
<td>13605</td>
<td>Potential</td>
</tr>
<tr>
<td></td>
<td>Dih_146.06</td>
<td>146.0600</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Dih_174.06</td>
<td>174.0600</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Dih_202.06</td>
<td>202.0600</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The detected signature ions are plotted in Figure 6, up to a maximum of 8 charts.

**Figure 6. Frequency Plot of Detected Signature Ions.**

![Signature Ion](image)

Expected Signature Ions are shown with green markers in the annotated spectrum chart.
All user interface related functions as described in the previous sections in this Chapter 6 are the same for the PEAKS PTM result. Additional options are added to view filters for the convenience of filtering for specific potential or expected signature ions.

### 6.11 SPIDER

SPIDER is an optional dedicated search tool for homology search after Database Search.


Open the SPIDER result by double-clicking on the node in the Project View.

In the Protein table, supporting peptides with mutations are shown as white coloured blocks.

All user interface related functions as described in the previous sections in this Chapter 6 are the same for the SPIDER result.

### 6.12 Identification Quality Control (QC)

The QC function was added in PEAKS Studio 11 upgrade. In the Identification workflow, it can be included as an optional step to produce an automated QC result node for sophisticated QC analysis on the data level and protein/peptide identification. This function has an additional use in analyzing protein/peptide quantification when enabled in the Label free quantification workflow. The function is used in both DDA and DIA data.
Double-click on the QC result node in the analysis to open the QC result node.

### QC Summary

**Table 1. Statistics of filtered results.**

<table>
<thead>
<tr>
<th>Selection</th>
<th>Count</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selected Samples</td>
<td>5</td>
</tr>
<tr>
<td>Passed Samples Count</td>
<td>2</td>
</tr>
<tr>
<td>Ratio of Passed Samples</td>
<td>40%</td>
</tr>
<tr>
<td>Failed Samples Count</td>
<td>3</td>
</tr>
<tr>
<td>Ratio of Failed Samples</td>
<td>0%</td>
</tr>
<tr>
<td>Total Failed Attributes</td>
<td>5</td>
</tr>
</tbody>
</table>

**Figure 1. Failed Attributes Chart.**

**Table 2. Statistics of I.FQ Result**

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>MS1</th>
<th>MS2</th>
<th>PSM</th>
<th>Identified Features</th>
<th>Quantified Features</th>
<th>Peptides</th>
<th>Quantified Peptides</th>
<th>Protein Groups</th>
<th>PSM</th>
<th>Peptides</th>
<th>Protein Groups</th>
<th>ID Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control Sample</td>
<td>4848</td>
<td>1669</td>
<td>16444</td>
<td>43989</td>
<td>23758</td>
<td>22388</td>
<td>14165</td>
<td>10013</td>
<td>22312</td>
<td>1945</td>
<td>3.08</td>
<td>53.70%</td>
</tr>
<tr>
<td>Average of Passed Samples</td>
<td>3889</td>
<td>16436</td>
<td>42231</td>
<td>28927</td>
<td>20351</td>
<td>14855</td>
<td>9750</td>
<td>2687</td>
<td>1945</td>
<td>2.81</td>
<td>52.63%</td>
<td></td>
</tr>
<tr>
<td>Ecolv0_Diagl_Mill</td>
<td>1738</td>
<td>1675</td>
<td>20489</td>
<td>23870</td>
<td>21640</td>
<td>9339</td>
<td>9052</td>
<td>1480</td>
<td>1947</td>
<td>3.54</td>
<td>24.83%</td>
<td></td>
</tr>
<tr>
<td>Ecolv0_Diagl_green</td>
<td>492</td>
<td>1698</td>
<td>29490</td>
<td>21784</td>
<td>16431</td>
<td>12763</td>
<td>8223</td>
<td>2694</td>
<td>1844</td>
<td>2.38</td>
<td>24.84%</td>
<td></td>
</tr>
<tr>
<td>Ecolv0_Diagl_red</td>
<td>4848</td>
<td>1675</td>
<td>20489</td>
<td>23870</td>
<td>21640</td>
<td>9339</td>
<td>9052</td>
<td>1480</td>
<td>1947</td>
<td>3.54</td>
<td>24.83%</td>
<td></td>
</tr>
</tbody>
</table>

**Figure 2. Error Distribution.** (a) MS1 Mass Error Distribution. (b) MS2 Mass Error Distribution.

### 6.12.1 Identification Quality Control (QC) - Summary

The QC Summary page gives an overview of the passed and failed samples with the selected attributes.

Click on Summary Filter to open the QC summary filter which provides the option to choose samples and attributes to check whether they pass the acceptance tolerance with respect to the Control sample.
To adjust the control sample or acceptance tolerance, another analysis will need to be run. Simply click on Modify Analysis and adjust those two settings in the QC step, leaving other parameters the same.

Table 1 provides a quick glance on the number of passed and failed samples from the selected samples.

Table 1. Statistics of filtered results.
Selected Samples Count  5
Passed Samples Count     2
Ratio of Passed Samples  40%
Failed Samples Count     3
Ratio of Failed Samples  60%
Total Failed Attributes  3

Figure 1 shows a histogram of the failed attributes and the number of samples with that failed attribute.
Table 2. Statistics of LFQ Result

<table>
<thead>
<tr>
<th>Sample Name</th>
<th># MS1</th>
<th># MS2</th>
<th># PSM</th>
<th># Identified Features</th>
<th># Quantified Features</th>
<th># Peptides</th>
<th># Quantified Peptides</th>
<th># Protein Groups</th>
<th># Quantified Protein Groups</th>
<th># PSM / # Peptides</th>
<th>ID Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control Sample</td>
<td>4848</td>
<td>16605</td>
<td>43989</td>
<td>31758</td>
<td>22988</td>
<td>14505</td>
<td>10013</td>
<td>2315</td>
<td>1945</td>
<td>2.83</td>
<td>53.76%</td>
</tr>
<tr>
<td>Average of Passed Samples</td>
<td>4889</td>
<td>16456</td>
<td>42231</td>
<td>28927</td>
<td>20351</td>
<td>14955</td>
<td>9750</td>
<td>2687</td>
<td>1945</td>
<td>2.83</td>
<td>52.61%</td>
</tr>
<tr>
<td>Ecoli30_Offgel_black</td>
<td>4738</td>
<td>16750</td>
<td>29489</td>
<td>23870</td>
<td>21640</td>
<td>8330</td>
<td>9052</td>
<td>1489</td>
<td>1847</td>
<td>3.54</td>
<td>34.88%</td>
</tr>
<tr>
<td>Ecoli30_Offgel_green</td>
<td>4892</td>
<td>16982</td>
<td>29400</td>
<td>21784</td>
<td>16431</td>
<td>12363</td>
<td>8223</td>
<td>2694</td>
<td>1844</td>
<td>2.38</td>
<td>34.84%</td>
</tr>
<tr>
<td>Ecoli30_Offgel_red</td>
<td>4884</td>
<td>16789</td>
<td>29939</td>
<td>21112</td>
<td>20661</td>
<td>10970</td>
<td>9682</td>
<td>2177</td>
<td>1939</td>
<td>2.73</td>
<td>34.84%</td>
</tr>
</tbody>
</table>

Table 2 provides more details on which particular samples have failed attributes, and the average values for those passed samples. The columns in this table were selected as attributes of interest, so they will not align with all attributes that were selected in the Summary filter.

Figure 2. Error Distribution. (a) MS1 Mass Error Distribution. (b) MS2 Mass Error Distribution.

Figure 2 shows how the MS and MS/MS distribution compares to the control sample.

Table 3. QC Parameters

Control Sample: Ecoli10_Offgel_black
Acceptance Tolerance: 20.0%
Total Sample Count: 6
QC Attributes:
# Identified Features
# Peptides
Quantified Peptides
Standard Peptides:
GFQVYFESPADAK, 2
GVAINMVTIEDK, 2
ITLDVAYMER, 2
VDQSAVGFEYQGK, 2
LLKPDIER, 3
LLKPDIER, 2
TFIAIKPDGVQR, 2
TFIAIKPDGVQR, 3
TPFLEVMQR, 2
ALAAQNIVDMEQR, 2

Table 3 simply shows the QC parameters that were set in this analysis. The Standard Peptides list here can be highlighted and copied, allowing for easy re-use of the standard peptides in another analysis.
6.12.2 Identification Quality Control (QC) - Data QC

The Data QC pertains to quality control measures related to the data itself. The following columns are in the table:

- **Sample Name**: Name of each sample.
- **All Attributes**: Pass/Fail flag. If a sample fails ANY attribute (not just in Data QC), it will be marked as Fail.
- **#MS1**: The number of MS scans in the last fraction in each sample.
- **#MS2**: The number of MS/MS scans in the last fraction in each sample.
- **MS2/MS1 Rate**: Calculated as #MS2/#MS1.
- **# Detected Feature**: The number of features in each sample.
- **Full Width (min)**: Measures the width of the median eluted peptide feature for each sample.
- **FWHM (sec)**: Abbreviation for Full Width at Half Maximum. This measures the number of seconds at half the maximum abundance of the median eluted peptide feature for each sample.
- **BPC**: Shows the BPC intensity of each sample.
- **TIC Correlation**: The Pearson correlation coefficient with respect to points on the control sample TIC.

The table has two selection controls. When selecting a cell, the column and row are highlighted. The highlighted column will update the histogram, showing the distribution between the control sample and all other samples.
The highlighted row is for comparing the control sample and the sample in the selected row. A Sample TIC chart is provided to examine how the TIC curve compares. Click on the button to change the view so that the curves overlap. The MS2 Count over Retention Time chart compares the cumulative MS/MS count as Retention time increases.

Right-click on the charts to export the image at any time.
6.12.3 Identification Quality Control (QC) - Identification QC

Similar to Data QC, the Identification QC tab also provides a table with statistics of interest and relevant charts from Identification results. If PEAKS PTM Search or SPIDER was included in the analysis, then the Identification QC will consider results from the last result generated in the workflow, otherwise it will use the DB Search result.

The table contains the following columns pertaining to the Identification result:

- **Sample Name**: Name of each sample.
- **All Attributes**: Pass/Fail flag. If a sample fails ANY attribute (not just in Identification QC), it will be marked as Fail.
- **# MS2**: The number of MS2 scans in each sample.
- **# Identified Feature**: The total number of features with identified spectra.
- **# PSM**: Total number of PSMs identified in the sample.
- **# Peptide**: Total number of distinct peptides identified in the sample.
- **# Sequence**: Total number of distinct peptide backbones identified in the sample.
- **# Protein Group**: Total number of protein groups identified in the sample.
- **# Top Protein**: Total number of top proteins identified in the sample.
- **# All Protein**: Total number of proteins identified in the sample.
- **Missed Cleavage Ratio**: The ratio of identified peptides with missed cleavages versus peptides without any missed cleavages.
- **Enzyme Specificity Ratio**: The ratio of identified peptides with specific digest versus peptides without.
- **# PSM / # Peptide Rate**: The ratio of # PSM and # Peptide for each sample.

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>All Attributes</th>
<th># MS2</th>
<th># Identified Feature</th>
<th># PSM</th>
<th># Peptide</th>
<th># Sequence</th>
<th># Protein Group</th>
<th># Top Protein</th>
<th># All Protein</th>
<th>Missed Cleavage Ratio</th>
<th>Enzyme Specificity Ratio</th>
<th># PSM / # Peptide Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>E.coli10.Offgel_black</td>
<td>Pass</td>
<td>81330</td>
<td>317150</td>
<td>63080</td>
<td>14395</td>
<td>13172</td>
<td>2051</td>
<td>2083</td>
<td>2772</td>
<td>0.01</td>
<td>0.01</td>
<td>6.98</td>
</tr>
<tr>
<td>E.coli10.Offgel_green</td>
<td>Pass</td>
<td>83988</td>
<td>31970</td>
<td>43622</td>
<td>16096</td>
<td>15540</td>
<td>2065</td>
<td>2772</td>
<td>3210</td>
<td>0.01</td>
<td>0.01</td>
<td>2.69</td>
</tr>
<tr>
<td>E.coli30.Offgel_red</td>
<td>Fail</td>
<td>77486</td>
<td>25978</td>
<td>41141</td>
<td>13624</td>
<td>13017</td>
<td>2689</td>
<td>2792</td>
<td>3124</td>
<td>0.04</td>
<td>0.01</td>
<td>2.68</td>
</tr>
<tr>
<td>E.coli30.Offgel_black</td>
<td>Fail</td>
<td>84599</td>
<td>23870</td>
<td>29489</td>
<td>8510</td>
<td>17865</td>
<td>1480</td>
<td>1543</td>
<td>1820</td>
<td>0.07</td>
<td>0.04</td>
<td>2.54</td>
</tr>
<tr>
<td>E.coli30.Offgel_green</td>
<td>Fail</td>
<td>84182</td>
<td>21784</td>
<td>29400</td>
<td>12360</td>
<td>11778</td>
<td>2694</td>
<td>2895</td>
<td>3194</td>
<td>0.08</td>
<td>0.04</td>
<td>2.39</td>
</tr>
<tr>
<td>E.coli30.Offgel_red</td>
<td>Fail</td>
<td>88625</td>
<td>21112</td>
<td>29039</td>
<td>10970</td>
<td>10483</td>
<td>2177</td>
<td>2265</td>
<td>2565</td>
<td>0.08</td>
<td>0.04</td>
<td>2.73</td>
</tr>
<tr>
<td>E.coli30.Offgel_black</td>
<td>Fail</td>
<td>83070</td>
<td>28803</td>
<td>36213</td>
<td>12546</td>
<td>12208</td>
<td>2340</td>
<td>2428</td>
<td>2776</td>
<td>0.06</td>
<td>0.03</td>
<td>1.30</td>
</tr>
</tbody>
</table>

The charts above for Protein Group Identification By Sample, Protein Overlap between Selected Samples, Peptide Sequence Overlap between Selected Samples, Peptide Frequency Distribution, and MS1 and MS2 Mass Error Distribution provide visual insights into the distribution and overlap of identified features and PSMs among different samples. The diagrams allow for a comparative analysis of sample performance and peptide specificity across the identified samples.
• MS1 Error Mean (SD): Shows the MS1 error mean in ppm and the standard deviation
• MS2 Error Mean (SD): Shows the MS/MS error mean in Da and the standard deviation
• ID Rate (%): Calculated as percent of #PSM / # MS2

Row and Column selection works the same as Data QC, where the selected Column will update the histogram to show the distribution of the column for each sample, and the selected Row will update the charts according to how they compare with the control sample. The charts include a venn diagram for protein and peptide overlap, a #PSM/#Peptide distribution, and the MS1 and MS2 error distributions.

Right-click on the charts to export the image at any time.

6.12.4 Identification Quality Control (QC) - Standard Peptides
When standard peptides were added to QC function, an additional tab is in the QC result node. A table will show which peptides are found in each sample, and the RT and m/z for those standard peptides can be viewed for quick comparison between samples.

Each sample and the standard peptides are listed in the table. The table can be sorted, but the control sample will always be the first index. Click on the columns to select/deselect that standard peptide for viewing in the RT and M/Z reproducibility charts. Multiple columns can be selected, and if none are selected then all standard peptides will be displayed. For those standard peptides of interest, there are checkboxes on the right side of the charts which can be toggled to show missing values. Those samples that are missing will have a vertical bar drawn, so that it is easier to tell which samples are missing the standard peptides.
Right-click on the charts to export the image at any time.

6.12.5 Identification Quality Control (QC) - QC Result Exports
As mentioned, all images in the QC result node can be exported by right-clicking on the chart and selecting Export image.

The Export Chart pop-up will appear, and the image can be exported at different scales.

Besides image exporting, the tables can be exported in CSV files and each QC view can be exported in HTML format. Double-click the node to find the QC Export options.
7. DIA Workflow (Spectral Library, DB Search, De Novo, Quantification)

PEAKS Library Search offers an accurate and sensitive identification method for complex spectra, such as DIA spectra. In addition to annotated spectra along with precursor m/z and charge, PEAKS Library search also uses predicted retention time and ion-mobility collisional cross section (CCS) data for improved sensitivity.

To start a DIA analysis, ensure all samples selected for the workflow have DIA Acquisition mode.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Enzyme</th>
<th>Instrument</th>
<th>Fragment</th>
<th>Acquisition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>Trypsin</td>
<td>Orbitrap (Orbi-Orbi)</td>
<td>HCD</td>
<td>DIA</td>
</tr>
</tbody>
</table>

7.1 DIA Analysis Workflow and Parameter Settings

To run a DIA Identification Analysis, Select Identification in Workflow Selection

The Spectral Library, Database Search, and De Novo are all optional steps in the DIA workflow. Any combination of these analysis can be run, but at least one of these steps must be included in the analysis.

7.1.1 DIA Identification - Spectral Library Search parameters

The first step is the optional Spectral Library Search.
The following parameter settings are provided:

**Precursor Mass Error Tolerance:** The maximum mass difference between the peptide and the monoisotopic mass of the precursor ion.

**Precursor Mass Tolerance unit:** PEAKS generates peptides within the precursor mass error tolerance. Precursor mass error tolerance can be specified in either Daltons or ppm.

**Fragment Mass Error Tolerance:** PEAKS uses this value when scoring peptide sequences. PEAKS Studio considers a fragment ion to be matched if the calculated m/z is within the specified error tolerance. Units are Daltons.

Both precursor and fragment error tolerance parameters need to be set consistently with the mass accuracy of the instrument.

**Optimize Tolerance:** When enabled, PEAKS Studio will determine the optimal values to use for Precursor and Fragment Mass Error Tolerance for each fraction in your analysis. These values will replace the values provided in the previous Precursor and Fragment Mass Tolerance sections.

**Library Selection:** Select a PEAKS-generated library for the search. Select one from the list of libraries that have been configured in PEAKS Studio and filter for the peptide length, precursor m/z, and charge when necessary. To configure a new library, click on the button. For more information on Spectral Library Configuration, refer to 2.10 Spectral Library Configuration. Also of interest is the chapter on Library Viewer.

**Library Filters:** The spectra that will be used from the library can be filtered based on peptide length, precursor m/z, and charge. This is optional. Each filter can be enabled by checking the checkbox to the left of the filter inputs.

Protein inference can optionally be enabled to search peptides found in the selected library against a provided database.

**Target Database:** Select a reference database from the list of configured databases or create a new one, and set the taxonomy if applicable. This database is the database used for protein inference.

**Contaminant Database:** A contaminant database can optionally be provided as well. #CONTAM# will be added to the beginning of the accession of all identifications from the contaminant database.

### 7.1.2 DIA Identification - DIA DB Search parameters
PEAKS DIA DB can be launched as a standalone workflow, or launched on top of a PEAKS Library Search workflow. It is an optional step in the workflow.
The Precursor and Fragment Mass Error Tolerance are set and shared in the previous Spectral Library step.

**Enzyme Settings:** Select the enzyme used for protein digestion from the enzyme drop-down menu.

**Note:** "None" is a special enzyme allowing non-specific cleavage at both ends of the peptide. It is the recommended option when no digestion enzyme was used or when the digestion enzyme exhibits a high degree of non-specificity.

**Note:** When the selected dataset is digested with different enzymes, "Specified by each sample" allows samples to be analyzed separately using their respective enzymes specified during project creation.

**Digest Mode:** This specifies the type of in silico cleavages allowed at a peptide termini based on the method used for protein digestion. If "Specific" is selected, the specificity of the selected enzyme is strictly enforced at both termini of a peptide. "Semispecific" allows one terminus to disobey the enzyme specificity rules.

**Note:** If the enzyme is specified as "None", then no matter which mode is selected for "Digest Mode", the "None" enzyme digest rule will be applied.

**Missed Cleavages:** This specifies the number of missed cleavage sites allowed in a peptide.
**PTM Settings:** Click “Set PTM” to open the **PTM Options** dialog and specify the fixed PTMs and a few common variable PTMs expected in the experiment. Highlight PTMs and click on the button to place them as either Fixed or Variable PTM. From there, highlighted PTMs can be removed by clicking the “Remove” button. Clicking “Remove All” will reset the PTM options. Highlighting and clicking “Switch Type” will move the Fixed PTM in into the Variable PTM list and vice-versa. Click on “New” to create a new custom PTM. While a PTM is selected, click on “View” to see more details.

**Note:** Multiple PTMs can be highlighted by holding down the CTRL key or SHIFT key on the keyboard.

**Note:** For a specific residue, only one fixed modification is allowed.

**Note:** For DIA Database search, currently only the following PTMs are supported: Carbamidomethylation, Oxidation (M), Deamidation (NQ), and Acetylation (Protein N-term).

Once the PTMs are selected, the maximum number of variable PTMs per peptide can be specified and a value of less than 4 is recommended.

**Target Database:** Select a protein sequence database for the search. Select one from the list of databases that have been configured in PEAKS and set the taxonomy by clicking on , if applicable. To configure a new database, click on next to the Target Database dropdown to open the Database Configuration dialog.

Note: If running Spectral Library search, the database selected here will be used for protein inference, regardless if it was checked or not.

**Contaminant Database:** To search for contaminants, select a contaminant database from the list of databases that have been configured in PEAKS. Keep the selection as N/A if no contaminant database is required for the search.

**Database search filters:** The peptide length, fragment m/z, precursor m/z, and charge are filters applied to the final database search result.

The maximum peptide length for the DIA DB Search algorithm is 30. Setting a maximum length beyond 30 will still report peptides with maximum length of 30.

**7.1.3 DIA Identification - DIA De Novo parameters**

PEAKS DIA De Novo is an optional step that can be launched as a standalone workflow, or launched on top of a PEAKS Library Search, PEAKS DIA Database Search, or PEAKS Library Search + PEAKS Database Search workflow.
The Precursor and Fragment Mass Error Tolerance are set and shared in the Spectral Library step.

Refer to the previous section 7.1.2 DIA Identification - DIA DB Search parameters for details on configuring the Enzyme and PTM settings.

### 7.1.4 DIA Quantification (DIA LFQ) Analysis Workflow and Parameter Settings

DIA Label Free Quantification requires at least 2 samples in the analysis. It is recommended to only run LFQ with 1 data file per sample, but if you choose to have fractionated samples, ensure each sample has the same number of fractions. All samples must have DIA Acquisition mode set.
To run the DIA Label Free Quantification workflow, choose the Quantification option in Workflow Selection.

The Spectral Library Search step can be selected and details are provided in section 7.1.1 DIA Identification - Spectral Library Search parameters.

The DIA DB Search step can be selected and details are provided in above section 7.1.2 DIA Identification - DIA DB Search parameters.

Select one or both of these identification steps in order to run DIA Label free quantification.

After deciding on the DIA Identification parameters, click on to proceed to the DIA label free quantification parameters page.

The following parameters are provided for Label Free Quantification (LFQ):

**Sample Group:** LFQ requires sample grouping. All available samples are listed in the left window. After selecting a sample, it can be added to a new group by clicking the button or to an existing group by clicking the button and selecting that option from the drop-down menu. If you select multiple files, you can add each
one individually to create multiple new groups by clicking the button. Samples and Groups can be removed by clicking the button next to each element. The names and colours for each group can be edited by clicking on the coloured square dropdown, respectively.

**Match Between Run:**

**Retention Time Shift Tolerance:** The retention time shift tolerance is the maximum elution time difference that is considered for the quantification of an identified peptide after RT alignment. Select “Auto Detect” to let PEAKS Studio detect the optimal Retention Time Shift Tolerance for your data based on the observed trend.

**Ion Mobility Tolerance (1/k0) (if applicable):** The maximum 1/k0 differential for features to transfer IDs. 

*Note:* This parameter is only available when the analysis contains timsTOF data.

**Feature intensity:** Set a value for the minimum intensity a peptide feature must exhibit.

**RT Range:** Feature vectors within the selected retention time range will be included in the results while everything else will be removed.

**Base Sample:** The base sample will act as the reference. The peptide and protein abundance corresponding to this reference label will be the denominator in all of the abundance ratio calculations.

**Peptide Feature Filters:**

- **Avg. Area:** The minimum average area of all peptide features associated with the peptide. Features with higher abundance have been shown to be more reproducible across replicates. Only peptide features with an average area above this threshold will be used for quantification.
• **Quality:** The quality score for the peptide feature. Factors that affect the quality score include m/z difference, RT difference, XIC shape similarity, and the feature intensities.

• **Peptide ID Count:** This filter sets the minimum limit for how many times a peptide must be identified within a group to be included in the LFQ results.

• **Charge between:** Only peptides with feature vectors that all fall within this range will be used in protein quantification.

• **Have at least ___ confident samples per group:** This filter sets the minimum number of sample(s) per group that the peptide feature has an abundance value. A peptide is more quantifiable when it is detected in more samples.

• **Use in group Coefficient of Variation Filter:** For each feature vector (set of peptide features associated with a peptide), at least one group must have a coefficient of variation (CV) less than the CV threshold. The CV threshold is automatically selected by the software based on the current data set.

**Protein Filters:**

- **Significance Method:** Protein abundance is used for the significance testing calculation. When there are replicates in each group, ANOVA is suggested to be used for significance testing. For more details on ANOVA, refer to the following paper: "On the comparison of several mean values: an alternative approach", *Biometrika*, 1951, 38(3/4): 330-336

- **Modified Form Exclusion:** The expression level of modified peptides might be different than the overall expression level of a protein. In such cases, including modified peptides for protein ratio calculation might lead to incorrect results. By checking this box, if a peptide has modified and unmodified forms, both will not be used as candidates for protein ratio calculation, if a peptide only has one modified form, it is still a candidate, but if a peptide has more than one modified form, all forms will not be used as candidates

- **Remove Outliers:** This filter removes supporting peptides that have a different variation trend compared to others.

- **Use Top # peptides:** This is the number of quantifiable supporting peptide that will be used to calculate the protein profile. Three is the default setting.
• **Significance:** Only protein groups with a significance above this threshold will be listed in the “Protein” view. The significance score is calculated as the $-10\log$ of the significance testing p-value (e.g. Significance score of 20 is equivalent to a p-value of 0.05). PEAKS provides the significance testing method: ANOVA.

• **FDR:** Adopt the Benjamin-Hochberg method to adjust the p-value to the false discovery rate for all protein groups that have already passed the other filters. Only protein groups with significance scores passing the calculated FDR will be listed in the "Protein" view. Either this or "Significance" can be selected to set a significance threshold.

• **Fold Change:** Only protein groups above the lower threshold or below the upper threshold value will be listed in the "Protein" view.

• **Used Peptides:** Only protein groups that were quantified with this number of peptides (or more) will be listed in the "Protein" view.

**Normalization Method:**

• **Use TIC:** By default, PEAKS Studio uses the total ion current (TIC) of the samples to calculate the normalization factors. Normalized abundance for each sample is calculated from the raw abundance divided by the normalization factor for that sample.

• **Manual input:** This option performs the same default calculation as auto normalization ("Use TIC") but also allows for the manual change of each sample’s expected ratios. This option should be used if the expected ratio is not 1:1 for each sample as the software attempts to normalize to a 1:1 ratio.

• **Use internal standard proteins:** This option displays a list of identified proteins (when a database or library search has been completed) that can be selected to create a normalization ratio. A search bar is included to enable the quick location of select proteins.

• **No Normalization:** All samples are automatically assigned Factors of 1.0.

• **Local RT Normalization:** Normalization factors are calculated based on the TIC of the local RT region. As such each normalization factor will adjust accordingly and peptides at nearby retention times would have similar normalization factors.
7.1.5 DIA Identification and Quantification QC

In PEAKS Studio 11, the next optional step is the newly introduced QC function. See section 7.22 DIA Quality Control (QC) for further explanation on how this function is utilized.

When QC is checked, a QC result node is added in the analysis. Select a control sample and acceptance tolerance, where the tolerance will mark a sample as pass or fail depending on the relation to the control sample. These two cannot be adjusted in the QC result node, and a new analysis would have to be run. The Data, Identification, and LFQ QC attributes are the attributes of interest. For Identification workflow, there will not be any LFQ QC attributes to select. If a sample is outside the acceptance tolerance in any of these attributes with respect to the control sample, it will be marked as a failed sample. These attributes can be changed in the QC Summary filter, so they can be adjusted without needing to run another analysis should you want to change the selections. Lastly the Standard Peptides list is used to specify peptides and their charge in the format shown (e.g. LLDMGET, 2). These standard peptides will have their RT and m/z mapped for each sample that reported the peptide and charge. At this time, only peptides without modifications can be used (so adding in a modification to the sequence will not work).

Click on “Report” to proceed to the final step. Report filters can be set here which will filter the final result.

Both the DIA Identification and DIA Quantification workflows have the same Report filter step.
These filters are provided:

**PSM/Peptide**: this section is a mandatory section to filter out some results under the user-defined filter. Any changes to the filter will create a new analysis with old parameters and update the results and statistics.

**PSM -10lgP**: The PEAKS peptide score (-10lgP) is calculated for every peptide-spectrum match (PSM) reported by PEAKS DB, PEAKS PTM, and SPIDER. The score is derived from the p-value that indicates the statistical significance of the peptide-spectrum match. A peptide may be matched to many spectra, resulting in multiple PSMs. In that case, the peptide’s score is calculated as the maximum among all PSM scores. For details of the scoring algorithm, please refer to the publication, "PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification", Mol Cell Proteomics, 2011 Dec 20. A minimum requirement can be set and all identifications must pass this filter.

**PSM FDR (%)**: The PSM filter can also be set to use the false discovery rate. FDR is estimated using a "decoy-fusion" approach.

**Peptide FDR (%)**: The Peptides can also be set to use the false discovery rate. Peptides are scored based on their top PSM and using a “decoy-fusion” approach, the result is reported at that FDR.

**Note**: Decoy fusion is an enhanced target-decoy method for result validation with FDR. Decoy fusion appends a decoy sequence to each protein as the "negative control" for the search. See BSI's web tutorial (http://www.bioinfor.com/fdr-tutorial/) for more details.
From here, there is the option to specify an Analysis Name (which can be later renamed) and/or decide to save the Workflow to quickly re-use in future analyses.

7.2 Understanding PEAKS Spectral Library Search Results

After the library searches are complete, library result nodes are generated. Double-click the node to examine the analysis report. The analysis report is presented in three pages:

- **Summary**: This shows an outline of the library search results with key statistics. The overall quality of the experiment can be examined and the filters for peptide and protein identifications can be adjusted.
- **Protein**: This shows a list of protein identifications. This view also visualizes protein sequence coverage and helps with protein characterization. This tab only appears if Protein Inference was enabled for the PEAKS Spectral Library Search.
- **Peptide**: This shows a list of peptide identifications. This view also provides spectrum annotation and other detailed information for peptide precursor spectrum matches.

7.3 Spectral Library - Summary View

The Library result **Summary View** is similar to the PEAKS database identification result.

The Summary view reports key statistics as an overview of the result. In the report, several statistical charts can be examined to assess result quality. At the top of each summary result page, protein filters can be set. Identifications that fall below the filters will not be visible in the result pages or exports.

**Proteins -10lgP**: The PEAKS protein score (-10lgP) is calculated as the weighted sum of the -10lgP scores of the protein’s supporting peptides. After removing any redundant peptides, the supporting peptides are sorted by -10lgP scores in descending order, and the k-th ranked peptide contributes to the weighted sum with a weight of 1/k. A default threshold of 15 is set.

**Proteins FDR**: The Proteins can also be filtered in accordance to the false-discovery rate of the protein set.

**Proteins Unique Peptides**: The minimum number of unique supporting peptides for a protein identification. A unique peptide is defined as a peptide that can be mapped to only one protein group.

After changing the filters, click **Apply** to apply the new filter and update the results accordingly.
1. Notes

2. Result Statistics

Figure 1. False discovery rate (FDR) curve. X axis is the number of peptides being kept. Y axis is the corresponding FDR.

Click on [Notes] to open the Notes dialog to save useful information regarding this Analysis. The Notes will appear in Section 1 of the summary page at the top.

The Summary view reports key statistics as an overview of the result. In the report, several statistical charts can be examined to assess result quality.

7.3.1 Spectral Library result - Figure 1. False Discovery Rate (FDR) curve.

Figure 1 shows the FDR curve with respect to the number of Peptides (or PSMs) being kept after filtration by the $-10\lg P$ score. By lowering the score threshold, more Peptides are kept in the filtered result. Conversely, the FDR increases because more false positives are kept. In this figure, the vertical line indicates the current score threshold. The number of Peptides and the corresponding FDR by the current score threshold are shown in the top-left corner. PEAKS Search estimates FDR using the "decoy-fusion" approach. Decoy-fusion is an enhanced
target-decoy approach that makes more conservative FDR estimations. For details of the "decoy-fusion" approach, please refer to the publication, "PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification", Mol Cell Proteomics, 2011 Dec 20.

It is commonly recommended to set FDR at <1% by adjusting the score threshold. If a rapid growth of the FDR around the 1% FDR threshold is noticed, some PSMs may be sacrificed to significantly reduce the FDR.

Note: When counting the number of PSMs, PEAKS Studio keeps one peptide per spectrum at most. Thus, the number of PSMs is actually the number of spectra with assigned peptides.

Note: During Analysis creation, if Peptide FDR (%) filter was used in Report step, the Summary page FDR curve in the results will show the Peptide FDR curve and corresponding number of Identified peptides.

Note: Decoy matches are not counted in the number of PSMs. Unless otherwise specified, decoy matches are also excluded from the other statistical numbers shown in the Summary view.

7.3.2 Spectral Library result - Figure 2. RT Calibration

**Figure 2. RT Calibration**

Plots the RT regression function used during the spectral library search. Peptides from the first MS Run in the analysis are plotted against their indexed retention time (iRT) predicted using machine learning. The $R^2$ value is presented based on the regression function.

7.3.3 Spectral Library result - Boxplot for RT change

**Figure 3. Boxplot for median ΔRT across all MS runs**
A box plot showing the difference between the observed retention time and iRT peptides within set retention time bins across all fractions, showing the accuracy of retention time prediction.

7.3.3 Spectral Library result - Table 1. Statistics of data

Statistics of Data tabulates MS1 and MS2 information, as well as identification in the peptide and protein level. The protein level statistics will only be presented if protein inference was performed. This table in the Summary view provides a condensed statistical information for the analysis.

<table>
<thead>
<tr>
<th></th>
<th>#MS Run</th>
<th>#Scans</th>
<th>#PSMs</th>
<th>#Peptides</th>
<th>#Sequences</th>
<th>#Proteins</th>
<th>#PSM/#MS2</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>2</td>
<td>4678</td>
<td>196476</td>
<td>59460</td>
<td>27883</td>
<td>27104</td>
</tr>
<tr>
<td>Sample 1</td>
<td>1</td>
<td>2339</td>
<td>98238</td>
<td>29541</td>
<td>26931</td>
<td>26225</td>
<td>3967</td>
</tr>
<tr>
<td>Sample 2</td>
<td>1</td>
<td>2339</td>
<td>98238</td>
<td>29919</td>
<td>27277</td>
<td>26552</td>
<td>3971</td>
</tr>
</tbody>
</table>

#MS Run: Total # of runs in each sample.

#MS1: Total # of MS1 spectra for each sample. For timsTOF data this shows the # MS1 frames.

#MS/MS: Total # of MS2 spectra for each sample. This does not include chimeric scans. For timsTOF data this shows the # MS2 frames.

#PSMs: The number of peptide-spectrum matches.

#Peptides: The unique number of peptide sequences with modifications.

#Sequences: The unique number of peptide sequences not including modifications.

#Proteins Groups: PEAKS Studio groups proteins identified by a common set of peptides. This number shows the number of protein groups in the filtered result based on All proteins.

#All Proteins: Total number of proteins with significant peptides, according to current protein filters.

#Top Proteins: Total number of top proteins with significant peptides, according to current protein filters.

#PSM/#MS2: The identification rate by dividing #PSM by #MS2 for the sample.

7.3.4 Spectral Library result - Result filtration parameters

Tables 2-4 show the statistical numbers of the data and results.

<table>
<thead>
<tr>
<th>Peptide -10 logP</th>
<th>≥ 0</th>
<th>≥ 35.9</th>
<th>≥ 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proteins unique peptides</td>
<td>≥ 1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Name</th>
<th>ΔMass</th>
<th>Position</th>
<th>#PSM</th>
<th>#Peptides</th>
<th>-10 logP</th>
<th>Area</th>
<th>AScore</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbamidomethyl</td>
<td>57.02</td>
<td>C</td>
<td>6310</td>
<td>2874</td>
<td>45.13</td>
<td>1.7E7</td>
<td></td>
</tr>
<tr>
<td>Deamidation</td>
<td>.98</td>
<td>NQ</td>
<td>1664</td>
<td>775</td>
<td>45.09</td>
<td>4.78E6</td>
<td></td>
</tr>
<tr>
<td>Oxidation</td>
<td>13.99</td>
<td>M</td>
<td>1452</td>
<td>690</td>
<td>45.07</td>
<td>4.92E6</td>
<td></td>
</tr>
</tbody>
</table>

| FDR (Peptide-Spectrum Matches) | 0.7% |
| FDR (Peptide Sequences)        | 1.0% |
| FDR (Protein Group)            | 1.0% |
Table 2. Result filtration parameters: Table 2 summarizes the result filtration parameters set at the top of the Summary View page. This information provides a helpful reference for exported data.

Table 3. Statistics of filtered results: Table 3 presents the statistics based on the data analysis results after setting the desired filters. See below for further explanation:

- **FDR (Peptide-Spectrum Matches):** The total number of decoy database assignments to spectra relative to the total number of target database assignments to spectra represented as a percentage.
- **FDR (Peptide Sequences):** The total number of decoy database assignments to unique peptide sequences relative to the total number of target database assignments to unique peptide sequences represented as a percentage.
- **FDR (Protein Group):** The total number of decoy protein groups where the top hit in the protein group is a decoy database protein relative to the total number of target database protein groups where the top hit is from the target database.

Table 4. PTM Profile: shows modifications identified in the filtered result and the number of PSMs containing each modification

7.3.5 Spectral Library result - Experiment Control

Figure 4 in the Summary View show the precursor mass errors of PSMs in filtered results. The precursor mass error is calculated in ppm as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$. Calibration is for display purposes only and PEAKS corrects m/z value for each PSM to correct systematic biases and instrument measurement drift.

![Figure 4. Distribution of precursor mass error of filtered PSM in ppm.](image)

Table 5 shows the number of identified peptides in each sample with the number of missed cleavages, indicating the enzyme digestion efficiency.

Table 5. Number of identified peptides in each sample by the number of missed cleavages.

<table>
<thead>
<tr>
<th>Missed Cleavages</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4+</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>26015</td>
<td>878</td>
<td>32</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>Sample 2</td>
<td>26352</td>
<td>888</td>
<td>31</td>
<td>6</td>
<td>0</td>
</tr>
</tbody>
</table>
7.4 Spectral Library result - Proteins View

The Proteins View is only displayed if Protein Inference was selected in Spectral Library parameter step (shown below) in Project Wizard. It will automatically be selected if DIA Database Search was performed. It will also have to be performed if running DIA Quantification workflow.

The Protein View lists protein identifications that have been filtered by the current settings in Summary View. It also visualizes the protein sequence coverage for identified proteins. The view has four main components:

- **Protein table**: This lists protein identifications that have been filtered by the current settings in the Summary View and the “Protein View Filters”.
- **Coverage**: This visualizes the protein sequence coverage of the selected protein, showing the mapping of support peptides and de novo tags to the protein sequence.
- **Peptides**: This lists supporting peptides mapped to the selected protein.

### 7.4.1 Proteins View - Protein Table controls

At the top of the Proteins View, there are the options to filter, show/hide columns, search, and export the protein table.
7.4.1.1 Protein Table controls - Protein View Filters

Click on the button to open the Protein View Filter.

The following options are available for filtering the protein table:

**Show Top, All, or First proteins in each group:** Adjust the protein list based on Protein grouping, top is selected by default. Proteins are grouped based on parsimony.

- **Top:** Show only the top proteins in each group. These proteins are supported by the most unique peptides in the group. Proteins in the group that share a subset of the unique peptides that support the top protein (subproteins) will not be displayed.
- **All:** Show all proteins in each group. Proteins are grouped together if they are supported by the same set or a sub-set of the top protein in the group. If a protein is supported by a peptide not supporting the top protein it will be added to a new group.

**Protein accession/name contains:** Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

**Protein description contains:** Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

**Note:** In these above two filters, multiple entries can be separated by a semi-colon.

**Protein sample area:** Filter proteins based on total protein area found in each sample. Sample areas are calculated using the total of all peptide features from unique supporting peptides. If a protein is identified and has areas from multiple samples, only one of the samples must pass this filter to be included.

**PTM contains:** Filter protein results by the presence of selected PTMs. Proteins that are supported by peptides containing modifications that match the selected PTMs will be included. If multiple PTMs match the modification name, they will all be included in the table. For SPIDER results, the mutations will also be listed in the PTM contains filter.
Click on the **Reset** button to remove all filters.

Click on **OK** to apply the Protein View Filter to the Protein table.

Click on **Cancel** to discard changes and close the dialog.

### 7.4.1.2 Protein Table controls - Protein count

At the top of the protein table shows the count of protein groups and number of proteins in the table based on the current Protein View Filters.

57 protein groups, total 142 proteins

### 7.4.1.3 Protein Table controls - Optional columns

Click on the **button** to see a list of sample-based optional columns.

Select the corresponding sample-based Coverage, Spectra count, or Area to add into the Protein Table view.

- **Coverage by Sample**: The protein coverage as a percentage will be displayed for the individual sample when selected.
- **#Spec by Sample**: The total number of spectra identified that support the given protein. Totals are given for each sample included in the search.
- **Area by Sample**: The total area of peptide features from unique supporting peptides in each sample will be displayed.

The exports will always include the contents of these optional columns even if they are hidden.

### 7.4.1.4 Protein Table controls - Protein Table Search Function

An easy-to-use search function is available for the protein table. The protein accession or description substring can be searched for, and all search results can be navigated through with the Up and Down arrows.
Protein accession/name contains: Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

Protein description contains: Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

7.4.1.5 Protein Table controls - Protein Table Export

Click on the button to export the contents of the current table. This considers the Protein View Filters, the sorting applied to the table and is consistent with the view. There is the option to export the Protein table and/or the Supporting peptide table in CSV format.

7.4.2 Proteins View - Protein Table

Each row in the table represents a group of proteins that are supported by a common set of peptides. To expand the group, click the right arrow button at the left.

The columns in the Protein Table are:

- **Accession**: The accession number of the protein as seen in the FASTA database.
- **Cluster**: A unique identifier that classifies which protein group the protein belongs to.
- **Top**: Indicates whether this protein is a top protein.
- **-10lgP**: The protein confidence score.
- **Coverage (%)**: The percentage of the protein sequence that is covered by supporting peptides. The coverage is visualized by a colored bar. Light blue blocks indicate the parts of the sequence covered by low-confidence peptides. Dark blue blocks indicate the parts covered by high-confidence peptides. This is the total coverage including results from all samples.
• **#Peptides**: The number of high-confidence supporting peptides.
• **#Unique**: The number of high-confidence supporting peptides that are mapped to only one protein group. Unique peptides with same sequence but different modifications are only counted once in this number.
• **PTM**: The identified modifications displayed with color-coded icons.
• **Avg. Mass**: The protein mass calculated using the average mass.
• **Description**: The protein's header information as seen in the FASTA database.

**Note**: For #Peptides and #Unique, two peptides with the same starting and ending positions in the protein are counted as one, regardless of their PTM forms. This is to follow the MCP (Molecular & Cellular Proteomics) guidelines.

### 7.4.3 Proteins View - Coverage Pane
The Protein Coverage view visually maps the supporting peptides and de novo tags to the protein selected in the Protein table. It also shows all identified sites with modifications or mutations to assist with protein characterization at the amino acid level. The "Protein Coverage" view is composed of three major components - Protein Sequence Coverage, Coverage Control Panel, and Protein Tools.

#### 7.4.3.1 Coverage Pane - Protein Sequence Coverage
This area visualizes the coverage of the protein sequence.

Regions in the protein sequence that are covered by supporting peptides are displayed in bold font with a grey background. Confident modifications and mutations identified in supporting peptides are displayed as icons above the protein sequence. Modifications are represented by colored icons with the first letter of its modification name. If a residue is modified by more than one modification in the same supporting peptide, "**" is used instead of a letter. The header information of the protein is shown on the top of the protein sequence.

The supporting peptides can be shown as colored bars under the protein sequence.

Placing the cursor over a bar shows a tooltip with detailed information of the peptide. Left clicking on a bar shows the annotated spectrum from which the supporting peptide is identified.
Right clicking on a peptide bar shows additional options. “Show in spectrum” has the same behaviour as left clicking on the peptide bar which is to open the annotated spectrum chart. “Show in table” has the function to jump to the supporting peptides table.

7.4.3.2 Coverage Pane - Coverage Control Panel
This panel controls what to display in the protein sequence coverage view.

Sequence Display Mode: The protein sequence can be displayed in either "outline" or "coverage" mode.

Outline mode: In outline mode, the protein sequence is displayed without supporting peptides and matched de novo tags. However, the sequence coverage at a specific position can be examined by left-clicking a residue.

Covered mode: In coverage mode, all the supporting peptides and matched de novo tags are shown.

AAs per line and 10AA gap: Specify the number of amino acids per line and whether to show a gap for every 10 amino acids in the protein sequence.

Sequence Display Option: Select whether to view the protein sequence as Fasta or a specific enzyme used in the project. When a particular enzyme is selected, the protein sequence will be highlighted to emphasize expected peptides generated by the selected enzyme's cleavage specificity.
Modification table: The modification table shows the modifications identified in supporting peptides on the protein. For each modification, the number of supporting PSMs with this particular modification is shown. The checkbox on the left controls whether to indicate the modification in the protein coverage.

<table>
<thead>
<tr>
<th>AM</th>
<th>PTM</th>
<th>#</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>Carboxymethylation</td>
<td>40</td>
</tr>
<tr>
<td>O</td>
<td>Oxidation (M)</td>
<td>4</td>
</tr>
<tr>
<td>D</td>
<td>Deamidation (NQ)</td>
<td>4</td>
</tr>
</tbody>
</table>

7.4.3.3 Proteins View - Protein Tools
The Proteins View toolbar is at the upper-right corner of the protein sequence coverage tab and contains helpful protein tools.

Full screen: When selected, the protein coverage window expands in full screen mode. Click to return from full screen mode.

Tool box: The tool box contains the following tools:
- Copy protein sequence. This copies the protein sequence of the current selected protein into the system clipboard.
- Save protein coverage as image. Different scale options are provided to increase the image size.
- Coverage statistics. This provides useful statistics based on the current selected protein.

7.4.3.4 Proteins View - Supporting Peptides
The "Peptides" tab displays a table of supporting peptides for the protein of interest. This table is similar to the peptide table in the Peptide view, except that "Accession" is excluded and the columns "(Checkbox)", "Unique", "Start", and "End" are included. Descriptions of extra columns are given in the following sections.

- (Checkbox): This controls whether a peptide is shown in the protein coverage as a blue bar.
- Unique: This shows whether the peptide is a unique supporting peptide to the protein group.
- Start: This shows the peptide's starting position in the protein.
- End: This shows the peptide's ending position (inclusive) in the protein.
7.5 Spectral Library Result - Peptide View

The "Peptide View" displays the resulting peptide identifications that have been filtered by the current settings in Summary View and "Peptide View Filters". The table on the top shows the details of each peptide identification. The bottom section provides additional information about the peptide-spectrum matches for the native peptide selected in the table.
7.5.1 Peptide View - Peptide Table
The Peptide Table shows the filtered native peptide identification results. Each row in the table is a native peptide identification represented by its highest-scoring PSM. When there are more than 1000 peptides, the table is split into multiple pages.

Note: If there are multiple PSMs matched to a peptide, the table displays the information for the top-scoring PSM. Peptides with the same sequence but different modifications are considered different and shown in different rows.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence[s]”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

The following list describes the contents in each of the default columns:

- **Peptide**: The amino acid sequence of the peptide, as determined by the PEAKS search workflow. A modified residue is followed by a pair of parentheses enclosing the modification.
- **-10lgP**: The peptide -10lgP score. The score indicates the scoring significance of a peptide-precursor spectrum match.
- **Mass**: The monoisotopic mass of the peptide.
- **Length**: The number of amino acids in the sequence backbone.
- **ppm**: The precursor mass error, calculated as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$.
- **m/z**: The precursor mass-to-charge ratio.
- **RT**: The retention time (elution time) of the spectrum as recorded in the data.
- **delta RT**: The difference between the query and library RT.
- **Area per Sample**: The combined area for all features in that sample associated with this peptide.
- **Scan**: The scan number of the spectrum that matches the peptide sequence with the highest -10lgP.
- **#Spec**: Number of MS2 spectra assigned to the peptide.
- **Accession**: The accessions of proteins that contain this peptide.
- **PTM**: The types and the numbers of modifications present in the peptide shown in color-coded icons.
- **Found By**: The name of the PEAKS search workflow that identifies the peptide; this can only be Spectral Library for this result.
7.5.2 Peptide View - Peptide Table controls
The menu above the peptide table provides options to scroll through the table of peptides, filter the table, show optional columns and search for specific entries.

7.5.2.1 Peptide Table controls - Pagination
For easy viewing and to improve performance, the peptide table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.

In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

7.5.2.2 Peptide Table controls - View Filter
Click on the Peptide View Filter button to open the Peptide View Filter which is used to filter the Peptide table. The following are the filter options provided:

- **Peptide sequence contains**: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains for further filtering if necessary.

- **Peptide score (-10lgP)**: Filters out all peptides with -10lgP scores below this threshold.
• Peptide length: Filters for peptides of length within this min/max threshold.

• Peptide sample area: Filters based on total peptide area found in each sample. Sample areas are calculated using the total of all peptide features from all spectra that identified the native peptide. If the peptide is found in multiple samples, only one sample must pass this filter for the peptide to be included.

• PTM contains: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text they will all be included in the table.

Note: In all filters, multiple entries can be separated by a semi-colon.

Click on the Reset button to reset all filters to their default setting.

Click on OK to apply the filters to the Peptide table.

Click on Cancel to discard the changes in the View Filter.

7.5.2.3 Peptide Table controls - Optional Columns

Click on the button to open the optional columns dropdown. By default the sample area columns are displayed. Sample Spec count columns can also be shown. These sample spec counts are always from identification, and does not consider whether there are feature vectors present. For projects with many samples, it might be useful to hide these optional columns and reduce the width of the peptide table.

7.5.2.4 Peptide Table controls - Search function

On the top right of menu, there is a search function with 4 options. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

• **Scan =:** Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction.

• **m/z =:** Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit. For example, searching for m/z = 560.21 will find all m/z between [560.20, 560.22]

• **RT =:** Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/- 1 of the last significant digit set. For example, searching for RT = 55.5 will find all m/z between [55.4, 55.6]

• **Seq contains:** Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence
with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered. Additionally, since the peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”.  

7.5.3 Peptide View - Feature selection breadcrumb
There is a sample and feature selection breadcrumb which allows you to switch between features associated with the peptide. Click on the Sample in the breadcrumb to reveal other samples and the features in that sample related to the current peptide selection. From here you can select other samples, fractions, or features.

By clicking on the fraction level, all fractions in this sample are revealed and the features in each fraction associated with this peptide. From here you can select other fractions and features to view.

Lastly, clicking on the feature level will show all features in the currently selected fraction. Features with Feature Id = 0 are known as fake features and are constructed to enhance identification.

7.5.4 Peptide View - PSM selection
Underneath the feature breadcrumb will shown the currently selected PSM. Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.

All other components in the Peptide tab interface will update according to the currently selected PSM.

For diaPASEF instruments, Frame is used rather than Scan number.

7.5.4.1 Peptide View - Protein jump button
Click on the protein jump button to jump to a protein that contains the currently selected PSM. If there are more than 1 protein, a dropdown is revealed which allows the user to select which protein to jump to. If a protein is filtered out due to a
protein summary filter or summary page protein filter, then the option to jump will be grayed out. If there is no protein tab in the result, then the button will always be disabled.

7.5.5 Peptide View - Annotated Spectrum Chart

Click on button to switch from the Mirror plot view and annotated spectrum chart view.

For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

7.5.5.1 Annotated Spectrum Chart - Chart Navigation

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.

Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.

Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.

See the whole spectrum: Double click in the spectrum or click the 1:1 button.

7.5.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings

Click on the button to show the spectrum annotation settings.
The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places**: Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation**: Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated**: Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation**: Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info**: Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity**: Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

### 7.5.3.5 Annotated Spectrum Chart - Additional Chart Controls

Click on 📊 or 📈 to toggle between absolute and percentage intensity values in the Y axis.

Click on 🔍 to reset the zoom level of the annotated spectrum chart to the default.

Click on 📈 to zoom into the X axis by a factor of 2.

Click on 📈 to zoom into the Y axis by a factor of 2.
The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

7.5.5.4 Annotated Spectrum Chart - Mirror Plot

Spectral Library results have a Mirror Plot view. Click on the button to switch to the Mirror Plot.

This view shows the ions from the query spectrum compared to the library spectrum.

Hold down CTRL when scrolling the mouse wheel to zoom only on the top half or bottom half of the mirror plot.

7.5.6 Peptide View - Ion Match Table

The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings".

Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

7.5.6.1 Ion Match Table - Library matching table

While in mirror plot view, the Ion Match table updates to display the ion matches between fragment query ions and library ions.
Library intensity and Query intensity are reported relative to the highest intensity ion. All columns can be rearranged and sorted.

### 7.5.7 Peptide View - LC/MS Snapshot

The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The peptide feature is highlighted in purple and the selected PSM marker is blue. The span of the orange line in the LC-MS Snapshot designates the isolation window.

![LC-MS Snapshot](image)

Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button 🤡 in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button 📄 in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking 🎉 to reveal a search option to specify a $m/z$ $RT$ (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

### 7.5.8 Peptide View - Precursor Profile

The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with the retention time along the x-axis and the intensity along the y-axis.
7.5.9 Peptide View - Fragment Ion XIC

An eXtracted Ion Chromatogram (XIC) chart that displays the shape of each identified fragment ion across all scans for the selected PSM. Select the number of ions to display in the chart using the “Show top Fragment Ions” dropdown.
7.6 Spectral Library Search Result - Exporting

In Project View, double-click on 🔄 export to open the Export node.

7.6.1 Spectral Library Search - Exporting - comma separated values (CSV) format

There are several options for exporting the PEAKS Identification results in CSV format:

- **Protein CSV**: The list of protein identifications, filtered by the protein filters in the Summary view and will be saved to proteins.csv.
- **Peptide CSV**: All of the identified peptides and details will be saved to peptides.csv.
- **Support peptides CSV**: A list of supporting peptides of each protein identification, filtered by the protein filters in the "Summary" view and will be exported to protein-peptides.csv. This usually contains more entries than the Peptide table as a peptide can be identified in several proteins and, therefore, can be reported multiple times in this file.
- **PSM CSV**: The peptide-spectrum matches (PSMs) will be exported to DB search psms.csv. Peptides differentiated with only I/L isoform are represented by separate entries. As a result, the number of entries in this file might be bigger than the number of PSMs mentioned in the Summary view.

7.6.2 Spectral Library Search - Exporting - HTML format

**Export in HTML**: This will generate single or multiple HTML reports in the specified location.

The following exporting options are available:

- **Summary view**: The “Summary” view page will be saved as a summary.html file in HTML format in the specified location.
- **Protein coverage**: The coverage pane will be saved for each protein.
- **Supporting peptides**: A list of supporting peptides will be saved for each protein.
- **Best unique PSM**: The best unique PSM will be saved for each protein.
- **Export peptide list in HTML**: creates a separate HTML-containing peptide identifications.

An individual protein will have its own HTML output file, where the corresponding protein coverage, the supporting peptides, and the best unique PSM are gathered.

7.7 Understanding PEAKS DIA Database Search Results

After the DIA database search is complete, DB Search result nodes are generated. Double-click the node to examine the analysis report. The analysis report is presented in three pages:

- **Summary**: This shows an outline of the library search results with key statistics. The overall quality of the experiment can be examined and the filters for peptide and protein identifications can be adjusted.
- Protein: This shows a list of protein identifications. This view also visualizes protein sequence coverage and helps with protein characterization.
- Peptide: This shows a list of peptide identifications. This view also provides spectrum annotation and other detailed information for peptide precursor spectrum matches.

### 7.8 DIA Database Search - Summary View
The DIA DB Search result **Summary View** is similar to the PEAKS Spectral Library Search identification result.

The Summary view reports key statistics as an overview of the result. In the report, several statistical charts can be examined to assess result quality. At the top of each summary result page, protein filters can be set. Identifications that fall below the filters will not be visible in the result pages or exports.

**Proteins -10lgP**: The PEAKS protein score (-10lgP) is calculated as the weighted sum of the -10lgP scores of the protein's supporting peptides. After removing any redundant peptides, the supporting peptides are sorted by -10lgP scores in descending order, and the k-th ranked peptide contributes to the weighted sum with a weight of 1/k. A default threshold of 15 is set.

**Proteins FDR**: The Proteins can also be filtered in accordance to the false-discovery rate of the protein set.

**Proteins Unique Peptides**: The minimum number of unique supporting peptides for a protein identification. A unique peptide is defined as a peptide that can be mapped to only one protein group.

After changing the filters, click **Apply** to apply the new filter and update the results accordingly.

#### 1. Notes

#### 2. Result Statistics

**Figure 1**: False discovery rate (FDR) curve. X axis is the number of peptides being kept. Y axis is the corresponding FDR.

**Figure 2**: RT Calibration

**Figure 3**: Boxplot for median ΔRT across all MS runs
Click on [Notes] to open the Notes dialog to save useful information regarding this Analysis. The Notes will appear in Section 1 of the summary page at the top.

The Summary view reports key statistics as an overview of the result. In the report, several statistical charts can be examined to assess result quality.

### 7.8.1 DIA Database Search - Figure 1. False Discovery Rate (FDR) curve.

**Figure 1.** False discovery rate (FDR) curve. X axis is the number of peptides being kept. Y axis is the corresponding FDR.

![False Discovery Rate (FDR) curve](image)

Figure 1 shows the FDR curve with respect to the number of Peptides (or PSMs) being kept after filtration by the $-10\lg P$ score. By lowering the score threshold, more Peptides are kept in the filtered result. Conversely, the FDR increases because more false positives are kept. In this figure, the vertical line indicates the current score threshold. The number of Peptides and the corresponding FDR by the current score threshold are shown in the top-left corner. PEAKS Search estimates FDR using the "decoy-fusion" approach. Decoy-fusion is an enhanced target-decoy approach that makes more conservative FDR estimations. For details of the "decoy-fusion" approach, please refer to the publication, "PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification", *Mol Cell Proteomics*, 2011 Dec 20.

It is commonly recommended to set FDR at <1% by adjusting the score threshold. If a rapid growth of the FDR around the 1% FDR threshold is noticed, some PSMs may be sacrificed to significantly reduce the FDR.

**Note:** When counting the number of PSMs, PEAKS Studio keeps one peptide per spectrum at most. Thus, the number of PSMs is actually the number of spectra with assigned peptides.

**Note:** During Analysis creation, if Peptide FDR (%) filter was used in Report step, the Summary page FDR curve in the results will show the Peptide FDR curve and corresponding number of Identified peptides.

**Note:** Decoy matches are not counted in the number of PSMs. Unless otherwise specified, decoy matches are also excluded from the other statistical numbers shown in the Summary view.
7.8.2 DIA Database Search result - Figure 2. RT Calibration

Figure 2. RT Calibration

Plots the RT regression function used during the DIA database search. Peptides from the first MS Run in the analysis are plotted against their indexed retention time (iRT) predicted using machine learning. The $R^2$ value is presented based on the regression function.

7.8.3 DIA Database Search result - Boxplot for RT change

Figure 3. Boxplot for median ΔRT across all MS runs

A box plot showing the difference between the observed retention time and iRT peptides within set retention time bins across all fractions, showing the accuracy of retention time prediction.

7.8.4 DIA Database Search result - Table 1. Statistics of data

Statistics of Data tabulates MS1 and MS2 information, as well as identification in the peptide and protein level. The protein level statistics will only be presented if protein inference was performed. This table in the Summary view provides a condensed statistical information for the analysis.

<table>
<thead>
<tr>
<th>Table 1. Statistics of data</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>#MS Run</strong></td>
</tr>
<tr>
<td>---------------</td>
</tr>
<tr>
<td>Total</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Sample 1</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Sample 2</td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>
#MS Run: Total # of runs in each sample.

#MS1: Total # of MS1 spectra for each sample. For timsTOF data this shows the # MS1 frames.

#MS/MS: Total # of MS2 spectra for each sample. This does not include chimeric scans. For timsTOF data this shows the # MS2 frames.

#PSMs: The number of peptide-spectrum matches.

#Peptides: The unique number of peptide sequences with modifications.

#Sequences: The unique number of peptide sequences not including modifications.

#Proteins Groups: PEAKS Studio groups proteins identified by a common set of peptides. This number shows the number of protein groups in the filtered result based on All proteins.

#All Proteins: Total number of proteins with significant peptides, according to current protein filters.

#Top Proteins: Total number of top proteins with significant peptides, according to current protein filters.

#PSM/#MS2: The identification rate by dividing #PSM by #MS2 for the sample.

7.8.5 DIA Database Search result - Result filtration parameters

Tables 2-4 show the statistical numbers of the data and results.

### Table 2. Result filtration parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peptide -10lgP ≤ 0</td>
<td></td>
</tr>
<tr>
<td>Protein -10lgP ≥ 41.5</td>
<td></td>
</tr>
<tr>
<td>Proteins unique peptides ≥</td>
<td>1</td>
</tr>
</tbody>
</table>

### Table 3. Statistics of filtered result

<table>
<thead>
<tr>
<th>FDR</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDR (Peptide-Spectrum Matches)</td>
<td>0.5%</td>
</tr>
<tr>
<td>FDR (Peptide Sequences)</td>
<td>1.0%</td>
</tr>
<tr>
<td>FDR (Protein Group)</td>
<td>100.0%</td>
</tr>
</tbody>
</table>

### Table 4. PTM profile

<table>
<thead>
<tr>
<th>Name</th>
<th>ΔMass</th>
<th>Position</th>
<th>#PSM</th>
<th>#Peptides</th>
<th>-10lgP</th>
<th>Area</th>
<th>AScore</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deamidation</td>
<td>0.98</td>
<td>NQ</td>
<td>20897</td>
<td>5405</td>
<td>51.59</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carboxymethyl</td>
<td>5.07</td>
<td>C</td>
<td>17500</td>
<td>8620</td>
<td>51.30</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oxidation</td>
<td>15.99</td>
<td>M</td>
<td>4787</td>
<td>1713</td>
<td>51.36</td>
<td>9.09E6</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Result filtration parameters: Table 2 summarizes the result filtration parameters set at the top of the Summary View page. This information provides a helpful reference for exported data.

Table 3. Statistics of filtered results: Table 3 presents the statistics based on the data analysis results after setting the desired filters. See below for further explanation:

- **FDR (Peptide-Spectrum Matches):** The total number of decoy database assignments to spectra relative to the total number of target database assignments to spectra represented as a percentage.
- **FDR (Peptide Sequences):** The total number of decoy database assignments to unique peptide sequences relative to the total number of target database assignments to unique peptide sequences represented as a percentage.
- **FDR (Protein Group):** The total number of decoy protein groups where the top hit in the protein group is a decoy database protein relative to the total number of target database protein groups where the top hit is from the target database.

Table 4. PTM Profile: shows modifications identified in the filtered result and the number of PSMs containing each modification.
7.8.6 DIA Database Search result - Experiment Control

Figure 4 in the Summary View show the precursor mass errors of PSMs in filtered results. The precursor mass error is calculated in ppm as $10^6 \times \frac{\text{precursor mass} - \text{peptide mass}}{\text{peptide mass}}$. Calibration is for display purposes only and PEAKS corrects m/z value for each PSM to correct systematic biases and instrument measurement drift.

**Figure 4.** Distribution of precursor mass error of filtered PSM in ppm.

![Distribution of precursor mass error of filtered PSM in ppm.](image)

Table 5 shows the number of identified peptides in each sample with the number of missed cleavages, indicating the enzyme digestion efficiency.

**Table 5.** Number of identified peptides in each sample by the number of missed cleavages.

<table>
<thead>
<tr>
<th>Missed Cleavages</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4+</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>26015</td>
<td>878</td>
<td>32</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>Sample 2</td>
<td>26352</td>
<td>888</td>
<td>31</td>
<td>6</td>
<td>0</td>
</tr>
</tbody>
</table>

7.9 DIA Database Search - Proteins View

The Protein View lists protein identifications that have been filtered by the current settings in Summary View. It also visualizes the protein sequence coverage for identified proteins. The view has four main components:

- **Protein table:** This lists protein identifications that have been filtered by the current settings in the Summary View and the “Protein View Filters”.
- **Coverage:** This visualizes the protein sequence coverage of the selected protein, showing the mapping of support peptides and de novo tags to the protein sequence.
- **Peptides:** This lists supporting peptides mapped to the selected protein.
7.9.1 Proteins View - Protein Table controls
At the top of the Proteins View, there are the options to filter, show/hide columns, search, and export the protein table.

7.9.1.1 Protein Table controls - Protein View Filters
Click on the button to open the Protein View Filter.
The following options are available for filtering the protein table:

**Show Top, All, or First proteins in each group:** Adjust the protein list based on Protein grouping, top is selected by default. Proteins are grouped based on parsimony.

- **Top:** Show only the top proteins in each group. These proteins are supported by the most unique peptides in the group. Proteins in the group that share a subset of the unique peptides that support the top protein (subproteins) will not be displayed.
- **All:** Show all proteins in each group. Proteins are grouped together if they are supported by the same set or a sub-set of the top protein in the group. If a protein is supported by a peptide not supporting the top protein it will be added to a new group.

**Protein accession/name contains:** Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

**Protein description contains:** Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

**Note:** In these above two filters, multiple entries can be separated by a semi-colon.

**Protein sample area:** Filter proteins based on total protein area found in each sample. Sample areas are calculated using the total of all peptide features from unique supporting peptides. If a protein is identified and has areas from multiple samples, only one of the samples must pass this filter to be included.

**PTM contains:** Filter protein results by the presence of selected PTMs. Proteins that are supported by peptides containing modifications that match the selected PTMs will be included. If multiple PTMs match the modification name, they will all be included in the table. For SPIDER results, the mutations will also be listed in the PTM contains filter.

Click on the **Reset** button to remove all filters.

Click on **OK** to apply the Protein View Filter to the Protein table.

Click on **Cancel** to discard changes and close the dialog.
7.9.1.2 Protein Table controls - Protein count
At the top of the protein table shows the count of protein groups and number of proteins in the table based on the current Protein View Filters.

57 protein groups, total 142 proteins

7.9.1.3 Protein Table controls - Optional columns
Click on the button to see a list of sample-based optional columns.

Select the corresponding sample-based Coverage, Spectra count, or Area to add into the Protein Table view.

- **Coverage by Sample**: The protein coverage as a percentage will be displayed for the individual sample when selected.
- **#Spec by Sample**: The total number of spectra identified that support the given protein. Totals are given for each sample included in the search.
- **Area by Sample**: The total area of peptide features from unique supporting peptides in each sample will be displayed.

The exports will always include the contents of these optional columns even if they are hidden.

7.9.1.4 Protein Table controls - Protein Table Search Function
An easy-to-use search function is available for the protein table. The protein accession or description substring can be searched for, and all search results can be navigated through with the Up and Down arrows.

**Protein accession/name contains**: Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.
Protein description contains: Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

7.9.1.5 Protein Table controls - Protein Table Export

Click on the button to export the contents of the current table. This considers the Protein View Filters, the sorting applied to the table and is consistent with the view. There is the option to export the Protein table and/or the Supporting peptide table in CSV format.

7.9.2 Proteins View - Protein Table

Each row in the table represents a group of proteins that are supported by a common set of peptides. To expand the group, click the right arrow button at the left.

The columns in the Protein Table are:

- **Accession**: The accession number of the protein as seen in the FASTA database.
- **Cluster**: A unique identifier that classifies which protein group the protein belongs to.
- **Top**: Indicates whether this protein is a top protein.
- **-10lgP**: The protein confidence score.
- **Coverage (%)**: The percentage of the protein sequence that is covered by supporting peptides. The coverage is visualized by a colored bar. Light blue blocks indicate the parts of the sequence covered by low-confidence peptides. Dark blue blocks indicate the parts covered by high-confidence peptides. This is the total coverage including results from all samples.
- **#Peptides**: The number of high-confidence supporting peptides.
- **#Unique**: The number of high-confidence supporting peptides that are mapped to only one protein group. Unique peptides with same sequence but different modifications are only counted once in this number.
- **PTM**: The identified modifications displayed with color-coded icons.
- **Avg. Mass**: The protein mass calculated using the average mass.
- **Description**: The protein’s header information as seen in the FASTA database.
Note: For #Peptides and #Unique, two peptides with the same starting and ending positions in the protein are counted as one, regardless of their PTM forms. This is to follow the MCP (Molecular & Cellular Proteomics) guidelines.

7.9.3 Proteins View - Coverage Pane

The Protein Coverage view visually maps the supporting peptides and de novo tags to the protein selected in the Protein table. It also shows all identified sites with modifications or mutations to assist with protein characterization at the amino acid level. The "Protein Coverage" view is composed of three major components - Protein Sequence Coverage, Coverage Control Panel, and Protein Tools.

7.9.3.1 Coverage Pane - Protein Sequence Coverage

This area visualizes the coverage of the protein sequence.

Regions in the protein sequence that are covered by supporting peptides are displayed in bold font with a grey background. Confident modifications and mutations identified in supporting peptides are displayed as icons above the protein sequence. Modifications are represented by colored icons with the first letter of its modification name. If a residue is modified by more than one modification in the same supporting peptide, "*" is used instead of a letter. The header information of the protein is shown on the top of the protein sequence.

The supporting peptides can be shown as colored bars under the protein sequence.

Placing the cursor over a bar shows a tooltip with detailed information of the peptide. Left clicking on a bar shows the annotated spectrum from which the supporting peptide is identified.

Right clicking on a peptide bar shows additional options. “Show in spectrum” has the same behaviour as left clicking on the peptide bar which is to open the annotated spectrum chart. “Show in table” has the function to jump to the supporting peptides table.
7.9.3.2 Coverage Pane - Coverage Control Panel

This panel controls what to display in the protein sequence coverage view.

Outline mode: In outline mode, the protein sequence is displayed without supporting peptides and matched de novo tags. However, the sequence coverage at a specific position can be examined by left-clicking a residue.

Coverage mode: In coverage mode, all the supporting peptides and matched de novo tags are shown.

AAs per line and 10AA gap: Specify the number of amino acids per line and whether to show a gap for every 10 amino acids in the protein sequence.

Sequence Display Option: Select whether to view the protein sequence as Fasta or a specific enzyme used in the project. When a particular enzyme is selected, the protein sequence will be highlighted to emphasize expected peptides generated by the selected enzyme's cleavage specificity.
Modification table: The modification table shows the modifications identified in supporting peptides on the protein. For each modification, the number of supporting PSMs with this particular modification is shown. The checkbox on the left controls whether to indicate the modification in the protein coverage.

<table>
<thead>
<tr>
<th>AM</th>
<th>PTM</th>
<th>#</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Carbamidomethylation</td>
<td>40</td>
</tr>
<tr>
<td></td>
<td>Oxidation (M)</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Deamidation (NQ)</td>
<td>4</td>
</tr>
</tbody>
</table>

7.9.3.3 Proteins View - Protein Tools
The Proteins View toolbar is at the upper-right corner of the protein sequence coverage tab and contains helpful protein tools.

Full screen: When selected, the protein coverage window expands in full screen mode. Click to return from full screen mode.

Tool box: The tool box contains the following tools:
- Copy protein sequence. This copies the protein sequence of the current selected protein into the system clipboard.
- Save protein coverage as image. Different scale options are provided to increase the image size.
- Coverage statistics. This provides useful statistics based on the current selected protein.

7.9.3.4 Proteins View - Supporting Peptides
The "Peptides" tab displays a table of supporting peptides for the protein of interest. This table is similar to the peptide table in the Peptide view, except that "Accession" is excluded and the columns "(Checkbox)", "Unique", "Start", and "End" are included. Descriptions of extra columns are given in the following sections.

- (Checkbox): This controls whether a peptide is shown in the protein coverage as a blue bar.
- Unique: This shows whether the peptide is a unique supporting peptide to the protein group.
- Start: This shows the peptide's starting position in the protein.
- End: This shows the peptide's ending position (inclusive) in the protein.
7.10 DIA Database Search Result - Peptide View

The "Peptide View" displays the resulting peptide identifications that have been filtered by the current settings in Summary View and "Peptide View Filters". The table on the top shows the details of each peptide identification. The bottom section provides additional information about the peptide-spectrum matches for the native peptide selected in the table.

---

7.10.1 Peptide View - Peptide Table

The Peptide Table shows the filtered native peptide identification results. Each row in the table is a native peptide identification represented by its highest-scoring PSM. When there are more than 1000 peptides, the table is split into multiple pages.
Note: If there are multiple PSMs matched to a peptide, the table displays the information for the top-scoring PSM. Peptides with the same sequence but different modifications are considered different and shown in different rows.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

The following list describes the contents in each of the default columns:

- **Peptide:** The amino acid sequence of the peptide, as determined by the PEAKS search workflow. A modified residue is followed by a pair of parentheses enclosing the modification.
- **-10lgP:** The peptide -10lgP score. The score indicates the scoring significance of a peptide-precursor spectrum match.
- **Mass:** The monoisotopic mass of the peptide.
- **Length:** The number of amino acids in the sequence backbone.
- **ppm:** The precursor mass error, calculated as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$.
- **m/z:** The precursor mass-to-charge ratio.
- **RT:** The retention time (elution time) of the spectrum as recorded in the data.
- **delta RT:** The difference between the query and library RT.
- **Area per Sample:** The combined area for all features in that sample associated with this peptide.
- **Scan:** The scan number of the spectrum that matches the peptide sequence with the highest -10lgP.
- **#Spec:** number of MS2 spectra assigned to the peptide.
- **Accession:** The accessions of proteins that contain this peptide.
- **PTM:** The types and the numbers of modifications present in the peptide shown in color-coded icons.
- **Found By:** The name of the PEAKS search workflow that identifies the peptide; this can only be Spectral Library or DB Search for this result.

### 7.10.2 Peptide View - Peptide Table controls

The menu above the peptide table provides options to scroll through the table of peptides, filter the table, show optional columns and search for specific entries.
7.10.2.1 Peptide Table controls - Pagination
For easy viewing and to improve performance, the peptide table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.

In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

7.10.2.2 Peptide Table controls - View Filter
Click on the button to open the Peptide View Filter which is used to filter the Peptide table. The following are the filter options provided:

- **Peptide sequence contains**: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains for further filtering if necessary.

- **Peptide score (-10lgP)**: Filters out all peptides with -10lgP scores below this threshold.

- **Peptide length**: Filters for peptides of length within this min/max threshold.

- **Peptide sample area**: Filters based on total peptide area found in each sample. Sample areas are calculated using the total of all peptide features from all spectra that identified the native peptide. If the peptide is found in multiple samples, only one sample must pass this filter for the peptide to be included.

- **PTM contains**: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text they will all be included in the table.

- **Found By**: The dropdown gives the option to show all peptides or only peptides found by the Spectral Library Search or DIA Database Search algorithm.
Note: In all filters, multiple entries can be separated by a semi-colon.

Click on the Reset button to reset all filters to their default setting.

Click on OK to apply the filters to the Peptide table.

Click on Cancel to discard the changes in the View Filter.

### 7.10.2.3 Peptide Table controls - Optional Columns

Click on the [button to open the optional columns dropdown. By default the sample area columns are displayed. Sample Spec count columns can also be shown. These sample spec counts are always from identification, and does not consider whether there are feature vectors present. For projects with many samples, it might be useful to hide these optional columns and reduce the width of the peptide table.

![Show/Hide Columns]

### 7.10.2.4 Peptide Table controls - Search function

On the top right of menu, there is a search function with 4 options. Search results can be navigated between using the Up or Down arrows.

- **Scan :=** Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction.
- **m/z ≈:** Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit. For example, searching for m/z = 560.21 will find all m/z between \([560.20, 560.22]\)
- **RT ≈:** Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/- 1 of the last significant digit set. For example, searching for RT = 55.5 will find all m/z between \([55.4, 55.6]\)
- **Seq contains:** Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered. Additionally, since the peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVIK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”

![m/z](560.2)
7.10.3 Peptide View - Feature selection breadcrumb

There is a sample and feature selection breadcrumb which allows you to switch between features associated with the peptide. Click on the Sample in the breadcrumb to reveal other samples and the features in that sample related to the current peptide selection. From here you can select other samples, fractions, or features.

By clicking on the fraction level, all fractions in this sample are revealed as well as the features in each fraction associated with this peptide. From here you can select other fractions and features to view.

Lastly, clicking on the feature level will show all features in the currently selected fraction. Features with Feature Id $\leq 0$ are known as fake features and are constructed to enhance identification.

7.10.4 Peptide View - PSM selection

Underneath the feature breadcrumb will shown the currently selected PSM. Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.

All other components in the Peptide tab interface will update according to the currently selected PSM.

For diaPASEF instruments, Frame is used rather than Scan number.
7.10.4.1 Peptide View - Protein jump button

Click on \[
\text{Protein jump button}
\]
to jump to a protein that contains the currently selected PSM. If there are more than 1 protein, a dropdown is revealed which allows the user to select which protein to jump to. If a protein is filtered out due to a protein summary filter or summary page protein filter, then the option to jump will be grayed out.

7.10.5 Peptide View - Annotated Spectrum Chart

Click on button \[
\text{Mirror plot view and annotated spectrum chart view}
\]
to switch from the Mirror plot view and annotated spectrum chart view.

For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

7.10.5.1 Annotated Spectrum Chart - Chart Navigation

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.

Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.

Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.

See the whole spectrum: Double click in the spectrum or click the 1:1 button.
7.10.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings

Click on the button to show the spectrum annotation settings.

The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places**: Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation**: Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated**: Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation**: Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info**: Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity**: Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

7.10.5.3 Annotated Spectrum Chart - Additional Chart Controls

Click on or to toggle between absolute and percentage intensity values in the Y axis.

Click on to reset the zoom level of the annotated spectrum chart to the default.
Click on $\times$ to zoom into the X axis by a factor of 2.

Click on $\div$ to zoom into the Y axis by a factor of 2.

The "alignment" check box $\square$ allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The "error map" check box $\square$ is used to show/hide the error map underneath the chart.

7.10.5.4 Annotated Spectrum Chart - Mirror Plot
Spectral Library results have a Mirror Plot view. Click on the $\square$ button to switch to the Mirror Plot.

This view shows the ions from the query spectrum compared to the library spectrum.

Hold down CTRL when scrolling the mouse wheel to zoom only on the top half or bottom half of the mirror plot.

7.10.6 Peptide View - Ion Match Table
The "Ion Match" tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings".

Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.
7.10.6.1 Ion Match Table - Library matching table
While in mirror plot view, the Ion Match table updates to display the ion matches between fragment query ions and library ions.

Library intensity and Query intensity are reported relative to the highest intensity ion. All columns can be rearranged and sorted.

7.10.7 Peptide View - LC/MS Snapshot
The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The peptide feature is highlighted in purple and the selected PSM marker is blue. The span of the orange line in the LC-MS Snapshot designates the isolation window.

Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a m/z RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

7.10.8 Peptide View - Precursor Profile
The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with the retention time along the x-axis and the intensity along the y-axis.
An eXtracted Ion Chromatogram (XIC) chart that displays the shape of each identified fragment ion across all scans for the selected PSM. Select the number of ions to display in the chart using the “Show top Fragment Ions” dropdown.
7.11 DIA Database Search Result - Exporting

In Project View, double-click on to open the Export node.

7.11.1 DIA Database Search - Exporting - comma separated values (CSV) format

There are several options for exporting the PEAKS Identification results in CSV format:

- **Protein CSV**: The list of protein identifications, filtered by the protein filters in the Summary view and will be saved to proteins.csv.
- **Peptide CSV**: All of the identified peptides and details will be saved to peptides.csv.
- **Support peptides CSV**: A list of supporting peptides of each protein identification, filtered by the protein filters in the "Summary" view and will be exported to protein-peptides.csv. This usually contains more entries than the Peptide table as a peptide can be identified in several proteins and, therefore, can be reported multiple times in this file.
- **PSM CSV**: The peptide-spectrum matches (PSMs) will be exported to DB search psm.csv. Peptides differentiated with only I/L isoform are represented by separate entries. As a result, the number of entries in this file might be bigger than the number of PSMs mentioned in the Summary view.

7.11.2 DIA Database Search - Exporting - HTML format

**Export in HTML**: This will generate single or multiple HTML reports in the specified location.

The following exporting options are available:

- **Summary view**: The “Summary” view page will be saved as a summary.html file in HTML format in the specified location.
- **Protein coverage**: The coverage pane will be saved for each protein.
- **Supporting peptides**: A list of supporting peptides will be saved for each protein.
- **Best unique PSM**: The best unique PSM will be saved for each protein.
- **Export peptide list in HTML**: creates a separate HTML-containing peptide identifications.

An individual protein will have its own HTML output file, where the corresponding protein coverage, the supporting peptides, and the best unique PSM are gathered.

7.11.3 DIA Database Search - Exporting - Spectral Library generation

**Spectral Library TSV**: PEAKS DIA Database Search results can be exported as a spectral library that can be used by the Library Search engine. Select “Use original RT” to keep original RT values in the library. Select “Map to iRT” to save iRT values in the library. The map to iRT function maps experimental retention times for peptides to an indexed RT (iRT) to account for chromatographic differences between runs (e.g. different gradient lengths).

For more information on how to configure a PEAKS generated spectral library, refer to Section 2.9 Spectral Library Configuration.
7.12 Understanding PEAKS DIA De Novo Results

After the DIA de novo sequencing is complete, De Novo result nodes are generated. Double-click the node to examine the analysis report. The analysis report is presented in two pages:

- **Summary**: This shows an outline of the De Novo results with key statistics. The overall quality of the experiment can be examined and the filters for de novo ALC can be adjusted.
- **De novo**: This shows a list of peptide sequences that pass the ALC threshold detected by de novo sequencing.

The de novo view displays the de novo sequencing results. The table on the top displays de novo sequences, while the bottom section provides additional information about the peptide-spectrum match. The results can be filtered and exported.

7.13 DIA De Novo - Summary View

The DIA De Novo **Summary View** is similar to the PEAKS Deep Novo sequencing result.

In the summary tab, a PEAKS Studio DIA de novo sequencing result can be filtered using Average Local Confidence (ALC) score. Low quality deep novo sequences can be filtered out by specifying a minimal threshold of the ALC score. The purpose of filtering is to remove poor sequences in which residues are incorrectly identified. By default, the ALC threshold is set to 50%. After changing the threshold, click **Apply** to apply the new filter. The result in the De novo result view will be updated accordingly.

### 1. Notes

### 2. Result Statistics

**Table 1. Statistics of data and result.**

<table>
<thead>
<tr>
<th>Sample Name</th>
<th># MS2</th>
<th># PSM</th>
<th># PSM (qk &gt; 50)</th>
<th># PSM (qk &gt; 50)</th>
<th># PSM (qk &gt; 70)</th>
<th># PSM / # MS2</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>190460</td>
<td>235829</td>
<td>187169</td>
<td>97363</td>
<td>30827</td>
<td>128.8%</td>
</tr>
<tr>
<td>Sample 1</td>
<td>98238</td>
<td>129486</td>
<td>96385</td>
<td>10069</td>
<td>10640</td>
<td>132.0%</td>
</tr>
<tr>
<td>Sample 2</td>
<td>98238</td>
<td>123383</td>
<td>91184</td>
<td>47262</td>
<td>14767</td>
<td>125.0%</td>
</tr>
</tbody>
</table>

**Table 2. Result filtration parameters.**

De novo score (%): ≥ 50%

### 3. Other Information

**Table 3. Search parameters.**
- **Search Engine Name**: PEAKS DENOVO
- **Parent Mass Error Tolerance**: 10.0 ppm
- **Fragment Mass Error Tolerance**: 0.02 Da
- **Enzyme**: Specified by each sample
- **Fixed Modifications**: Carbamidomethylation (+57.02)
- **Variable Modifications**: Deamidation (Q) (<7.98)
- **Oxidation (M) (<15.99)
- **Max Variable PTM Per Peptide**: 2

Click on **Notes** to open the Notes dialog to save useful information regarding this Analysis. The Notes will appear in Section 1 of the summary page at the top.
Section 2 Result Statistics gives key statistics in Table 1 on the number of MS2 scans and # PSMs total as well as with different ALC thresholds for each sample. The #PSM / #MS2 provides a de novo sequencing identification rate metric for each sample.

Table 2 simply shows what is the current De novo ALC filter set. This is placed here so that when the Summary page is exported, the viewer will know what the ALC threshold was at the time.

Section 3 Other Information shows the parameters that were set when the analysis was ran.

7.14 DIA De Novo result - De Novo View

The De novo View displays the DIA de novo sequencing results in greater detail, as shown in the next figure. The table on the top displays de novo sequences, while the bottom section provides additional information about the peptide-spectrum match.

<table>
<thead>
<tr>
<th>Scan</th>
<th>Peptide</th>
<th>ALC/CO</th>
<th>Length</th>
<th>r/a</th>
<th>RT</th>
<th>Area</th>
<th>Mass</th>
<th>ppp</th>
<th>FTY</th>
<th>Mode</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ABCDE</td>
<td>5040</td>
<td>1000</td>
<td>2</td>
<td>0.5</td>
<td>50</td>
<td>500</td>
<td>0.5</td>
<td>0.5</td>
<td>HCD</td>
</tr>
<tr>
<td>2</td>
<td>FGHJ</td>
<td>6040</td>
<td>1000</td>
<td>2</td>
<td>0.5</td>
<td>50</td>
<td>500</td>
<td>0.5</td>
<td>0.5</td>
<td>HCD</td>
</tr>
</tbody>
</table>

7.14.1 De Novo View - De novo table

The following list describes the contents in each of the columns of the Denovo table:

- **Scan**: The scan number and data file it belongs to. The scan number is a unique index for tandem mass spectra in the data.
- **Peptide**: The amino acid sequence of the peptide as determined by de novo sequencing. A modified residue is followed by a pair of parentheses enclosing the mass of that modification.
• **ALC(%):** The average local confidence. ALC is calculated as the total of the residue local confidence scores in the peptide divided by the peptide length. The higher the score the more confident the de novo candidate.

• **Frame:** This column is only displayed for diaPASEF data. A unique identifier is assigned for each frame obtained from the raw data.

• **Length:** The number of amino acids in the sequence backbone.

• **m/z:** The precursor mass-to-charge ratio.

• **z:** The precursor charge.

• **RT:** The retention time (elution time) of the spectrum as recorded in the data.

• **1/k0 Range:** This shows the ion mobility range associated with precursor. This column is only shown if the analysis contains diaPASEF data.

• **Area:** The area under the curve of the peptide feature found at the same m/z and retention time as the MS/MS scan. This can be used as an indicator of the abundance of the peptide.

• **Mass:** The monoisotopic mass of the peptide.

• **ppm:** The precursor mass error, calculated as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$.

• **PTM:** The types and the numbers of modifications present in the peptide shown in color-coded icons.

• **Mode:** The fragmentation mode in which de novo sequencing is performed by the algorithm.

Click on any of the column headers to sort the column.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

In the table, Amino acids in deep novo sequences are color-coded according to their local confidence scores. Red represents a very high confidence (greater than 90%); purple represents a high confidence (80 to 90%); blue represents a medium confidence (60 to 80%); and black represents low confidence (less than 60%). Hover the cursor over the sequence to view local confidence scores of each individual amino acid.

### 7.14.2 De Novo View - De Novo table controls

The menu above the De Novo table provides options to scroll through the table of deep novo peptides, filter for mass tags, switch between samples, search for specific entries, and export the table.

#### 7.14.2.1 De Novo table controls - Sample Selection

Result for different samples can be viewed by selecting the sample using the "Show Results for Specified Sample" drop-down menu.

#### 7.14.2.2 De Novo table controls - Pagination

1000 rows are visible at any one time in the De Novo table. Use the top left Pagination options to switch between result records.
7.14.2.3 De Novo table controls - Mass Tag Filter

Click on \( \pm 0\% \) to use the mass tag confidence slider filter which will replace amino acids in the sequence with their mass tag values if their local confidence does not meet the threshold.

7.14.2.4 De Novo table controls - Search function

Use the search function to search for scan # or precursor ID, m/z, RT, or sequence contains. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- **Scan**: Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction. For timsTOF, there is the option to search for Precursor ID instead.

- **m/z**: Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit.
  
  For example, searching for m/z = 580.8 will find all m/z between [580.7, 580.9]

- **RT**: Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/- 1 of the last significant digit set.
  
  For example, searching for RT = 64 will find all m/z between [63, 65]

- **Seq contains**: Search for the sequence in the table. If specifying a sequence without glycans or PTMs, the results will include that sequence and any sequences with glycans or PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered.

  Additionally, since the glycan peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”

7.14.2.5 De Novo table controls - Table exporting

Click on the \( \pm 0\% \) button to export the contents of the current table. This considers sorting applied to the table and is consistent with the view. There is the option to export the De Novo table and/or the De novo all candidates table in CSV format.
7.14.3 De Novo View - Deep Novo All Candidates

Click on the button to view the All de novo candidates for that spectrum. By default, the top 5 candidates are reported. The highest ALC scoring candidate is the one displayed in the De novo table.

### All de novo Candidates

<table>
<thead>
<tr>
<th>Peptide</th>
<th>ALC(%)</th>
<th>Length</th>
<th>ppm</th>
<th>Mode</th>
</tr>
</thead>
<tbody>
<tr>
<td>ETH:Q(+0.08)LLC(−37.02)K</td>
<td>99.9</td>
<td>9</td>
<td>1.3</td>
<td>HCD</td>
</tr>
<tr>
<td>Q(+0.08)TVHEL(−37.02)K</td>
<td>99.9</td>
<td>9</td>
<td>1.3</td>
<td>HCD</td>
</tr>
<tr>
<td>TEYHEL(−37.02)K</td>
<td>81.5</td>
<td>9</td>
<td>1.3</td>
<td>HCD</td>
</tr>
<tr>
<td>TV(+0.08)YHEL(−37.00)K</td>
<td>80.5</td>
<td>9</td>
<td>1.3</td>
<td>HCD</td>
</tr>
</tbody>
</table>

7.14.4 PSM selection

Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.

All other components in the De novo View interface will update according to the currently selected PSM.

For diaPASEF instruments, Frame is used rather than Scan number.

#### 7.14.4.1 Show spectrum in Data View

Click on the button to open the Data Refine MS/MS tab (if not already opened) and jump to the corresponding selected Scan.

#### 7.14.4.2 Show spectrum in LC/MS View

Click on the button to open the Data Refine LC/MS tab (if not already opened) and jump to the corresponding selected Scan, highlighting the De novo marker and the feature if present.

#### 7.14.4.3 Show Raw Spectrum View

Click on the button to show the raw spectrum chart of the currently selected Scan.

7.14.5 Annotated Spectrum Chart

Click on the button to switch from the Mirror plot view and annotated spectrum chart view.
For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

7.14.5.1 Annotated Spectrum Chart - Chart Navigation
Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

- **Zoom to a m/z region**: Drag horizontally from the start m/z and to the end m/z with left mouse button.
- **Zoom in/out smoothly**: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.
- **Increase/Decrease peak intensity**: Place the cursor in the spectrum and scroll the mouse wheel.
- **See the whole spectrum**: Double click in the spectrum or click the 1:1 button.

7.14.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings
Click on the button to show the spectrum annotation settings.
The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places:** Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation:** Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated:** Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation:** Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info:** Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity:** Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

### 7.14.5.3 Annotated Spectrum Chart - Additional Chart Controls

Click on ⬇️ or ⬆️ to toggle between absolute and percentage intensity values in the Y axis.

Click on ⬇️ to reset the zoom level of the annotated spectrum chart to the default.

Click on ⬇️ to zoom into the X axis by a factor of 2.

Click on ⬇️ to zoom into the Y axis by a factor of 2.
The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

7.14.5.4 Annotated Spectrum Chart - Mirror Plot
Deep Novo results have a Mirror Plot view. Click on the button to switch to the Mirror Plot.

This view shows the ions from the query spectrum compared to the predicted spectrum.

Hold down CTRL when scrolling the mouse wheel to zoom only on the top half or bottom half of the mirror plot.

7.14.5.5 Annotated Spectrum Chart - Pseudo Scan
Click on the button to show the Pseudo Scan.

7.14.6 DIA De Novo Ion Match Table
The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table
can be configured in "Spectrum Annotation Settings" (see 7.14.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings).

<table>
<thead>
<tr>
<th>#</th>
<th>b</th>
<th>b-H2O</th>
<th>b-NH3</th>
<th>b(2+)-</th>
<th>Seq</th>
<th>y</th>
<th>y-H2O</th>
<th>y-NH3</th>
<th>y(2+)-</th>
<th>#</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>101.05</td>
<td>112.04</td>
<td>113.02</td>
<td>65.55</td>
<td>E</td>
<td>908.54</td>
<td>931.52</td>
<td>991.50</td>
<td>502.37</td>
<td>9</td>
</tr>
<tr>
<td>2</td>
<td>231.10</td>
<td>213.09</td>
<td>214.07</td>
<td>116.05</td>
<td>T</td>
<td>899.49</td>
<td>930.48</td>
<td>991.45</td>
<td>489.75</td>
<td>8</td>
</tr>
<tr>
<td>3</td>
<td>330.17</td>
<td>312.16</td>
<td>313.14</td>
<td>165.58</td>
<td>V</td>
<td>898.49</td>
<td>930.48</td>
<td>991.45</td>
<td>489.75</td>
<td>7</td>
</tr>
<tr>
<td>4</td>
<td>407.23</td>
<td>389.22</td>
<td>390.20</td>
<td>234.13</td>
<td>H</td>
<td>799.42</td>
<td>831.41</td>
<td>891.39</td>
<td>400.31</td>
<td>6</td>
</tr>
<tr>
<td>5</td>
<td>596.27</td>
<td>578.26</td>
<td>579.24</td>
<td>288.63</td>
<td>E</td>
<td>663.36</td>
<td>694.35</td>
<td>745.33</td>
<td>331.68</td>
<td>5</td>
</tr>
<tr>
<td>6</td>
<td>709.36</td>
<td>691.35</td>
<td>692.33</td>
<td>335.18</td>
<td>L</td>
<td>533.32</td>
<td>565.30</td>
<td>616.28</td>
<td>267.15</td>
<td>4</td>
</tr>
<tr>
<td>7</td>
<td>822.44</td>
<td>804.43</td>
<td>805.41</td>
<td>411.72</td>
<td>L</td>
<td>420.23</td>
<td>452.22</td>
<td>502.21</td>
<td>210.61</td>
<td>3</td>
</tr>
<tr>
<td>8</td>
<td>982.47</td>
<td>964.46</td>
<td>965.45</td>
<td>491.73</td>
<td>C(+17.02)</td>
<td>307.15</td>
<td>339.13</td>
<td>390.12</td>
<td>154.07</td>
<td>2</td>
</tr>
<tr>
<td>9</td>
<td>1229.51</td>
<td>1211.50</td>
<td>1212.49</td>
<td>631.62</td>
<td>K</td>
<td>147.11</td>
<td>129.10</td>
<td>120.09</td>
<td>74.06</td>
<td>1</td>
</tr>
</tbody>
</table>

Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

7.14.6.1 Ion Match Table - Predicted matching table
While in mirror plot view, the Ion Match table updates to display the ion matches between fragment query ions and predicted ions.

<table>
<thead>
<tr>
<th>#</th>
<th>Label</th>
<th>Predicted M/Z</th>
<th>Predicted Intensity (%)</th>
<th>Query M/Z</th>
<th>Query Intensity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>b2</td>
<td>258.15</td>
<td>0.67</td>
<td>258.14</td>
<td>4.088 (58.85%)</td>
</tr>
<tr>
<td>2</td>
<td>y2</td>
<td>310.17</td>
<td>0.38</td>
<td>310.17</td>
<td>2.022 (5.91%)</td>
</tr>
<tr>
<td>3</td>
<td>b3</td>
<td>345.18</td>
<td>0.29</td>
<td>345.17</td>
<td>5.344 (7.55%)</td>
</tr>
<tr>
<td>4</td>
<td>y3</td>
<td>409.24</td>
<td>0.13</td>
<td>409.24</td>
<td>1.512 (2.72%)</td>
</tr>
<tr>
<td>5</td>
<td>b4</td>
<td>508.24</td>
<td>0.79</td>
<td>508.24</td>
<td>3.238 (34.66%)</td>
</tr>
<tr>
<td>6</td>
<td>y4</td>
<td>572.31</td>
<td>8.81</td>
<td>572.36</td>
<td>3.583 (53.31%)</td>
</tr>
<tr>
<td>7</td>
<td>b5</td>
<td>595.27</td>
<td>12.40</td>
<td>595.27</td>
<td>7.17E (16.35%)</td>
</tr>
<tr>
<td>8</td>
<td>y5</td>
<td>671.37</td>
<td>14.42</td>
<td>671.37</td>
<td>8.888 (21.14%)</td>
</tr>
<tr>
<td>9</td>
<td>b6</td>
<td>694.34</td>
<td>55.14</td>
<td>694.33</td>
<td>2.81E (40.55%)</td>
</tr>
<tr>
<td>10</td>
<td>y6</td>
<td>758.41</td>
<td>41.06</td>
<td>758.40</td>
<td>2.888 (61.56%)</td>
</tr>
<tr>
<td>11</td>
<td>b7</td>
<td>857.40</td>
<td>63.27</td>
<td>857.40</td>
<td>4.37E (63.06%)</td>
</tr>
</tbody>
</table>

Predicted intensity and Query intensity are reported relative to the highest intensity ion. All columns can be rearranged and sorted.

7.14.7 DIA De Novo View - LC/MS Snapshot
The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The span of the orange line in the LC-MS Snapshot designates the isolation window.
Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a m/z RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

7.14.8 DIA De Novo View - Precursor Profile

The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with the retention time along the x-axis and the intensity along the y-axis. The first two isotopes are also drawn as M+1 and M+2 respectively.
7.14.9 DIA De Novo View - Fragment Ion XIC

An eXtracted Ion Chromatogram (XIC) chart that displays the shape of each identified fragment ion across all scans for the selected PSM. Select the number of ions to display in the chart using the “Show top Fragment Ions” dropdown.

7.15 DIA De Novo result - Exporting

In Project View, double-click on Export to open the Export node.

There are several options for exporting the Deep Novo results:

- **De novo Summary HTML**: Exports the Summary page including all figures in HTML format. The content will be the same as shown in PEAKS Studio.
- **De novo CSV**: Exports the De Novo table. This export does not consider any sorting and will export a separate file for each sample.
- **De novo CSV with all Candidates**: Exports the De novo all candidates. This export does not consider any sorting and will export a separate file for each sample.
- **De novo Features CSV**: Exports the features list with de novo sequencing results. Each sample will have its own export file.
- **Spectral Library TSV**: PEAKS DIA De Novo results can be exported as a spectral library that can be used by the Library Search engine.
  For more information on how to configure a PEAKS generated spectral library, refer to Section 2.9 Spectral Library Configuration.
7.16 DIA Label Free Quantification result

The DIA LFQ result is very similar to the following chapter 8. Label Free Quantification (LFQ)

Label free quantification can be used as a quick profiling tool to examine the relative abundance of proteins in large proteomic datasets. In PEAKS Studio this quantification method is based on the relative abundance of peptide features detected in multiple samples. PEAKS Studio performs feature detection separately on each sample, and uses an EM (expectation-maximization) algorithm to detects and deconvolute overlapped features. Importantly, features of the same peptide from different samples are reliably aligned together (i.e. matched between runs) using a high performance retention time alignment algorithm to improve overall protein quantification rates.

**Note:** For details of the retention alignment algorithm, refer to the paper "A Combinatorial Approach to the Peptide Feature Matching Problem for Label-Free Quantification", Bioinformatics, 2013, 10.1093.

To generate a DIA LFQ result, follow the steps in above section 7.1.4 DIA Quantification (DIA LFQ) Analysis Workflow and Parameter Settings

7.17 Understanding DIA LFQ Results

Once completed, a label free quantification result node will be added to the project tree. Double-click on this \( \text{LFQ} \) node to open the LFQ result that contains multiple tabs as described below:

- **Summary:** The outline of the PEAKS LFQ search results with statistics. This is the place to examine the overall performance of the experiment and adjust filters.
- **Proteins:** The quantified proteins with a list of supporting peptide features for each protein.
- **Peptides:** The quantified peptides listed in a table. Extracted ion chromatogram (XIC), sample features and retention time alignment also displayed for each peptide.
1. Notes

2. Result Statistics

**Figure 1:** Protein profile heatmap (Cell colour represents the log2(ratio) to the average abundance across different samples)

---

### 7.18.1 Summary View - Protein Filters

Filter settings used in the screenshot above means that no protein filter is applied, thus giving all quantifiable protein.

Alternately, parameters can be adjusted for additional analysis by updating the protein filters above.

**Significance:** Only protein groups with a significance above this threshold will be listed in the Protein View. The significance score is calculated as the $-10\log_{10}$ of the significance testing p-value. PEAKS provides ANOVA significance testing methods A significance score threshold of 20 is recommended, which equals to a significance testing p value of 0.01. Either this or “FDR (adjusted p-value)” can be selected to set a significance threshold.

**Fold Change:** Only proteins at or above this fold change threshold will be listed in the Protein View.

**Used Peptides:** Only proteins with at least this many number of used peptides will be listed in the Protein View.

Click on [Apply] to apply these filters changes to the result. This will update charts on the Summary View and will update the Protein table accordingly.

### 7.18.2 Summary View - Notes

Click on [Notes] to open the Notes dialog and save information about the result and saved in section 1. Notes on the Summary page.
1. Notes

Example 1: Notes
The heat map displays the protein groups that passed the filters. The relative protein abundance is represented as a heat map of the representative proteins of each protein group. These representative proteins are clustered if they exhibit a similar expression trend across the samples. The hierarchical clustering is generated using a neighbour-joining algorithm with a Euclidean distance similarity measurement of the log2 ratios of the abundance of each sample relative to the average abundance. Similarly, the conditions in different samples are clustered if they exhibit a similar expression trend across the protein groups. An interactive protein profile heat map is opened in a new window when the button is clicked options.

Mouse over the dendogram in a section of the heatmap and click on it to open the Variation Trend chart.
The Variation Trend chart displays the expression level of the selected proteins in the samples.

Mouse over points on the variation trend chart to see a tooltip of the protein, sample, and log2ratio.

Click on the Export Data button to export the variation trend chart data in a text file format. When Displayed is selected, the log2 ratios will be in the export. When Original is selected, the original ratio values will be in the export instead.
To view the group ratio distribution of the quantified proteins, click the Density-Ratio button. By default, all proteins will be shown. Click Define Species to group the different species together.

The proteins can be searched from the list by typing into the search box, marked and then added as a species group. The expected ratio of the species in the "Ratio" column can be edited to match with the experiment design. Once all the species are added and organized, click Ok to finish. In the plot window, click Update to apply the changes.
7.18.5 Summary View - Sample Correlation plot

To view the abundance correlation plot for the detected features and proteins, click the Sample Correlation button from the Summary View title bar. A Correlation Plot Options window will pop-up to select which samples to include.

By default, the first 5 samples are selected. At most 5 samples can be used to generate the correlation plot, and at least 2 samples must be selected.
The Protein Correlation and Peptide Correlation plots will be displayed for the samples selected. The Pearson’s correlation score indicates the reproducibility of the experiment based on a selected pair of samples. To export the image right click on the plots and click on “Export Image”.

7.18.6 Summary View - Group Correlation plot

To view the abundance correlation plot for the detected features and proteins, click the button from the Summary View title bar. A Correlation Plot Options window will pop-up to select which groups to include.

By default, the first 5 groups are selected. At most 5 groups can be used to generate the correlation plot, and at least 2 groups must be selected.
The Protein Correlation and Peptide Correlation plots will be displayed for the samples selected. The Pearson’s correlation score indicates the reproducibility of the experiment based on a selected pair of samples. To export the image right click on the plots and click on “Export Image”.

7.18.7 Summary View - Volcano plot

Figure 2. volcano plot combines a statistical test with the magnitude of the change enabling quick visual identification of those data-points that display large-magnitude changes and that are also statistically significant. The volcano plot in Figure 2(a) plots significance versus fold-change of the quantified proteins.

Horizontal broken grey line: The selected significance threshold

Vertical broken grey lines: The selected fold change thresholds

Plotting points in this way results in two regions of interest in the plot: those points that are found towards the top of the plot that are far to either the left or the right side. These represent values that display large magnitude fold changes (hence being left or right of center) as well as high statistical significance (hence being towards the top).
In this figure, the ratio is the group ratio set with respect to the base group. If more than two groups are present, the max ratio is used instead.

The Protein View contains an interactive volcano plot that can be hidden or displayed. Markers for the proteins that are above the set significance threshold will be displayed in colour. Scrolling the cursor over a marker will highlight it and, subsequently, clicking on this marker will highlight the protein in the interface.
7.18.8 Summary View - RT and m/z Shift Distribution
Figure 3(a) plots the retention time shift distribution to the base sample. Figure 3(b) plots the distribution of the m/z shift (in ppm) of the data with respect to the base sample. These two figures can be used to refine the parameter settings for label free quantification.

Figure 3. (a) RT shift distribution, (b) M/Z shift distribution.

7.18.9 Summary View - Missing Value percentage chart
Figure 4 shows the percentage of missing values before and after ID transfer for each sample.

Figure 4. Percentage of missing values in each sample with or without ID transfer.
7.18.10 Summary View - Filtration parameters and statistics
Tables 1 and 2 list the filtration parameters and filtered results, respectively.

**Table 1. Result filtration parameters.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Significance</td>
<td>≥ 0</td>
</tr>
<tr>
<td>Min Fold Change</td>
<td>≥ 1</td>
</tr>
<tr>
<td>Max Fold Change</td>
<td>≤ 64</td>
</tr>
<tr>
<td>Used peptides</td>
<td>≥ 1</td>
</tr>
</tbody>
</table>

**Table 2. Statistics of filtered result.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Features</td>
<td>203883</td>
</tr>
<tr>
<td>Features with ID</td>
<td>177325</td>
</tr>
<tr>
<td>Feature vectors with ID</td>
<td>67961</td>
</tr>
<tr>
<td>Protein groups</td>
<td>5633</td>
</tr>
</tbody>
</table>

7.18.11 Summary View - Search parameters
Table 3 lists the parameters as set in the workflow.

**Table 3. Search Parameters**

Search Engine Name: PEAKS
Quantification type: Label free quantification
LFQ method: Identification directed quantification
Retention time shift tolerance: Auto detected
Feature intensity ≥ 100.0
Retention time range: [0.0 - Max]
Base sample: Sample 1
Peptide Feature Filter:
  - Avg area ≥ 200.0
  - Quality ≥ 20.0
  - Charge between: [2 - 5]
  - Peptide ID count ≥ 0 per group
  - Detected in at least: 1 samples per group
Protein Filter:
  - Significance method: ANOVA
  - Modified Form Exclusion
  - Remove Outlier
  - Use Top 3 peptide
  - Significance ≥ 0.0
  - Fold change between: [1.0 - 64.0]
  - Has at least: 1 used peptide
Normalization method: Use TIC
Samples: 3 samples in 3 groups
  - Group 1:
    - Sample 1
  - Group 2:
    - Sample 2
  - Group 3:
    - Sample 3
7.19 DIA LFQ - Protein View

The **Proteins View** lists all the quantified proteins present in the sample, characterizes each protein at the amino acid level, and lists the supporting peptide features of each protein. It has six components:

**Protein Table**: Lists all the quantified proteins present in the sample.

**Protein View Filters**: Allows filtering the protein table to select specific proteins.

**Protein Volcano Plot**: Click on a marker that is above the significance threshold (the horizontal dashed line) set in the Summary View. Upon selection, the corresponding protein in the Protein Table will be highlighted.

**PTM Profiling**: Click on to open PTM profiling.

**Coverage**: Characterizes the protein sequence at the amino acid level. All the PTMs occurred on the protein sequence will be displayed together with the MS/MS data supporting the inference.

**Peptides**: Lists the supporting peptides identified from the selected proteins in the protein table.

7.19.1 Proteins View - Protein Table controls

At the top of the Proteins View, there are the options to filter, show/hide columns, search, and export the protein table.

![Protein View Filter](image)

**141 protein groups, total 160 proteins**

7.19.1.1 Protein Table controls - Protein View Filters

Click on the **Protein View Filter** button to open the Protein View Filter.
The following options are available for filtering the protein table:

**Show All or Top proteins in each group:** Adjust the protein list based on Protein grouping, top is selected by default. Proteins are grouped based on parsimony.

- **Top:** Show only the top proteins in each group. These proteins are supported by the most unique peptides in the group. Proteins in the group that share a subset of the unique peptides that support the top protein (subproteins) will not be displayed.
- **All:** Show all proteins in each group. Proteins are grouped together if they are supported by the same set or a sub-set of the top protein in the group. If a protein is supported by a peptide not supporting the top protein it will be added to a new group.

**Protein accession/name contains:** Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

**Protein description contains:** Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

**Note:** In these above two filters, multiple entries can be separated by a semi-colon.

**PTM contains:** Filter protein results by the presence of selected PTMs. Proteins that are supported by peptides containing modifications that match the selected PTMs will be included. If multiple PTMs match the modification name, they will all be included in the table. For SPIDER results, the mutations will also be listed in the PTM contains filter.

Click on the **Reset** button to remove all filters.

Click on **OK** to apply the Protein View Filter to the Protein table.

Click on **Cancel** to discard changes and close the dialog.
7.19.1.2 Protein Table controls - Protein count
At the top of the protein table shows the count of protein groups and number of proteins in the table based on the current Protein View Filters.

57 protein groups, total 142 proteins

7.19.1.3 Protein Table controls - Optional columns
Click on the button to see a list of sample-based optional columns.

Select the corresponding sample-based and group-based ratio in the Protein Table view.

The exports will always include the contents of these optional columns even if they are hidden.

7.19.1.4 Protein Table controls - Protein Table Search Function
An easy-to-use search function is available for the protein table. The protein accession or description substring can be searched for, and all search results can be navigated through with the Up and Down arrows.

**Protein accession/name contains:** Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

**Protein description contains:** Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

7.19.1.5 Protein Table controls - Protein Table Export
Click on the button to export the contents of the current table. This considers the Protein View Filters, the sorting applied to the table and is consistent with the view. There is the option to export the Protein table and/or the Supporting peptide table in CSV format.
7.19.2 Proteins View - Protein Table

Each row in the table represents a group of proteins that are supported by a common set of peptides. To expand the group, click the right arrow button at the left.

The columns in the Protein Table are:

- **Accession**: The accession number of the protein as seen in the FASTA database.
- **Cluster**: A unique identifier that classifies which protein group the protein belongs to.
- **Top**: Indicates whether this protein is a top protein.
- **Significance**: The protein significance.
- **Coverage (%)**: The percentage of the protein sequence that is covered by supporting peptides. The coverage is visualized by a colored bar. Light blue blocks indicate the parts of the sequence covered by low-confidence peptides. Dark blue blocks indicate the parts covered by high-confidence peptides. This is the total coverage including results from all samples.
- **#Peptides**: The number of high-confidence supporting peptides.
- **#Unique**: The number of high-confidence supporting peptides that are mapped to only one protein group. Unique peptides with same sequence but different modifications are only counted once in this number.
- **PTM**: The identified modifications displayed with color-coded icons.
- **Sample Profile**: The protein abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the sample channels, abundances, and ratios with respect to the base sample.
- **Group Profile**: The protein abundance among the groups is depicted as a heat map. The Group Profile is determined by calculated the total abundance (area) of supporting peptides within a group. Hold the cursor on a profile to view the group channels, abundances, and ratios compared to the group that contains the base sample.
- **Avg. Mass**: The protein mass calculated using the average mass.
- **Description**: The protein's header information as seen in the FASTA database.

**Note**: For #Peptides and #Unique, two peptides with the same starting and ending positions in the protein are counted as one, regardless of their PTM forms. This is to follow the MCP (Molecular & Cellular Proteomics) guidelines.
7.19.3 Proteins View - Coverage Pane

The Protein Coverage view visually maps the supporting peptides and de novo tags to the protein selected in the Protein table. It also shows all identified sites with modifications or mutations to assist with protein characterization at the amino acid level. The "Protein Coverage" view is composed of three major components - Protein Sequence Coverage, Coverage Control Panel, and Protein Tools.

7.19.3.1 Protein Sequence Coverage

This area visualizes the coverage of the protein sequence.

Regions in the protein sequence that are covered by supporting peptides are displayed in bold font with a grey background. Confident modifications identified in supporting peptides are displayed as icons above the protein sequence. Modifications are represented by colored icons with the first letter of its modification name. If a residue is modified by more than one modification in the same supporting peptide, "*" is used instead of a letter. The header information of the protein is shown on the top of the protein sequence.

The supporting peptides and matched de novo tags can be shown as colored bars under the protein sequence.

Placing the cursor over a bar shows a tooltip with detailed information of the peptide.
Left clicking on a bar shows the details from which the supporting peptide is identified.

Right clicking on a peptide bar shows additional options. “Show in spectrum” has the same behaviour as left clicking on the peptide bar which is to open the annotated spectrum chart. “Show in table” has the function to jump to the supporting peptides table.

7.19.3.2 Coverage Control Panel
This panel controls what to display in the protein sequence coverage view.
Sequence Display Mode: The protein sequence can be displayed in either "outline" or "coverage" mode.

Outline mode: In outline mode, the protein sequence is displayed without supporting peptides and matched de novo tags. However, the sequence coverage at a specific position can be examined by left-clicking a residue.

Coverage mode: In coverage mode, all the supporting peptides and matched de novo tags are shown.

AAs per line and 10AA gap: Specify the number of amino acids per line and whether to show a gap for every 10 amino acids in the protein sequence.

Sequence Display Option: Select whether to view the protein sequence as Fasta or a specific enzyme used in the project. When a particular enzyme is selected, the protein sequence will be highlighted to emphasize expected peptides generated by the selected enzyme's cleavage specificity.
Modification minimal ion intensity / Ascore: A modification or mutation is considered confident if there are fragment ions supporting the modified residue with the minimal peak intensity above the specified threshold. Only confident modifications and mutations are displayed as icons above the protein sequence.

- Show confident modification site
  - Minimal ion intensity
    - Threshold
    - Percentage
  - Minimal Ascore
    - Threshold

Modification table: The modification table shows the modifications identified in supporting peptides on the protein. For each modification, the number of supporting PSMs with this particular modification is shown. The checkbox on the left controls whether to indicate the modification in the protein coverage.

<table>
<thead>
<tr>
<th>AA</th>
<th>PTM</th>
<th>#</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>Carbamidomethyl... 63</td>
<td>3</td>
</tr>
<tr>
<td>D</td>
<td>Deamidation (N)</td>
<td>7</td>
</tr>
<tr>
<td>O</td>
<td>Oxidation (M)</td>
<td>5</td>
</tr>
<tr>
<td>T</td>
<td>2-amino-3-oxo-butyric...</td>
<td>3</td>
</tr>
<tr>
<td>F</td>
<td>Formylation</td>
<td>3</td>
</tr>
<tr>
<td>K</td>
<td>Acetylation (Protein...)</td>
<td>2</td>
</tr>
<tr>
<td>S</td>
<td>Sodium adduct</td>
<td>2</td>
</tr>
<tr>
<td>A</td>
<td>Amidation</td>
<td>1</td>
</tr>
</tbody>
</table>

7.19.3.3 Proteins View - Protein Tools
The Proteins View toolbar is at the upper-right corner of the protein sequence coverage tab and contains helpful protein tools.

- Full screen: When selected, the protein coverage window expands in full screen mode. Click to return from full screen mode.

- Tool box: The tool box contains the following tools:
  - Copy protein sequence. This copies the protein sequence of the current selected protein into the system clipboard.
  - Save protein coverage as image. Different scale options are provided to increase the image size.
  - Coverage statistics. This provides useful statistics based on the current selected protein.

7.19.3.4 Proteins View - Supporting Peptides
The supporting peptides assigned to the protein are shown in a separate tab beside the Protein Coverage view. The sequence can be clicked and a pop-up window will show up to display the feature details.
This table contains the following information of the quantifiable support peptides for the selected protein:

- **Peptide:** The amino acid sequence of the peptide, as determined in PEAKS Database or Spectral Library search. A modified residue is followed by a pair of parentheses enclosing the modification mass. All non-unique peptides and those not passing the filter will be excluded automatically.

- **Used:** Shows whether this peptide is used to calculate the protein abundance. The top three peptides with highest abundance are used to calculate the protein abundance.

- **Quality:** The feature vector quality score.

- **Significance:** The -10\(\log p\)-value represents the likelihood that the observed change between conditions is caused by random chance. The peptides are first separated into groups based on similar quality score. Significance is then calculated for each quality score group, or bin, using a two-tail T-test that assumes log normal distribution but does not assume equal variance. The -10\(\log p\)-value is then calculated from this result and then displayed in the significance column.

- **Avg. ppm:** A peptide can have one or more feature vectors. Each feature vector has its own ppm calculated from the mass error of each feature. The average mass error of a peptide is the average of mass errors of all the feature vectors. This column displays the average mass error in ppm.

- **Avg. Area:** Each feature in the feature vector has its own area under the XIC curve. The average area of a feature vector is the average area of all the features associated with the peptide.

- **Sample Profile:** The peptide abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the samples, the peptide areas in samples, and the corresponding ratios with respect to the base sample.

- **Group Profile:** The peptide abundance among the groups is depicted as a heat map. Hold the cursor on a profile to view the groups, the peptide areas in groups, and the corresponding ratios with respect to the group which contains the base sample.

- **Group Area:** This column shows the peptide area for a group. It is calculated by summing up the areas of feature vectors within a group.

- **Max Ratio:** This shows the maximum of the fold change values compared to the reference sample.

- **RT Mean:** Each feature in the feature vector has its own retention time center. This column displays the average retention time center of the feature vector.

- **#Vector:** This shows the number of quantifiable feature vectors of a peptide.

- **Start:** This shows the protein position of the first residue of the peptide.

- **End:** This shows the protein position of the last residue (inclusive) of the peptide.

- **PTM:** The types and the numbers of modifications present in the peptide shown using color-coded icons. Scroll over each icon to see the modification name and mass.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Used</th>
<th>Quality</th>
<th>Significance</th>
<th>Avg. ppm</th>
<th>Avg. Area</th>
<th>Sample Profile</th>
<th>Group Profile</th>
<th>Group Area</th>
<th>Max Ratio</th>
<th>RT Mean</th>
<th>#Vector</th>
<th>Start</th>
<th>End</th>
<th>PTM</th>
</tr>
</thead>
<tbody>
<tr>
<td>DILYDGDK</td>
<td>3.46</td>
<td>0.00</td>
<td>0.7</td>
<td>4.71</td>
<td>7.30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>KARQDK</td>
<td>3.46</td>
<td>0.00</td>
<td>0.7</td>
<td>4.71</td>
<td>7.30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DEYIK</td>
<td>3.46</td>
<td>0.00</td>
<td>0.7</td>
<td>4.71</td>
<td>7.30</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IDK</td>
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<td>0.00</td>
<td>0.7</td>
<td>4.71</td>
<td>7.30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IDK</td>
<td>3.46</td>
<td>0.00</td>
<td>0.7</td>
<td>4.71</td>
<td>7.30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IDK</td>
<td>3.46</td>
<td>0.00</td>
<td>0.7</td>
<td>4.71</td>
<td>7.30</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IDK</td>
<td>3.46</td>
<td>0.00</td>
<td>0.7</td>
<td>4.71</td>
<td>7.30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IDK</td>
<td>3.46</td>
<td>0.00</td>
<td>0.7</td>
<td>4.71</td>
<td>7.30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IDK</td>
<td>3.46</td>
<td>0.00</td>
<td>0.7</td>
<td>4.71</td>
<td>7.30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
- **Feature Detail**: Double click on a supporting peptide to go to the peptide page where the eXtracted Ion Chromatogram (XIC) curves are displayed (as seen below).

### 7.20 DIA LFQ result - Peptides View

Quantified proteins are displayed in the Peptide View along with their detailed information.

#### Peptide Table

The peptides are grouped together based on quantifiable feature vectors with the same sequence. When there are more than 1000 peptides, the table is split into multiple pages.

#### Peptide Table Entries

The view is similar to the support peptide table described in the previous section Section 7.19.3.4 Proteins View - Supporting Peptides. The Peptide table entries may be filtered by using the Peptide View Filters on top of the peptide table. To reiterate, the columns are:
- **Peptide**: The amino acid sequence of the peptide, as determined in PEAKS Database or Spectral Library search. A modified residue is followed by a pair of parentheses enclosing the modification mass. All non-unique peptides and those not passing the filter will be excluded automatically.

- **Avg. Area**: Each feature in the feature vector has its own area under the XIC curve. The average area of a feature vector is the average area of all the features associated with the peptide.

- **Quality**: The feature vector quality score.

- **Significance**: The -10lg of a p-value represents the likelihood that the observed change between conditions is caused by random chance. The peptides are first separated into groups based on similar quality score. Significance is then calculated for each quality score group, or bin, using a two-tail T-test that assumes log normal distribution but does not assume equal variance. The -10log p-value is then calculated from this result and then displayed in the significance column.

- **Avg. ppm**: A peptide can have one or more feature vectors. Each feature vector has its own ppm calculated from the mass error of each feature. The average mass error of a peptide is the average of mass errors of all the feature vectors. This column displays the average mass error in ppm.

- **Sample Profile**: The peptide abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the samples, the peptide areas in samples, and the corresponding ratios with respect to the base sample.

- **Group Profile**: The peptide abundance among the groups is depicted as a heat map. Hold the cursor on a profile to view the groups, the peptide areas in groups, and the corresponding ratios with respect to the group which contains the base sample.

- **Group Area**: This column shows the peptide area for a group. It is calculated by summing up the areas of feature vectors within a group

- **Max Ratio**: This shows the maximum of the fold change values compared to the reference sample.

- **RT Mean**: Each feature in the feature vector has its own retention time center. This column displays the average retention time center of the feature vector.

- **#Vector**: This shows the number of quantifiable feature vectors of a peptide.

- **Accession**: The proteins that the peptide was found in.

- **PTM**: The types and the numbers of modifications present in the peptide shown using color-coded icons. Scroll over each icon to see the modification name and mass.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

### 7.20.2 Peptide Table controls

The menu above the peptide table provides options to scroll through the table of peptides, filter the table, show optional columns and search for specific entries.

### 7.20.2.1 Peptide Table controls - Pagination

For easy viewing and to improve performance, the peptide table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.
In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

**7.20.2.2 Peptide Table controls - View Filter**

Click on the **Peptide View Filter** button to open the Peptide View Filter which is used to filter the Peptide table. The following are the filter options provided:

- **Peptide sequence contains**: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains for further filtering if necessary.

- **Significance >=**: Only peptides that pass this Significance threshold will remain in the table.

- **Fold Change >=**: Peptides with Max Ratio below this threshold will be removed from the table.

- **PTM contains**: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text they will all be included in the table.

  **Note**: In all filters, multiple entries can be separated by a semi-colon.

Click on the **Reset** button to reset all filters to their default setting.

Click on **OK** to apply the filters to the Peptides table.

Click on **Cancel** to discard the changes in the View Filter.
7.20.2.3 Peptide Table controls - Optional Columns

Click on the button to open the optional columns dropdown. The sample ratios and group ratios can be displayed in the peptide table for easy view and sorting. For projects with many samples, it might be useful to hide these optional columns and reduce the width of the peptide table.

7.20.2.4 Peptide Table controls - Search function

On the top right of menu, there is a search function with 2 options. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- \( m/z \approx \): Search for the approximate \( m/z \). The search will consider a range of \( m/z \) equal to the search value +/- 1 of the last significant digit. For example, searching for \( m/z \approx 560.21 \) will find all \( m/z \) between \([560.20, 560.22]\).

- **Seq contains**: Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered. Additionally, since the peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”.

7.20.3 Feature vector selection

Click on the Feature Vector dropdown to open a dropdown to display the feature vectors that correspond to the selected peptide. The peptide -10lgP score is the highest among the -10lgP scores of its feature vectors. The peptide quality score corresponds to the average of the quality scores of all feature vectors.
7.20.4 Protein jump button

Click on the protein jump button to jump to a protein that contains the currently selected PSM. If there are more than 1 protein, a dropdown is revealed which allows the user to select which protein to jump to. If a protein is filtered out due to a protein summary filter or summary page protein filter, then the option to jump will be grayed out.

7.20.5 Feature Details

The Feature Details contains the eXtracted Ion Chromatogram (XIC) chart. The eXtracted Ion Chromatogram (XIC) chart displays the shape of the selected peptide feature vector over the retention time range associated with the identification. The table on the left displays both the area and the ratio of the features in the highest average abundance feature vector. This table also works as a control to select the features that will be used to draw the XIC plot. By default, the first sample in each group is selected.

Right click on the chart area to export the XIC charts.

7.21 DIA LFQ Exporting

In Project View, double-click on the export button to open the Export node.
7.21.1 LFQ result - Exporting - comma separated values (CSV) format

There are several options for exporting the PEAKS Label Free Quantification results in CSV format:

- **Proteins CSV**: The list of protein quantifications, filtered by the protein filters in the Summary view and will be saved to proteins.csv.
- **Peptide CSV**: A complete list of all quantified peptides that pass the peptide feature filters set on the "Summary" page and their associated details.
- **Support peptides CSV**: These are the supporting peptides of the quantified proteins and their associated details.
- **Feature Vector CSV**: All predicted peptide features and their associated quantification information. Both identified and unidentified peptide features are included. To include unidentified peptide features, set the Peptide Id Count equal to 0 in the Peptide feature filter.

7.21.2 LFQ result - Exporting - HTML format

**Export in HTML**: This will generate single or multiple HTML reports in the specified location.

The following exporting options are available:

- **Summary view**: The “Summary” view page will be saved as a summary.html file in HTML format in the specified location.
- **Protein coverage**: The coverage pane will be saved for each protein.
- **Supporting peptides**: A list of supporting peptides will be saved for each protein.
- **Best unique PSM**: The best unique PSM will be saved for each protein.
- **Export peptide list in html**: creates a separate HTML-containing peptide identifications.

An individual protein will have its own HTML output file, where the corresponding protein coverage, the supporting peptides, and the best unique PSM are gathered.

7.22 DIA Quality Control (QC)

The QC function was added in PEAKS Studio 11 update. In the DIA Identification and Quantification workflow, it can be included as an optional step to produce an automated QC result node for sophisticated QC analysis on the data level and protein/peptide identification and quantification. The function is used in both DDA and DIA data.
Double-click on "QC" in the analysis to open the QC result node.

**QC Summary**

Table 1. Statistics of filtered results.

<table>
<thead>
<tr>
<th></th>
<th>Count</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selected Samples</td>
<td>5</td>
</tr>
<tr>
<td>Passed Samples Count</td>
<td>2</td>
</tr>
<tr>
<td>Ratio of Passed Samples</td>
<td>40%</td>
</tr>
<tr>
<td>Failed Samples Count</td>
<td>3</td>
</tr>
<tr>
<td>Ratio of Failed Samples</td>
<td>60%</td>
</tr>
<tr>
<td>Total Failed Attributes</td>
<td>5</td>
</tr>
</tbody>
</table>

Figure 1. Failed Attributes Chart.

Table 2. Statistics of LFQ Result

<table>
<thead>
<tr>
<th>Sample Name</th>
<th># MS1</th>
<th># MS2</th>
<th># PSM</th>
<th># Identified Features</th>
<th># Quantified Features</th>
<th># Peptides</th>
<th># Quantified Peptides</th>
<th># Protein Groups</th>
<th># Quantified Protein Groups</th>
<th># PSM / Peptides</th>
<th>ID Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>1644</td>
<td>1640</td>
<td>223,584</td>
<td>228,988</td>
<td>143,052</td>
<td>10113</td>
<td>2383</td>
<td>1945</td>
<td>3.08</td>
<td>53.76%</td>
<td></td>
</tr>
<tr>
<td>Average of Passed Samples</td>
<td>1643.5</td>
<td>1641.5</td>
<td>219,283</td>
<td>220,277</td>
<td>149,527</td>
<td>203,511</td>
<td>1945</td>
<td>2687</td>
<td>2.83</td>
<td>52.63%</td>
<td></td>
</tr>
<tr>
<td>Ecol03_0ffgel_black</td>
<td>2738</td>
<td>3940</td>
<td>238,700</td>
<td>216,400</td>
<td>9393</td>
<td>9022</td>
<td>1480</td>
<td>1847</td>
<td>3.54</td>
<td>94.88%</td>
<td></td>
</tr>
<tr>
<td>Ecol03_0ffgel_green</td>
<td>492</td>
<td>2840</td>
<td>217,844</td>
<td>164,313</td>
<td>12,263</td>
<td>8223</td>
<td>2694</td>
<td>1844</td>
<td>2.38</td>
<td>94.84%</td>
<td></td>
</tr>
<tr>
<td>Ecol03_0ffgel_red</td>
<td>4884</td>
<td>1878</td>
<td>211,178</td>
<td>209,611</td>
<td>19970</td>
<td>9682</td>
<td>2177</td>
<td>1939</td>
<td>2.71</td>
<td>94.84%</td>
<td></td>
</tr>
</tbody>
</table>

Figure 2. Error Distribution. (a) MS1 Mass Error Distribution. (b) MS2 Mass Error Distribution.

7.22.1 DIA Quality Control (QC) - Summary

The QC Summary page gives an overview of the passed and failed samples with the selected attributes.

Click on "Summary Filter" to open the QC summary filter which provides an option to choose samples and attributes to check whether they pass the acceptance tolerance with respect to the Control sample.
To adjust the control sample or acceptance tolerance, another analysis will need to be run. Simply click on Modify Analysis and adjust those two settings in the QC step, leaving other parameters the same.

**Figure 1. Failed Attributes Chart.**

<table>
<thead>
<tr>
<th>Table 1. Statistics of filtered results.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selected Samples Count</td>
</tr>
<tr>
<td>Passed Samples Count</td>
</tr>
<tr>
<td>Ratio of Passed Samples</td>
</tr>
<tr>
<td>Failed Samples Count</td>
</tr>
<tr>
<td>Ratio of Failed Samples</td>
</tr>
<tr>
<td>Total Failed Attributes</td>
</tr>
</tbody>
</table>

Table 1 provides a quick glance on the number of passed and failed samples from the selected samples.

Figure 1 shows a histogram of the failed attributes and the number of samples with that failed attribute.
Table 2. Statistics of LFQ Result

<table>
<thead>
<tr>
<th>Sample Name</th>
<th># MS1</th>
<th># MS2</th>
<th># PSM</th>
<th># Identified Precursors</th>
<th># Quantified Precursors</th>
<th># Peptides</th>
<th># Quantified Peptides</th>
<th># Protein Groups</th>
<th># Quantified Protein Groups</th>
<th>ID Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control Sample</td>
<td>5628</td>
<td>45017</td>
<td>80317</td>
<td>94075</td>
<td>28031</td>
<td>74758</td>
<td>27302</td>
<td>8408</td>
<td>4121</td>
<td>178.41%</td>
</tr>
<tr>
<td>Average of Passed Samples</td>
<td>5627</td>
<td>45016</td>
<td>80227</td>
<td>92469</td>
<td>28032</td>
<td>74376</td>
<td>27285</td>
<td>8491</td>
<td>4125</td>
<td>178.22%</td>
</tr>
<tr>
<td>Sample 4</td>
<td>5628</td>
<td>45016</td>
<td>79007</td>
<td>89830</td>
<td>28004</td>
<td>73264</td>
<td>27259</td>
<td>8288</td>
<td>4121</td>
<td>175.51%</td>
</tr>
<tr>
<td>Sample 5</td>
<td>5628</td>
<td>45016</td>
<td>78723</td>
<td>87116</td>
<td>27917</td>
<td>70454</td>
<td>27180</td>
<td>8013</td>
<td>4113</td>
<td>168.55%</td>
</tr>
<tr>
<td>Sample 6</td>
<td>5627</td>
<td>45016</td>
<td>73942</td>
<td>84902</td>
<td>27857</td>
<td>68764</td>
<td>27126</td>
<td>7871</td>
<td>4126</td>
<td>164.26%</td>
</tr>
<tr>
<td>Sample 7</td>
<td>5626</td>
<td>45000</td>
<td>81239</td>
<td>93302</td>
<td>27815</td>
<td>75476</td>
<td>27076</td>
<td>8076</td>
<td>4115</td>
<td>180.53%</td>
</tr>
</tbody>
</table>

Table 2 provides more details on which particular samples have failed attributes, and the average values for those passed samples. Samples with failed attributes will be listed as separate rows below the Average of Passed Samples. The columns in this table were selected as attributes of interest, so they will not align with all attributes that were selected in the Summary filter.

Figure 2. Error Distribution. (a) MS1 Mass Error Distribution. (b) MS2 Mass Error Distribution.

Figure 2 shows how the MS and MS/MS distribution compares to the control sample.

Table 3. QC Parameters.

<table>
<thead>
<tr>
<th>Control Sample: Sample 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acceptance Tolerance: 10.0%</td>
</tr>
<tr>
<td>Total Sample Count: 7</td>
</tr>
<tr>
<td>QC Attributes:</td>
</tr>
<tr>
<td># Identified Precursors</td>
</tr>
<tr>
<td># Peptides</td>
</tr>
<tr>
<td>Quantified Peptides</td>
</tr>
</tbody>
</table>

Table 3 simply shows the QC parameters that were set in this analysis. The Standard Peptides list here can be highlighted and copied, allowing for easy re-use of the standard peptides in another analysis.
The Data QC pertains to quality control measures related to the data itself. The following columns are in the table:

- **Sample Name**: Name of each sample.
- **All Attributes**: Pass/Fail flag. If a sample fails ANY attribute (not just in Data QC), it will be marked as Fail.
- **#MS1**: The number of MS scans in the last fraction in each sample.
- **#MS2**: The number of MS/MS scans in the last fraction in each sample.
- **# Detected Precursor**: The number of detected precursors in each sample.
- **Full Width (min)**: Measures the width of the median eluted peptide feature for each sample.
- **FWHM (sec)**: Abbreviation for Full Width at Half Maximum. This measures the number of seconds at half the maximum abundance of the median eluted peptide feature for each sample.
- **BPC**: Shows the BPC intensity of each sample.
- **TIC Correlation**: The Pearson correlation coefficient with respect to points on the control sample TIC.

The table has two selection controls. When selecting a cell, the column and row are highlighted. The highlighted column will update the line plot, showing the distribution between the control sample and all other samples.
The highlighted row is for comparing the control sample and the sample in the selected row. A Sample TIC chart is provided to examine how the TIC curve compares. Click on the button to change the view so that the curves overlap. The MS2 Count over Retention Time chart compares the cumulative MS/MS count as Retention time increases.

Right-click on the charts to export the image at any time.

7.22.3 DIA Quality Control (QC) - Identification QC
Similar to Data QC, the Identification QC tab also provides a table with statistics of interest and relevant charts from Identification results. QC will use a combination of the Spectral Library search result and the DIA Database Search result, depending on whether one or both of them were ran in the analysis.
The table contains the following columns pertaining to the DIA Identification result:

- **Sample Name**: Name of each sample.
- **All Attributes**: Pass/Fail flag. If a sample fails ANY attribute (not just in Identification QC), it will be marked as Fail.
- **# MS2**: The number of MS2 scans in each sample.
- **# Identified Precursor**: The total number of precursors with identified spectra.
- **# PSM**: Total number of PSMs identified in the sample.
- **# Peptide**: Total number of distinct peptides identified in the sample.
- **# Sequence**: Total number of distinct peptide backbones identified in the sample.
- **# Protein Groups**: Total number of protein groups identified in the sample.
- **# Top Protein**: Total number of top proteins identified in the sample.
- **# All Protein**: Total number of proteins identified in the sample.
- **Missed Cleavage Ratio**: The ratio of identified peptides with missed cleavages versus peptides without any missed cleavages.
- **Enzyme Specificity Ratio**: The ratio of identified peptides with specific digest versus peptides without.
- **# PSM / # Peptide Rate**: The ratio of # PSM and # Peptide for each sample.
- **MS1 Error Mean (SD)**: Shows the MS1 error mean in ppm and the standard deviation
- **MS2 Error Mean (SD)**: Shows the MS/MS error mean in Da and the standard deviation
- **ID Rate (%)**: Calculated as percent of #PSM / # MS2

Row and Column selection works the same as Data QC, where the selected Column will update the line chart to show the distribution of the column for each sample, and the selected Row will update the charts according to how they compare with the control sample. The charts include a venn diagram for protein and peptide overlap, a #PSM/#Peptide distribution, and the MS1 and MS2 error distributions violin plots.
7.22.4 DIA Quality Control (QC) - Quantification QC

For DIA Quantification workflow only, the Quantification QC tab provides a table with statistics of interest and relevant charts from DIA LFQ results.

The table contains the following columns pertaining to the Quantification result:

- **Sample Name**: Name of each sample.
- **All Attributes**: Pass/Fail flag. If a sample fails ANY attribute (not just in Quantification QC), it will be marked as Fail.
- **# Identified Precursor**: The total number of features with identified spectra. Same number as in Identification QC.
- **# Quantified Precursor**: The total number of features used for feature vectors from the sample.
- **Missing Value (%)**: Displays the missing value after ID Transfer for each sample. This is the same number as in the LFQ result.
- **Precursor Correlation**: Measures the pearson correlation coefficient of the precursors compared to the control sample. This is the same value as the precursor correlation in the LFQ result sample correlation pop-up.
- **# Identified Peptide**: Total number of distinct peptides identified in the sample. This is the same number as in Identification QC. When selecting this column, the # Quantified Peptide column is also selected.
- **# Quantified Peptide**: Total number of distinct peptides quantified in the sample. When selecting this column, the # Identified peptide column is also selected.
- **# Identified Protein Group**: Total number of protein groups identified in the sample. This is the same number as in Identification QC. When selecting this column, the # Quantified protein group column is also selected.
- **# Quantified Protein Group**: Total number of protein groups quantified in the sample. When selecting this column, the # Identified protein group column is also selected.

Row and Column selection works the same as Data QC, where the selected Column will update the line chart to show the distribution of the column for each sample, and the selected Row will update the charts according to how they compare with the control sample. The distribution chart for Peptides and Protein groups are always selected together and displayed on the same chart. A correlation plot between top protein, peptide, and feature is drawn for the selected sample and control sample. An RT Alignment chart is provided to show the retention time alignment between selected sample and control sample.

Right-click on the charts to export the image at any time.

### 7.22.5 DIA Quality Control (QC) - Standard Peptides
When standard peptides were added to QC function, an additional tab is in the QC result node. A table will show which peptides are found in each sample, and the RT and m/z for those standard peptides can be viewed for quick comparison between samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>EcoID:00000000</th>
<th>EcoID:00000001</th>
<th>EcoID:00000002</th>
<th>EcoID:00000003</th>
<th>EcoID:00000004</th>
<th>EcoID:00000005</th>
<th>EcoID:00000006</th>
<th>EcoID:00000007</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALAQQKQVTEHMDQIQ</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>GRAVYTTEPKDGKE</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>GAVAVNGTEK</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>ITLNAYYKMK</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>LILPPGR</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>LIPPPER</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>TILKHPODQYQV</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>TLYLEINR</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>VQDGAVTFQIGK</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Each sample and the standard peptides are listed in the table. The table can be sorted, but the control sample will always be the first index. Click on the columns to select/deselect that standard peptide for viewing in the RT and
M/Z reproducibility charts. Multiple columns can be selected, and if none are selected then all standard peptides will be displayed. For those standard peptides of interest, there are checkboxes on the right side of the charts which can be toggled to show missing values. Those samples that are missing will have a vertical bar drawn, so that it is easier to tell which samples are missing the standard peptides.

Right-click on the charts to export the image at any time.

8.7.6 Identification Quality Control (QC) - QC Result Exports
As mentioned, all images in the QC result node can be exported by right-clicking on the chart, and selecting Export image.

The Export Chart pop-up will appear, and the image can be exported at different scales.

Besides image exporting, the tables can be exported in CSV files and each QC view can be exported in HTML format. Double-click the node to find the QC Export options.
8. Label Free Quantification (LFQ)
Label free quantification can be used as a quick profiling tool to examine the relative abundance of proteins in large proteomic datasets. In PEAKS Studio this quantification method is based on the relative abundance of peptide features detected in multiple samples. PEAKS Studio performs feature detection separately on each sample, and uses an EM (expectation-maximization) algorithm to detect and deconvolute overlapped features. Importantly, features of the same peptide from different samples are reliably aligned together (i.e. matched between runs) using a high performance retention time alignment algorithm to improve overall protein quantification rates.

Note: For details of the retention alignment algorithm, refer to the paper "A Combinatorial Approach to the Peptide Feature Matching Problem for Label-Free Quantification", Bioinformatics, 2013, 10.1093.

8.1 Label Free Quantification Analysis Workflow and Settings
Label Free Quantification requires at least 2 samples in the analysis. It is recommended to only run LFQ with 1 data file per sample, but if you choose to have fractionated samples, ensure each sample has the same number of fractions.

Select the PEAKS Q workflow in the Workflow Selection step

The following steps are the same as in the PEAKS DB workflow (as in section 6.1 Database Search Workflow and Parameter Settings) and is reiterated below.

Click on Data Refine to proceed through the workflow.

After deciding the Data Refine option, proceed to the next step to see the DB Search parameters.
The following parameter settings are provided:

**Precursor Mass Error Tolerance:** The maximum mass difference between the peptide and the monoisotopic mass of the precursor ion.

**Precursor Mass Tolerance unit:** PEAKS generates peptides within the precursor mass error tolerance. Precursor mass error tolerance can be specified in either Daltons or ppm.

**Fragment Mass Error Tolerance:** PEAKS uses this value when scoring peptide sequences. PEAKS Studio considers a fragment ion to be matched if the calculated m/z is within the specified error tolerance. Units are Daltons.

Both precursor and fragment error tolerance parameters need to be set consistently with the mass accuracy of the instrument.

**Enzyme Settings:** PEAKS Studio digests the protein database *in silico* to generate peptide candidates. It is necessary to specify the enzyme for protein digestion from the Enzyme drop-down menu.

**Note:** When the selected dataset is digested with different enzymes, "Specified by each sample" allows samples to be analyzed separately using their respective enzymes specified during project creation.
Note: "None" is a special enzyme allowing non-specific cleavage at both ends of the peptide. It is the recommended option when no digestion enzyme was used or when the digestion enzyme exhibits a high degree of non-specificity.

Missed Cleavages: This specifies the number of missed cleavage sites allowed in a peptide.

Digest Mode: This specifies the type of in silico cleavages allowed at a peptide termini based on the method used for protein digestion. If "Specific" is selected, the specificity of the selected enzyme is strictly enforced at both termini of a peptide. "Semispecific" allows one terminus to disobey the enzyme specificity rules.

Note: If the enzyme is specified as "None", then no matter which mode is selected for "Digest Mode", the "None" enzyme digest rule will be applied.

PTM Settings: Click “Set PTM” to open the PTM Options dialog and specify the fixed PTMs and a few common variable PTMs expected in the experiment. Highlight PTMs and click on the button to place them as either Fixed or Variable PTM. From there, highlighted PTMs can be removed by clicking the “Remove” button. Clicking “Remove All” will reset the PTM options. Highlighting and clicking “Switch Type” will move the Fixed PTM in into the Variable PTM list and vice-versa. Click on “New” to create a new custom PTM. While a PTM is selected, click on “View” to see more details.

Note: Multiple PTMs can be highlighted by holding down the CTRL key or SHIFT key on the keyboard.

Note: For a specific residue, only one fixed modification is allowed.

Once the PTMs are selected, the maximum number of variable PTMs per peptide can be specified and a value of less than 4 is recommended.

Note: The use of variable modifications increases the size of the computational search space for the de novo sequencing algorithm. It is recommended not to select too many variable modifications in PEAKS de novo. See PEAKS PTM for alternate search options.

Target Database: Select a protein sequence database for the search. Select one from the list of databases that have been configured in PEAKS and set the taxonomy by clicking on , if applicable. To configure a new database, click on next to the Target Database dropdown to open the Database Configuration dialog.

Contaminant Database: To search for contaminants, select a contaminant database from the list of databases that have been configured in PEAKS. Keep the selection as N/A if no contaminant database is required for the search.

Peptide Length: The length for a peptide can be set when creating an analysis, where the default peptide length range is between 6 and 45.

Deep Learning Boost: Check this option to use PEAKS enhanced deep learning algorithm to better identify PSMs in the DB Search process.

Click on Quantification to proceed to the next step and select the Label Free option.
The following parameters are available:

**Sample Group:** LFQ requires sample grouping. All available samples are listed in the left window. After selecting a sample, it can be added to a new group by clicking the button or to an existing group by clicking the button and selecting that option from the drop-down menu. If you select multiple files, you can add each one individually to create multiple new groups by clicking the button. Samples and Groups can be removed by clicking the button next to each element. The names and colours for each group can be edited by clicking on the coloured square dropdown, respectively.

**Mass Error Tolerance:** The mass shift of a precursor ion between different runs. This can be set similar to the precursor mass error tolerance used for database search.

**Retention Time Shift Tolerance:** The retention time shift tolerance is the maximum elution time difference that is considered for the quantification of an identified peptide after RT alignment.
**Auto Detect:** Let PEAKS Studio detect the optimal Retention Time Shift Tolerance for your data based on the observed trend.

**Ion Mobility Tolerance (1/k0) (if applicable):** The maximum 1/k0 differential for features to transfer IDs.

*Note:* This parameter is only available when the analysis contains timsTOF data.

**Feature intensity:** Set a value for the minimum intensity a peptide feature must exhibit.

**RT Range:** Feature vectors within the selected retention time range will be included in the results while everything else will be removed.

**Base Sample:** The base sample will act as the reference. The peptide and protein abundance corresponding to this reference label will be the denominator in all of the abundance ratio calculations.

Click on the tool icon next to Peptide Feature Filters to open the Feature filter dialog.

**Avg. Area:** The minimum average area of all peptide features associated with the peptide. Features with higher abundance have been shown to be more reproducible across replicates. Only peptide features with an average area above this threshold will be used for quantification.

**Quality:** The quality score for the peptide feature. Factors that affect the quality score include m/z difference, RT difference, XIC shape similarity, and the feature intensities.

**Peptide ID Count:** This filter sets the minimum limit for how many times a peptide must be identified within a group to be included in the LFQ results.
**Charge between:** Only peptides with feature vectors that all fall within this range will be used in protein quantification.

**Have at least ___ confident samples per group:** This filter sets the minimum number of sample(s) per group that the peptide feature has an abundance value. A peptide is more quantifiable when it is detected in more samples.

**Use in group coefficient of variation filter:** For each feature vector (set of peptide features associated with a peptide), at least one group must have a coefficient of variation (CV) less than the CV threshold. The CV threshold is automatically selected by the software based on the current data set.

Click on the tool icon next to Protein to open the Protein filter dialog

**Significance Method:** Protein abundance is used for the significance testing calculation. When there are replicates in each group, ANOVA is suggested to be used for significance testing. For more details on ANOVA, refer to the following paper: "On the comparison of several mean values: an alternative approach ", Biometrika, 1951, 38(3/4): 330-336

**Modified Form Exclusion:** The expression level of modified peptides might be different than the overall expression level of a protein. In such cases, including modified peptides for protein ratio calculation might lead to incorrect results. By checking this box, if a peptide has modified and unmodified forms, both will not be used as candidates for protein ratio calculation, if a peptide only has one modified form, it is still a candidate, but if a peptide has more than one modified form, all forms will not be used as candidates.

**Remove Outliers:** This filter removes supporting peptides that have a different variation trend compared to others.
**Use Top # peptides:** This is the number of quantifiable supporting peptide that will be used to calculate the protein profile. Three is the default setting.

**Significance:** Only protein groups with a significance above this threshold will be listed in the “Protein” view. The significance score is calculated as the -10log of the significance testing p-value (e.g. Significance score of 20 is equivalent to a p-value of 0.05). PEAKS provides the significance testing method: ANOVA.

**FDR:** Adopt the Benjamin-Hochberg method to adjust the p-value to the false discovery rate for all protein groups that have already passed the other filters. Only protein groups with significance scores passing the calculated FDR will be listed in the "Protein" view. Either this or "Significance" can be selected to set a significance threshold.

**Fold Change:** Only protein groups above the lower threshold or below the upper threshold value will be listed in the "Protein" view.

**Used Peptides:** Only protein groups that were quantified with this number of peptides (or more) will be listed in the "Protein" view.

Click on the tool icon next to Normalization to open the Normalization dialog.

![Normalization Method](image)

**Use TIC:** By default, PEAKS Studio uses the total ion current (TIC) of the samples to calculate the normalization factors. Normalized abundance for each sample is calculated from the raw abundance divided by the normalization factor for that sample.
**Manual input:** This option performs the same default calculation as auto normalization ("Use TIC") but also allows for the manual change of each sample's expected ratios. This option should be used if the expected ratio is not 1:1 for each sample as the software attempts to normalize to a 1:1 ratio.

**Use internal standard proteins:** This option displays a list of identified proteins (when a database search has been completed) that can be selected to create a normalization ratio. A search bar is included to enable the quick location of select proteins.

**No Normalization:** All samples are automatically assigned Factors of 1.0.

In PEAKS Studio 11 update, the next optional step is the newly introduced QC function. See section 6.12 Identification Quality Control (QC) for further explanation on how this function is utilized.

When QC is checked, a QC result node is added in the analysis. Select a control sample and acceptance tolerance, where the tolerance will mark a sample as pass or fail depending on the relation to the control sample. These two cannot be adjusted in the QC result node, and a new analysis would have to be run. The Data, Identification, and LFQ QC attributes are the attributes of interest. If a sample is outside the acceptance tolerance in any of these attributes with respect to the control sample, it will be marked as a failed sample. These attributes can be changed in the QC Summary filter, so they can be adjusted without needing to run another analysis should you want to
change the selections. Lastly the Standard Peptides list is used to specify peptides and their charge in the format shown (e.g. LLMGET, 2). These standard peptides will have their RT and m/z mapped for each sample that reported the peptide and charge. At this time, only peptides without modifications can be used (so adding in a modification to the sequence will not work).

Click on “Report” to proceed to the final step. Report filters can be set here which will filter the final result.

These filters are provided:

**PSM/Peptide:** this section is a mandatory section to filter out some results under the user-defined filter. Any changes to the filter will create a new analysis with old parameters and update the results and statistics.

**PSM -10lgP:** The PEAKS peptide score (-10lgP) is calculated for every peptide-spectrum match (PSM) reported by PEAKS DB, PEAKS PTM, and SPIDER. The score is derived from the p-value that indicates the statistical significance of the peptide-spectrum match. A peptide may be matched to many spectra, resulting in multiple PSMs. In that case, the peptide's score is calculated as the maximum among all PSM scores. For details of the scoring algorithm, please refer to the publication, *"PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification ", Mol Cell Proteomics, 2011 Dec 20*. A minimum requirement can be set and all identifications must pass this filter.
**PSM FDR (%):** The PSM filter can also be set to use the false discovery rate. FDR is estimated using a "decoy-fusion" approach.

**Peptide FDR (%):** The Peptides can also be set to use the false discovery rate. Peptides are scored based on their top PSM and using a "decoy-fusion" approach, the result is reported at that FDR.

**Note:** Decoy fusion is an enhanced target-decoy method for result validation with FDR. Decoy fusion appends a decoy sequence to each protein as the "negative control" for the search. See BSI's web tutorial (http://www.bioinfor.com/fdr-tutorial/) for more details.

**De Novo ALC (%):** This filter controls the minimum ALC score required for display in the de novo only result. A default of 50% is set.

**De Novo Only Tag Sharing:** The number set here corresponds to the length of de novo only tags to be shown in the result Protein coverage. For de novo only peptides that are not fully matched, they must share a tag length with of minimum of this parameter with the protein residue to be displayed.

**De Novo Only Fully Matched:** Checking this option will show the de novo only peptides in the Protein coverage regardless of what tag sharing parameter is set if the entire de novo only sequence matches the protein residue.

In the Report step, there also is the option to specify an Analysis Name (which can be later renamed) and/or decide to save the Workflow to quickly re-use in future analyses.

### 8.2 Understanding LFQ Results

Once completed, a label free quantification result node will be added to the project tree. Double-click on this [LFQ](#) node to open the LFQ result that contains multiple tabs as described below:

**Summary:** The outline of the PEAKS LFQ search results with statistics. This is the place to examine the overall performance of the experiment and adjust filters.

**Proteins:** The quantified proteins with a list of supporting peptide features for each protein.

**Peptides:** The quantified peptides listed in a table. Extracted ion chromatogram (XIC), sample features and retention time alignment also displayed for each peptide.
8.3 LFQ - Summary View

8.3.1 Summary View - Protein Filters
Filter settings used in the screenshot above means that no protein filter is applied, thus giving all quantifiable protein.

Alternately, parameters can be adjusted for additional analysis by updating the protein filters above.

**Significance:** Only protein groups with a significance above this threshold will be listed in the Protein View. The significance score is calculated as the -10log10 of the significance testing p-value. PEAKS provides ANOVA significance testing methods. A significance score threshold of 20 is recommended, which equals to a significance testing p value of 0.01. Either this or “FDR (adjusted p-value)” can be selected to set a significance threshold.

**Fold Change:** Only proteins at or above this fold change threshold will be listed in the Protein View.

**Used Peptides:** Only proteins with at least this many number of used peptides will be listed in the Protein View.

Click on **Apply** to apply these filters changes to the result. This will update charts on the Summary View and will update the Protein table accordingly.

8.3.2 Summary View - Notes
Click on **Notes** to open the Notes dialog and save information about the result and saved in section 1. Notes on the Summary page.
1. Notes

Example 1: Notes
The heat map displays the protein groups that passed the filters. The relative protein abundance is represented as a heat map of the representative proteins of each protein group. These representative proteins are clustered if they exhibit a similar expression trend across the samples. The hierarchical clustering is generated using a neighbour-joining algorithm with a Euclidean distance similarity measurement of the log2 ratios of the abundance of each sample relative to the average abundance. Similarly, the conditions in different samples are clustered if they exhibit a similar expression trend across the protein groups. An interactive protein profile heat map is opened in a new window when the button is clicked.

Mouse over the dendogram in a section of the heatmap and click on it to open the Variation Trend chart.
The Variation Trend chart displays the expression level of the selected proteins in the samples. Mouse over points on the variation trend chart to see a tooltip of the protein, sample, and log2ratio.

Click on the Export Data button to export the variation trend chart data in a text file format. When Displayed is selected, the log2 ratios will be in the export. When Original is selected, the original ratio values will be in the export instead.
8.3.4 Summary View - Density-Ratio Plot

To view the group ratio distribution of the quantified proteins, click the Density-Ratio button. By default, all proteins will be shown. Click Define Species to group the different species together.

The proteins can be searched from the list by typing into the search box, marked and then added as a species group. The expected ratio of the species in the "Ratio" column can be edited to match with the experiment design. Once all the species are added and organized, click Ok to finish. In the plot window, click Update to apply the changes.
8.3.5 Summary View - Sample Correlation plot

To view the abundance correlation plot for the detected features and proteins, click the **Sample Correlation** button from the Summary View title bar. A Correlation Plot Options window will pop-up to select which samples to include.

By default, the first 5 samples are selected. At most 5 samples can be used to generate the correlation plot, and at least 2 samples must be selected.
The Protein Correlation and Peptide Correlation plots will be displayed for the samples selected. The Pearson’s correlation score indicates the reproducibility of the experiment based on a selected pair of samples. To export the image right click on the plots and click on “Export Image”.

8.3.6 Summary View - Group Correlation plot

To view the abundance correlation plot for the detected features and proteins, click the Group Correlation button from the Summary View title bar. A Correlation Plot Options window will pop-up to select which groups to include.
By default, the first 5 groups are selected. At most 5 groups can be used to generate the correlation plot, and at least 2 groups must be selected.

The Protein Correlation and Peptide Correlation plots will be displayed for the samples selected. The Pearson’s correlation score indicates the reproducibility of the experiment based on a selected pair of samples. To export the image right click on the plots and click on “Export Image”.

The Protein Correlation and Peptide Correlation plots will be displayed for the samples selected. The Pearson’s correlation score indicates the reproducibility of the experiment based on a selected pair of samples. To export the image right click on the plots and click on “Export Image”.
8.3.7 Summary View - Volcano plot

Figure 2. volcano plot combines a statistical test with the magnitude of the change enabling quick visual identification of those data-points that display large-magnitude changes and that are also statistically significant. The volcano plot in Figure 2(a) plots significance versus fold-change of the quantified proteins.

Horizontal broken grey line: The selected significance threshold
Vertical broken grey lines: The selected fold change thresholds

Plotting points in this way results in two regions of interest in the plot: those points that are found towards the top of the plot that are far to either the left or the right side. These represent values that display large magnitude fold changes (hence being left or right of center) as well as high statistical significance (hence being towards the top).

Figure 2. The volcano plot for proteins. 

In this figure, the ratio is the group ratio set with respect to the base group. If more than two groups are present, the max ratio is used instead.

The Protein View contains an interactive volcano plot that can be hidden or displayed. Markers for the proteins that are above the set significance threshold will be displayed in colour. Scrolling the cursor over a marker will highlight it and, subsequently, clicking on this marker will highlight the protein in the interface.
8.3.8 Summary View - RT and m/z Shift Distribution
Figure 3(a) plots the retention time shift distribution to the base sample. Figure 3(b) plots the distribution of the m/z shift (in ppm) of the data with respect to the base sample. These two figures can be used to refine the parameter settings for label free quantification.

Figure 3. (a) RT shift distribution, (b) M/Z shift distribution.

8.3.9 Summary View - Missing Value percentage chart
Figure 4 shows the percentage of missing values before and after ID transfer for each sample.

Figure 4. Percentage of missing values in each sample with or without ID transfer.
8.3.10 Summary View - Filtration parameters and statistics
Tables 1 and 2 list the filtration parameters and filtered results, respectively.

**Table 1. Result filtration parameters.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Significance</td>
<td>≥ 15</td>
</tr>
<tr>
<td>Min Fold Change</td>
<td>≥ 2</td>
</tr>
<tr>
<td>Max Fold Change</td>
<td>≤ 64</td>
</tr>
<tr>
<td>Used peptides</td>
<td>≥ 1</td>
</tr>
</tbody>
</table>

**Table 2. Statistics of filtered result.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Features</td>
<td>56866</td>
</tr>
<tr>
<td>Features with ID</td>
<td>37885</td>
</tr>
<tr>
<td>Feature vectors with ID</td>
<td>9535</td>
</tr>
<tr>
<td>Protein groups</td>
<td>141</td>
</tr>
</tbody>
</table>

8.3.11 Summary View - Search parameters
Table 3 lists the parameters as set in the workflow.

**Table 3. Search Parameters**

Search Engine Name: PEAKS  
Quantification type: Label free quantification  
LFQ method: Identification directed quantification  
Mass Error Tolerance: 20.0 ppm  
Retention time shift tolerance: Auto detected  
Feature intensity ≥ 10000.0  
Retention time range: [0.0 - Max]  
Base sample: WT_1  
Peptide Feature Filter:
  - Avg area ≥ 20000.0  
  - Quality ≥ 20.0  
  - Charge between: [2 - 5]  
  - Peptide ID count ≥ 0 per group  
  - Detected in at least: 1 samples per group  
Protein Filter:
  - Significance method: ANOVA  
  - Modified Form Exclusion  
  - Remove Outliers  
  - Use Top 3 peptide  
  - Significance ≥ 0.0  
  - Fold change between: [1.0 - 64.0]  
  - Has at least: 1 used peptide  
Normalization method: Use TIC  
Samples: 6 samples in 2 groups  
SMA:
  - SMA_1  
  - SMA_2  
  - SMA_3  
CTRL:
  - WT_1  
  - WT_2  
  - WT_3  

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8.4 LFQ - Protein View

The **Proteins View** lists all the quantified proteins present in the sample, characterizes each protein at the amino acid level, and lists the supporting peptide features of each protein. It has six components:

**Protein Table**: Lists all the quantified proteins present in the sample.

**Protein View Filters**: Allows filtering the protein table to select specific proteins.

**Protein Volcano Plot**: Click on a marker that is above the significance threshold (the horizontal dashed line) set in the Summary View. Upon selection, the corresponding protein in the Protein Table will be highlighted.

**PTM Profiling**: Click on to open PTM profiling.

**Coverage**: Characterizes the protein sequence at the amino acid level. All the PTMs occurred on the protein sequence will be displayed together with the MS/MS data supporting the inference.

**Peptides**: Lists the supporting peptides identified from the selected proteins in the protein table.

8.4.1 Proteins View - Protein Table controls

At the top of the Proteins View, there are the options to filter, show/hide columns, search, and export the protein table.

![Protein View Filter](image)

141 protein groups, total 1650 proteins

accession contains

Click on the **Protein View Filter** button to open the Protein View Filter.
The following options are available for filtering the protein table:

**Show All or Top proteins in each group:** Adjust the protein list based on Protein grouping, top is selected by default. Proteins are grouped based on parsimony.

- **Top:** Show only the top proteins in each group. These proteins are supported by the most unique peptides in the group. Proteins in the group that share a subset of the unique peptides that support the top protein (subproteins) will not be displayed.
- **All:** Show all proteins in each group. Proteins are grouped together if they are supported by the same set or a sub-set of the top protein in the group. If a protein is supported by a peptide not supporting the top protein it will be added to a new group.

**Protein accession/name contains:** Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

**Protein description contains:** Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

**Note:** In these above two filters, multiple entries can be separated by a semi-colon.

**PTM contains:** Filter protein results by the presence of selected PTMs. Proteins that are supported by peptides containing modifications that match the selected PTMs will be included. If multiple PTMs match the modification name, they will all be included in the table. For SPIDER results, the mutations will also be listed in the PTM contains filter.

Click on the **Reset** button to remove all filters.

Click on **OK** to apply the Protein View Filter to the Protein table.

Click on **Cancel** to discard changes and close the dialog.
8.4.1.2 Protein Table controls - Protein count
At the top of the protein table shows the count of protein groups and number of proteins in the table based on the current Protein View Filters.

57 protein groups, total 142 proteins

8.4.1.3 Protein Table controls - Optional columns
Click on the button to see a list of sample-based optional columns.
Select the corresponding sample-based and group-based ratio in the Protein Table view.

The exports will always include the contents of these optional columns even if they are hidden.

8.4.1.4 Protein Table controls - Protein Table Search Function
An easy-to-use search function is available for the protein table. The protein accession or description substring can be searched for, and all search results can be navigated through with the Up and Down arrows.

Protein accession/name contains: Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

Protein description contains: Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

8.4.1.5 Protein Table controls - Protein Table Export
Click on the button to export the contents of the current table. This considers the Protein View Filters, the sorting applied to the table and is consistent with the view. There is the option to export the Protein table and/or the Supporting peptide table in CSV format.
8.4.2 Proteins View - Protein Table

Each row in the table represents a group of proteins that are supported by a common set of peptides. To expand the group, click the right arrow button at the left.

The columns in the Protein Table are:

- **Accession**: The accession number of the protein as seen in the FASTA database.
- **Cluster**: A unique identifier that classifies which protein group the protein belongs to.
- **Top**: Indicates whether this protein is a top protein.
- **Significance**: The protein significance.
- **Coverage (%)**: The percentage of the protein sequence that is covered by supporting peptides. The coverage is visualized by a colored bar. Light blue blocks indicate the parts of the sequence covered by low-confidence peptides. Dark blue blocks indicate the parts covered by high-confidence peptides. This is the total coverage including results from all samples.
- **#Peptides**: The number of high-confidence supporting peptides.
- **#Unique**: The number of high-confidence supporting peptides that are mapped to only one protein group. Unique peptides with same sequence but different modifications are only counted once in this number.
- **PTM**: The identified modifications displayed with color-coded icons.
- **Sample Profile**: The protein abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the sample channels, abundances, and ratios with respect to the base sample.
- **Group Profile**: The protein abundance among the groups is depicted as a heat map. The Group Profile is determined by calculated the total abundance (area) of supporting peptides within a group. Hold the cursor on a profile to view the group channels, abundances, and ratios compared to the group that contains the base sample.
- **Avg. Mass**: The protein mass calculated using the average mass.
- **Description**: The protein’s header information as seen in the FASTA database.

**Note**: For #Peptides and #Unique, two peptides with the same starting and ending positions in the protein are counted as one, regardless of their PTM forms. This is to follow the MCP (Molecular & Cellular Proteomics) guidelines.
8.4.3 Proteins View - Coverage Pane

The Protein Coverage view visually maps the supporting peptides and de novo tags to the protein selected in the Protein table. It also shows all identified sites with modifications or mutations to assist with protein characterization at the amino acid level. The “Protein Coverage” view is composed of three major components - Protein Sequence Coverage, Coverage Control Panel, and Protein Tools.

8.4.3.1 Protein Sequence Coverage

This area visualizes the coverage of the protein sequence.

Regions in the protein sequence that are covered by supporting peptides are displayed in bold font with a grey background. Confident modifications identified in supporting peptides are displayed as icons above the protein sequence. Modifications are represented by colored icons with the first letter of its modification name. If a residue is modified by more than one modification in the same supporting peptide, "*" is used instead of a letter. The header information of the protein is shown on the top of the protein sequence.

The supporting peptides and matched de novo tags can be shown as colored bars under the protein sequence.

Placing the cursor over a bar shows a tooltip with detailed information of the peptide.
Left clicking on a bar shows the details from which the supporting peptide is identified.

Right clicking on a peptide bar shows additional options. “Show in spectrum” has the same behaviour as left clicking on the peptide bar which is to open the annotated spectrum chart. “Show in table” has the function to jump to the supporting peptides table.

8.4.3.2 Coverage Control Panel
This panel controls what to display in the protein sequence coverage view.
Sequence Display Mode: The protein sequence can be displayed in either "outline" or "coverage" mode.

Outline mode: In outline mode, the protein sequence is displayed without supporting peptides and matched de novo tags. However, the sequence coverage at a specific position can be examined by left-clicking a residue.

Coverage mode: In coverage mode, all the supporting peptides and matched de novo tags are shown.

AAs per line and 10AA gap: Specify the number of amino acids per line and whether to show a gap for every 10 amino acids in the protein sequence.

Sequence Display Option: Select whether to view the protein sequence as Fasta or a specific enzyme used in the project. When a particular enzyme is selected, the protein sequence will be highlighted to emphasize expected peptides generated by the selected enzyme’s cleavage specificity.
**Modification minimal ion intensity / Ascore:** A modification or mutation is considered confident if there are fragment ions supporting the modified residue with the minimal peak intensity above the specified threshold. Only confident modifications and mutations are displayed as icons above the protein sequence.

- **Show confident modification site**
  - **minimal ion intensity**
    - 5%
  - **minimal Ascore**
    - 20%

**Modification table:** The modification table shows the modifications identified in supporting peptides on the protein. For each modification, the number of supporting PSMs with this particular modification is shown. The checkbox on the left controls whether to indicate the modification in the protein coverage.

<table>
<thead>
<tr>
<th>AM</th>
<th>PTM</th>
<th>#</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>Carbamidomethyl...</td>
<td>02</td>
</tr>
<tr>
<td>D</td>
<td>Deamidation (NQ)</td>
<td>01</td>
</tr>
<tr>
<td>D</td>
<td>Oxidation (M)</td>
<td>02</td>
</tr>
<tr>
<td>T</td>
<td>2-amino-5-oxo-but...</td>
<td>01</td>
</tr>
<tr>
<td>F</td>
<td>Formylation</td>
<td>02</td>
</tr>
<tr>
<td>K</td>
<td>Acetylation (Protein...</td>
<td>01</td>
</tr>
<tr>
<td>S</td>
<td>Sodium adduct</td>
<td>01</td>
</tr>
<tr>
<td>A</td>
<td>Amidation</td>
<td>01</td>
</tr>
</tbody>
</table>

---

### 8.4.3.3 Proteins View - Protein Tools

The Proteins View toolbar is at the upper-right corner of the protein sequence coverage tab and contains helpful protein tools.

- **Full screen:** When selected, the protein coverage window expands in full screen mode. Click to return from full screen mode.

- **Tool box:** The tool box contains the following tools:
  - Copy protein sequence. This copies the protein sequence of the current selected protein into the system clipboard.
  - Save protein coverage as image. Different scale options are provided to increase the image size.
  - Coverage statistics. This provides useful statistics based on the current selected protein.

- **PTM Profiling:** Calculates the difference in abundance between peptides with confidently identified PTMs versus unmodified peptides. Peptide feature areas are used for this comparison.

  The PTM Profiling tool provides quantitative information of modified peptides compared with unmodified peptides for the modification sites of the protein across all MS samples.
The inputs for the profiling can be set using the drop-down lists at the top-left corner of the dialog.

- **Proteins**: The accession ID of the selected protein to be analyzed for PTM profiling is shown in the title of the dialog. The selected protein cannot be changed within the dialog. To run profiling on another protein, close the dialog, select the protein of interest from the protein table and run the profiling by clicking the button.

- **Modification**: The drop-down list contains the detected and selected confident PTMs for the protein. Select the type of modification to study a specific PTM or choose “All the PTMs” to visualize the profiling information of all PTM sites at together.

  **Note**: The fixed PTMs are not used for profiling. In addition to this, the labeling modifications are not considered either.

- **Peptides**: Data source used for the analysis. “All” considers all the peptides for this protein.

- **Apply protein-level normalization**: By default, the protein-level normalization is not applied. If checked, the displayed modified/unmodified area is obtained by dividing the true modified/unmodified area by the corresponding ratio of the selected protein.

**PTM Profile Table**
The PTM Profile Table shows the detailed quantitative information for the modification sites. Each row in the table shows the modified and unmodified area for each sample of the selected PTM at each detected modification site. The table can be sorted by any column. The following list describes the contents in each column:

- **(Check box):** Allow the selection of each table row. Only the marked rows are used for the PTM Profile Chart.
- **Protein Position:** The detected modification sites in the protein. The modified amino acid symbol together with the position in the protein sequence is shown.
- **PTM:** The type of PTM being profiled.
- **-10logP:** The best -10logP identification score for the corresponding modified peptides.
- **AScore/ion intensity:** The best PTM AScore / ion intensity associated with the corresponding PTM at the specified site of the modified peptides.
- **Group Modified:** Total area of the features associated with the corresponding modified peptides for each sample.
- **Group Unmodified:** Total area of the features associated with the corresponding unmodified peptides for each sample.

![PTM Profile Table](image)

**PTM Profile Heat Map**

*All the PTMs on Q8J91|ACTN2_MOUSE*

The PTM Profile Heat Map is located at the bottom left corner in the dialog. The relative abundance of modified peptides in all groups for the selected modification sites as a heat map. The color scheme is determined by the log2 ratios of the modified area of each group relative to the average abundance. For more details about the heat map, please refer to section 8.3.3 Summary View - Heat Map
**Spectrum Annotation View**

The Spectrum Annotation View provides a graphical representation for the best modified peptide and the best unmodified peptide when “All” is selected from the “Peptides” drop-down list. The top spectrum corresponds to the best modified peptide, whereas the bottom spectrum shows the best unmodified peptide.

**Peptides Table.** Peptides table lists the modified peptides that contain the selected PTM at the specified modification site.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Peptide Position</th>
<th>Ion Intensity(%)</th>
<th>SMA Area</th>
<th>CTRL Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>LM(+5.09)</td>
<td>2</td>
<td>1</td>
<td>2.65E4</td>
<td>5.91E4</td>
</tr>
</tbody>
</table>

**All Matches Table.** The supporting PSMs of the selected peptide in Peptides Table are listed in the All Matches Table. The headers are self-explanatory. The last column Area is the area of the feature that is associated with the corresponding PSM.

<table>
<thead>
<tr>
<th>m/z</th>
<th>RT Range</th>
<th>Quality</th>
<th>z</th>
<th>Ion Intensity(%)</th>
<th>SMA Area</th>
<th>CTRL Area</th>
</tr>
</thead>
<tbody>
<tr>
<td>664.8949</td>
<td>54.64-55.38</td>
<td>65.02</td>
<td>2</td>
<td>1</td>
<td>2.65E4</td>
<td>5.91E4</td>
</tr>
</tbody>
</table>

**XICs.** It shows the XICs figure for the selected modified match in the All Matches Table.
PTM Profile Export

The PTM profile can be exported as an image as well as in text format.

**Image export:** To save the profile chart, right click and choose Save to Image... menu item to save the displayed graph as a PNG file.

**Text format export:** Click the [Save to Text Format] button at the top-right corner to export the calculated PTM Profile data to a text file in CSV format. The PTM Profile of the selected modification on the selected protein will be exported by default. In addition, profiles on all proteins for the selected modification may also be exported, if “Export ptm profile for all proteins” is checked. The exported file contains all necessary information to construct the PTM profile graph. The PTM profile data are grouped together by protein, then by modification type, and then by the modification site on the protein sequence. The headers of the CSV column are similar to the PTM Profile Table and also self-explanatory. The following items are explained for clarity:

- **Modified Peptide Sequence Window:** protein sequence containing 10 AA ahead of the modification site and 10 AA after the modification site with padding as necessary.
- **Sample Best AScore / Ion Intensity:** best AScore or Ion Intensity for each sample.

*Note:* When "Export PTM profile for all proteins" is not checked, only the profiling data that is marked in the PTM Profile Table will be exported.

8.4.3.4 Proteins View - Supporting Peptides

The supporting peptides assigned to the protein are shown in a separate tab beside the Protein Coverage view. The sequence can be clicked and a pop-up window will show up to display the feature details.
This table contains the following information of the quantifiable support peptides for the selected protein:

- **Peptide**: The amino acid sequence of the peptide, as determined in PEAKS Database or Spectral Library search. A modified residue is followed by a pair of parentheses enclosing the modification mass. All non-unique peptides and those not passing the filter will be excluded automatically.

- **Used**: Shows whether this peptide is used to calculate the protein abundance. The top three peptides with highest abundance are used to calculate the protein abundance.

- **Quality**: The feature vector quality score.

- **Significance**: The -10lg of a p-value represents the likelihood that the observed change between conditions is caused by random chance. The peptides are first separated into groups based on similar quality score. Significance is then calculated for each quality score group, or bin, using a two-tail T-test that assumes log normal distribution but does not assume equal variance. The -10log p-value is then calculated from this result and then displayed in the significance column.

- **Avg. ppm**: A peptide can have one or more feature vectors. Each feature vector has its own ppm calculated from the mass error of each feature. The average mass error of a peptide is the average of mass errors of all the feature vectors. This column displays the average mass error in ppm.

- **Avg. Area**: Each feature in the feature vector has its own area under the XIC curve. The average area of a feature vector is the average area of all the features associated with the peptide.

- **Sample Profile**: The peptide abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the samples, the peptide areas in samples, and the corresponding ratios with respect to the base sample.

- **Group Profile**: The peptide abundance among the groups is depicted as a heat map. Hold the cursor on a profile to view the groups, the peptide areas in groups, and the corresponding ratios with respect to the group which contains the base sample.

- **Group Area**: This column shows the peptide area for a group. It is calculated by summing up the areas of feature vectors within a group.

- **Max Ratio**: This shows the maximum of the fold change values compared to the reference sample.

- **RT Mean**: Each feature in the feature vector has its own retention time center. This column displays the average retention time center of the feature vector.

- **#Vector**: This shows the number of quantifiable feature vectors of a peptide.

- **Start**: This shows the protein position of the first residue of the peptide.

- **End**: This shows the protein position of the last residue (inclusive) of the peptide.

- **PTM**: The types and the numbers of modifications present in the peptide shown using color-coded icons. Scroll over each icon to see the modification name and mass.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Quality</th>
<th>Significance</th>
<th>Ave. ppm</th>
<th>Ave. Area</th>
<th>Sample Profile</th>
<th>Group Profile</th>
<th>SMR Area</th>
<th>CTBL Area</th>
<th>Max Ratio</th>
<th>RT mean</th>
<th>#Vector</th>
<th>Start</th>
<th>End</th>
<th>PTM</th>
</tr>
</thead>
<tbody>
<tr>
<td>HIR18</td>
<td>95.55</td>
<td>90.02</td>
<td>0.0</td>
<td>5.1616</td>
<td>Green</td>
<td>Red</td>
<td>4.9659</td>
<td>9.9663</td>
<td>2.0869</td>
<td>32.73</td>
<td>1</td>
<td>122</td>
<td>227</td>
<td></td>
</tr>
<tr>
<td>NQKIP</td>
<td>88.22</td>
<td>34.33</td>
<td>0.0</td>
<td>1.7008</td>
<td>Blue</td>
<td>Green</td>
<td>1.2359</td>
<td>2.19E+05</td>
<td>1.89</td>
<td>24.56</td>
<td>2</td>
<td>193</td>
<td>262</td>
<td></td>
</tr>
<tr>
<td>BRSK2D</td>
<td>65.46</td>
<td>31.66</td>
<td>0.0</td>
<td>1.0391</td>
<td>Red</td>
<td>Green</td>
<td>4.9666</td>
<td>9.9865</td>
<td>1.3899</td>
<td>32.73</td>
<td>1</td>
<td>122</td>
<td>227</td>
<td></td>
</tr>
<tr>
<td>ACV56</td>
<td>65.94</td>
<td>31.66</td>
<td>0.0</td>
<td>1.0391</td>
<td>Green</td>
<td>Red</td>
<td>1.7256</td>
<td>7.5304</td>
<td>1.0399</td>
<td>26.47</td>
<td>1</td>
<td>193</td>
<td>262</td>
<td></td>
</tr>
<tr>
<td>ACT520</td>
<td>60.91</td>
<td>31.66</td>
<td>0.0</td>
<td>1.0391</td>
<td>Red</td>
<td>Green</td>
<td>8.87047</td>
<td>3.50E+05</td>
<td>4.0495</td>
<td>21.75</td>
<td>2</td>
<td>141</td>
<td>242</td>
<td></td>
</tr>
<tr>
<td>ATG1</td>
<td>74.67</td>
<td>16.14</td>
<td>0.0</td>
<td>1.9483</td>
<td>Green</td>
<td>Red</td>
<td>1.9063</td>
<td>1.9596</td>
<td>1.1993</td>
<td>22.28</td>
<td>1</td>
<td>563</td>
<td>572</td>
<td></td>
</tr>
<tr>
<td>ATG1A</td>
<td>95.47</td>
<td>22.27</td>
<td>1.0</td>
<td>3.4945</td>
<td>Red</td>
<td>Green</td>
<td>2.2924</td>
<td>4.85E+05</td>
<td>1.0958</td>
<td>36.46</td>
<td>1</td>
<td>563</td>
<td>572</td>
<td></td>
</tr>
<tr>
<td>ATG1B</td>
<td>79.67</td>
<td>29.20</td>
<td>0.0</td>
<td>1.9787</td>
<td>Red</td>
<td>Green</td>
<td>1.9063</td>
<td>1.9596</td>
<td>1.1993</td>
<td>22.28</td>
<td>1</td>
<td>563</td>
<td>572</td>
<td></td>
</tr>
<tr>
<td>ATG1C</td>
<td>78.60</td>
<td>20.68</td>
<td>0.4</td>
<td>1.4583</td>
<td>Green</td>
<td>Red</td>
<td>1.9063</td>
<td>1.9596</td>
<td>1.1993</td>
<td>22.28</td>
<td>1</td>
<td>563</td>
<td>572</td>
<td></td>
</tr>
<tr>
<td>ATG1D</td>
<td>85.65</td>
<td>0.73</td>
<td>1.4</td>
<td>6.2713</td>
<td>Red</td>
<td>Green</td>
<td>84.57</td>
<td>8.56E+04</td>
<td>1.1698</td>
<td>28.17</td>
<td>1</td>
<td>636</td>
<td>638</td>
<td></td>
</tr>
<tr>
<td>ATG1E</td>
<td>67.27</td>
<td>1.09</td>
<td>0.8</td>
<td>1.3273</td>
<td>Green</td>
<td>Red</td>
<td>1.7473</td>
<td>3.69E+04</td>
<td>1.5368</td>
<td>49.38</td>
<td>1</td>
<td>621</td>
<td>630</td>
<td></td>
</tr>
<tr>
<td>ATG1F</td>
<td>71.17</td>
<td>0.81</td>
<td>0.5</td>
<td>7.14E+04</td>
<td>Red</td>
<td>Green</td>
<td>7.1564</td>
<td>7.09E+04</td>
<td>1.1721</td>
<td>74.00</td>
<td>1</td>
<td>740</td>
<td>739</td>
<td></td>
</tr>
<tr>
<td>ATG1G</td>
<td>73.82</td>
<td>4.18</td>
<td>0.0</td>
<td>1.9352</td>
<td>Red</td>
<td>Green</td>
<td>1.9063</td>
<td>1.9596</td>
<td>1.1993</td>
<td>22.28</td>
<td>1</td>
<td>563</td>
<td>572</td>
<td></td>
</tr>
<tr>
<td>ATG1H</td>
<td>88.72</td>
<td>26.08</td>
<td>0.0</td>
<td>1.8175</td>
<td>Green</td>
<td>Red</td>
<td>9.2352</td>
<td>2.69E+05</td>
<td>1.7295</td>
<td>29.53</td>
<td>2</td>
<td>334</td>
<td>368</td>
<td></td>
</tr>
<tr>
<td>ATG1I</td>
<td>61.70</td>
<td>4.97</td>
<td>1.4</td>
<td>5.0641</td>
<td>Red</td>
<td>Green</td>
<td>5.49E+04</td>
<td>7.70E+04</td>
<td>1.7100</td>
<td>30.68</td>
<td>1</td>
<td>563</td>
<td>572</td>
<td></td>
</tr>
<tr>
<td>ATG1J</td>
<td>87.35</td>
<td>10.48</td>
<td>0.5</td>
<td>1.0163</td>
<td>Red</td>
<td>Green</td>
<td>8.62E+04</td>
<td>1.58E+05</td>
<td>1.3498</td>
<td>40.54</td>
<td>1</td>
<td>563</td>
<td>572</td>
<td></td>
</tr>
<tr>
<td>ATG1K</td>
<td>72.77</td>
<td>11.56</td>
<td>0.0</td>
<td>4.9955</td>
<td>Green</td>
<td>Red</td>
<td>2.53E+04</td>
<td>6.27E+04</td>
<td>2.8711</td>
<td>50.17</td>
<td>1</td>
<td>663</td>
<td>683</td>
<td></td>
</tr>
<tr>
<td>ATG1L</td>
<td>54.90</td>
<td>23.75</td>
<td>0.0</td>
<td>5.8866</td>
<td>Red</td>
<td>Green</td>
<td>6.87E+04</td>
<td>8.80E+04</td>
<td>1.7597</td>
<td>51.03</td>
<td>1</td>
<td>370</td>
<td>394</td>
<td></td>
</tr>
<tr>
<td>ATG1M</td>
<td>71.10</td>
<td>42.03</td>
<td>1.3</td>
<td>4.7633</td>
<td>Red</td>
<td>Green</td>
<td>2.04E+04</td>
<td>6.91E+04</td>
<td>2.6524</td>
<td>40.31</td>
<td>2</td>
<td>510</td>
<td>523</td>
<td></td>
</tr>
</tbody>
</table>
• **Feature Detail:** Double click on a supporting peptide to go to the peptide page where the eXtracted Ion Chromatogram (XIC) curves are displayed (as seen below).

### 8.5 LFQ Result - Peptides View
Quantified proteins are displayed in the Peptide View along with their detailed information.

#### 8.5.1 Peptides View - Peptide Table
The peptides are grouped together based on quantifiable feature vectors with the same sequence. When there are more than 1000 peptides, the table is split into multiple pages.

The view is similar to the support peptide table described in the previous section Section 8.4.3.4 Proteins View - Supporting Peptides. The Peptide table entries may be filtered by using the Peptide View Filters on top of the peptide table. To reiterate, the columns are:

- **Peptide:** The amino acid sequence of the peptide, as determined in PEAKS Database or Spectral Library search. A modified residue is followed by a pair of parentheses enclosing the modification mass. All non-unique peptides and those not passing the filter will be excluded automatically.

- **Avg. Area:** Each feature in the feature vector has its own area under the XIC curve. The average area of a feature vector is the average area of all the features associated with the peptide.
• **Quality**: The feature vector quality score.

• **Significance**: The \(-10\log\) of a p-value represents the likelihood that the observed change between conditions is caused by random chance. The peptides are first separated into groups based on similar quality score. Significance is then calculated for each quality score group, or bin, using a two-tail T-test that assumes log normal distribution but does not assume equal variance. The \(-10\log\) p-value is then calculated from this result and then displayed in the significance column.

• **Avg. ppm**: A peptide can have one or more feature vectors. Each feature vector has its own ppm calculated from the mass error of each feature. The average mass error of a peptide is the average of mass errors of all the feature vectors. This column displays the average mass error in ppm.

• **Sample Profile**: The peptide abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the samples, the peptide areas in samples, and the corresponding ratios with respect to the base sample.

• **Group Profile**: The peptide abundance among the groups is depicted as a heat map. Hold the cursor on a profile to view the groups, the peptide areas in groups, and the corresponding ratios with respect to the group which contains the base sample.

• **Group Area**: This column shows the peptide area for a group. It is calculated by summing up the areas of feature vectors within a group

• **Max Ratio**: This shows the maximum of the fold change values compared to the reference sample.

• **RT Mean**: Each feature in the feature vector has its own retention time center. This column displays the average retention time center of the feature vector.

• **#Vector**: This shows the number of quantifiable feature vectors of a peptide.

• **Accession**: The proteins that the peptide was found in.

• **PTM**: The types and the numbers of modifications present in the peptide shown using color-coded icons. Scroll over each icon to see the modification name and mass.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

### 8.5.2 Peptide Table controls
The menu above the peptide table provides options to scroll through the table of peptides, filter the table, show optional columns and search for specific entries.

#### 8.5.2.1 Peptide Table controls - Pagination
For easy viewing and to improve performance, the peptide table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.

In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.
8.5.2.2 Peptide Table controls - View Filter

Click on the button to open the Peptide View Filter which is used to filter the Peptide table. The following are the filter options provided:

- **Peptide sequence contains**: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains for further filtering if necessary.

- **Significance >=**: Only peptides that pass this Significance threshold will remain in the table.

- **Fold Change >=**: Peptides with Max Ratio below this threshold will be removed from the table.

- **PTM contains**: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text they will all be included in the table.

Note: In all filters, multiple entries can be separated by a semi-colon.

Click on the Reset button to reset all filters to their default setting.

Click on OK to apply the filters to the Peptides table.

Click on Cancel to discard the changes in the View Filter.

8.5.2.3 Peptide Table controls - Optional Columns

Click on the button to open the optional columns dropdown. The sample ratios and group ratios can be displayed in the peptide table for easy view and sorting. For projects with many samples, it might be useful to hide these optional columns and reduce the width of the peptide table.
8.5.2.4 Peptide Table controls - Search function

On the top right of menu, there is a search function with 2 options. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- **m/z ≈:** Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit. For example, searching for m/z ≈ 560.21 will find all m/z between [560.20, 560.22]

- **Seq contains:** Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered. Additionally, since the peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “LVIK”, and “LVIK”

8.5.3 Feature vector selection

Click on the Feature Vector dropdown to open a dropdown to display the feature vectors that correspond to the selected peptide. The peptide -10lgP score is the highest among the -10lgP scores of its feature vectors. The peptide quality score corresponds to the average of the quality scores of all feature vectors.
8.5.4 Protein jump button

Click on to jump to a protein that contains the currently selected PSM. If there are more than 1 protein, a dropdown is revealed which allows the user to select which protein to jump to. If a protein is filtered out due to a protein summary filter or summary page protein filter, then the option to jump will be grayed out.

8.5.5 Feature Details

The Feature Details contains the eXtracted Ion Chromatogram (XIC) chart. The eXtracted Ion Chromatogram (XIC) chart displays the shape of the selected peptide feature vector over the retention time range associated with the identification. The table on the left displays both the area and the ratio of the features in the highest average abundance feature vector. This table also works as a control to select the features that will be used to draw the XIC plot. By default, the first sample in each group is selected.

Click on to switch between raw and aligned RT view.

Click on to export the XIC chart.

8.5.6 Sample Features

This tab displays a close up LC/MS view of the detected features from the selected samples. The feature boundary is shown by the bounding polygon. The center of the feature is denoted by a solid red dot on the monoisotopic peak cluster. Placing the cursor on this feature center displays a tool tip that shows the m/z value, the mean retention time, the charge state, and the intensity. The centers of the features are used to calculate the retention time alignment.

The button at the top-right corner of the tab can be used to toggle between the 2D and the 3D LC/MS views. To open a closed sample select the sample using the drop down found on the right hand side.
8.5.7 RT Alignment
This tab displays the aligned TIC curves of the selected samples with the highlighted retention time range of the selected feature. Use the toggle switch in the top-right corner to switch between the before and after RT alignment view. Click the button to view the pairwise RT alignment in the pop-up window. Select the sample pair from the drop-down menu to view the desired pairwise RT alignment.

8.6 LFQ Exporting
In Project View, double-click on to open the Export node.
8.6.1 LFQ result - Exporting - comma separated values (CSV) format

There are several options for exporting the PEAKS Label Free Quantification results in CSV format:

- **Proteins CSV**: The list of protein quantifications, filtered by the protein filters in the Summary view and will be saved to proteins.csv.
- **Peptide CSV**: A complete list of all quantified peptides that pass the peptide feature filters set on the "Summary" page and their associated details.
- **Support peptides CSV**: These are the supporting peptides of the quantified proteins and their associated details.
- **Feature Vector CSV**: All predicted peptide features and their associated quantification information. Both identified and unidentified peptide features are included. To include unidentified peptide features, set the Peptide Id Count equal to 0 in the Peptide feature filter.

8.6.2 LFQ result - Exporting - HTML format

**Export in HTML**: This will generate single or multiple HTML reports in the specified location.

The following exporting options are available:

- **Summary view**: The “Summary” view page will be saved as a summary.html file in HTML format in the specified location.
- **Protein coverage**: The coverage pane will be saved for each protein.
- **Supporting peptides**: A list of supporting peptides will be saved for each protein.
- **Best unique PSM**: The best unique PSM will be saved for each protein.
- **Export peptide list in HTML**: creates a separate HTML-containing peptide identifications.

An individual protein will have its own HTML output file, where the corresponding protein coverage, the supporting peptides, and the best unique PSM are gathered.

8.7 Quantification Quality Control (QC)

In the Quantification workflow, the Quality Control function can be included as an optional step to produce an automated QC result node for sophisticated QC analysis on the data level and protein/peptide quantification. The function is used in both DDA and DIA data.
Double-click on in the analysis to open the QC result node.

**QC Summary**

Table 1. Statistics of filtered results.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Selected Samples Count</td>
<td>5</td>
</tr>
<tr>
<td>Passed Samples Count</td>
<td>2</td>
</tr>
<tr>
<td>Ratio of Passed Samples</td>
<td>40%</td>
</tr>
<tr>
<td>Failed Samples Count</td>
<td>3</td>
</tr>
<tr>
<td>Ratio of Failed Samples</td>
<td>0%</td>
</tr>
<tr>
<td>Total Failed Attributes</td>
<td>5</td>
</tr>
</tbody>
</table>

8.7.1 Quantification Quality Control (QC) - Summary

The QC Summary page gives an overview of the passed and failed samples with the selected attributes.

Click on to open the QC summary filter which provides to option to choose samples and attributes to check whether they pass the acceptance tolerance with respect to the Control sample.
To adjust the control sample or acceptance tolerance, another analysis will need to be run. Simply click on Modify Analysis and adjust those two settings in the QC step, leaving other parameters the same.

Table 1 provides a quick glance on the number of passed and failed samples from the selected samples.

Figure 1 shows a histogram of the failed attributes and the number of samples with that failed attribute.
Table 2. Statistics of LFQ Result

<table>
<thead>
<tr>
<th>Sample Name</th>
<th># MS1</th>
<th># MS2</th>
<th># PSM</th>
<th># Identified Features</th>
<th># Quantified Features</th>
<th># Peptides</th>
<th># Quantified Peptides</th>
<th># Protein Groups</th>
<th># Quantified Protein Groups</th>
<th># PSM / # Peptides</th>
<th>ID Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control Sample</td>
<td>4848</td>
<td>16605</td>
<td>43989</td>
<td>31758</td>
<td>22988</td>
<td>14305</td>
<td>10013</td>
<td>2315</td>
<td>1945</td>
<td>3.08</td>
<td>53.76%</td>
</tr>
<tr>
<td>Average of Passed Samples</td>
<td>4889</td>
<td>16456</td>
<td>42231</td>
<td>28927</td>
<td>20351</td>
<td>14955</td>
<td>9750</td>
<td>2687</td>
<td>1945</td>
<td>2.83</td>
<td>52.61%</td>
</tr>
<tr>
<td>Ecoli30_Offgel_black</td>
<td>4738</td>
<td>16750</td>
<td>29489</td>
<td>22870</td>
<td>21640</td>
<td>8330</td>
<td>9052</td>
<td>1480</td>
<td>1847</td>
<td>3.54</td>
<td>34.88%</td>
</tr>
<tr>
<td>Ecoli30_Offgel_green</td>
<td>4982</td>
<td>16982</td>
<td>29400</td>
<td>21784</td>
<td>16431</td>
<td>12363</td>
<td>8223</td>
<td>2694</td>
<td>1844</td>
<td>2.38</td>
<td>34.84%</td>
</tr>
<tr>
<td>Ecoli30_Offgel_red</td>
<td>4884</td>
<td>16789</td>
<td>29939</td>
<td>21112</td>
<td>20661</td>
<td>10970</td>
<td>9682</td>
<td>2177</td>
<td>1939</td>
<td>2.73</td>
<td>34.84%</td>
</tr>
</tbody>
</table>

Table 2 provides more details on which particular samples have failed attributes, and the average values for those passed samples. The columns in this table were selected as attributes of interest, so they will not align with all attributes that were selected in the Summary filter.

Figure 2. Error Distribution. (a) MS1 Mass Error Distribution. (b) MS2 Mass Error Distribution.

Figure 2 shows how the MS and MS/MS distribution compares to the control sample.

Table 3. QC Parameters

<table>
<thead>
<tr>
<th>Control Sample: Ecoli10_Offgel_black</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acceptance Tolerance: 20.0%</td>
</tr>
<tr>
<td>Total Sample Count: 6</td>
</tr>
<tr>
<td>QC Attributes:</td>
</tr>
<tr>
<td># Identified Features</td>
</tr>
<tr>
<td># Peptides</td>
</tr>
<tr>
<td>Quantified Peptides</td>
</tr>
<tr>
<td>Standard Peptides</td>
</tr>
<tr>
<td>GFAVYTFFSPADAK, 2</td>
</tr>
<tr>
<td>GVAINMVTEIDK, 2</td>
</tr>
<tr>
<td>ITLDNAYMEK, 2</td>
</tr>
<tr>
<td>VDQSAVGFEYQGK, 2</td>
</tr>
<tr>
<td>LLKPDIER, 3</td>
</tr>
<tr>
<td>LLKPDIER, 2</td>
</tr>
<tr>
<td>TFAIKPDGVQR, 2</td>
</tr>
<tr>
<td>TFAIKPDGVQR, 3</td>
</tr>
<tr>
<td>TPVLWEMQR, 2</td>
</tr>
<tr>
<td>ALAAQNVEDMBQR, 2</td>
</tr>
</tbody>
</table>

Table 3 simply shows the QC parameters that were set in this analysis. The Standard Peptides list here can be highlighted and copied, allowing for easy re-use of the standard peptides in another analysis.
8.7.2 Quantification Quality Control (QC) - Data QC

The Data QC pertains to quality control measures related to the data itself. The following columns are in the table:

- **Sample Name**: Name of each sample.
- **All Attributes**: Pass/Fail flag. If a sample fails ANY attribute (not just in Data QC), it will be marked as Fail.
- **#MS1**: The number of MS scans in the last fraction in each sample.
- **#MS2**: The number of MS/MS scans in the last fraction in each sample.
- **MS2/MS1 Rate**: Calculated as #MS2/#MS1.
- **# Detected Feature**: The number of features in each sample.
- **Full Width (min)**: Measures the width of the median eluted peptide feature for each sample.
- **FWHM (sec)**: Abbreviation for Full Width at Half Maximum. This measures the number of seconds at half the maximum abundance of the median eluted peptide feature for each sample.
- **BPC**: Shows the BPC intensity of each sample.
- **TIC Correlation**: The Pearson correlation coefficient with respect to points on the control sample TIC.

The table has two selection controls. When selecting a cell, the column and row are highlighted. The highlighted column will update the histogram, showing the distribution between the control sample and all other samples.
The highlighted row is for comparing the control sample and the sample in the selected row. A Sample TIC chart is provided to examine how the TIC curve compares. Click on the button to change the view so that the curves overlap. The MS2 Count over Retention Time chart compares the cumulative MS/MS count as Retention time increases.

Right-click on the charts to export the image at any time.
8.7.3 Quantification Quality Control (QC) - Identification QC

Similar to Data QC, the Identification QC tab also provides a table with statistics of interest and relevant charts from Identification results. If PEAKS PTM Search or SPIDER was included in the analysis, then the Identification QC will consider results from the last result generated in the workflow, otherwise it will use the DB Search result.

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>All Attributes</th>
<th># MS2</th>
<th># Identified Feature</th>
<th># PSM</th>
<th># Peptide</th>
<th># Sequence</th>
<th># Protein Group</th>
<th># Top Proteins</th>
<th># All Proteins</th>
<th>Missed Cleavage Ratio</th>
<th>Enzyme Specificity Ratio</th>
<th># PSM / # Peptide Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>EcoK10_Offgtel_black</td>
<td>Pass</td>
<td>8133</td>
<td>31770</td>
<td>63000</td>
<td>16300</td>
<td>13173</td>
<td>2051</td>
<td>2383</td>
<td>3716</td>
<td>0.03</td>
<td>0.63</td>
<td>3.08</td>
</tr>
<tr>
<td>EcoK10_Offgtel_green</td>
<td>Pass</td>
<td>8179</td>
<td>31776</td>
<td>63102</td>
<td>16396</td>
<td>15340</td>
<td>2063</td>
<td>2772</td>
<td>3210</td>
<td>0.04</td>
<td>0.61</td>
<td>2.49</td>
</tr>
<tr>
<td>EcoK10_Offgtel_red</td>
<td>Fail</td>
<td>7748</td>
<td>25978</td>
<td>41141</td>
<td>13824</td>
<td>13017</td>
<td>2689</td>
<td>2792</td>
<td>3124</td>
<td>0.04</td>
<td>0.62</td>
<td>2.08</td>
</tr>
<tr>
<td>EcoK10I_Offgtel_black</td>
<td>Fail</td>
<td>84359</td>
<td>23870</td>
<td>29489</td>
<td>8210</td>
<td>7965</td>
<td>1480</td>
<td>1543</td>
<td>1820</td>
<td>0.07</td>
<td>0.94</td>
<td>3.54</td>
</tr>
<tr>
<td>EcoK10I_Offgtel_green</td>
<td>Fail</td>
<td>84306</td>
<td>21784</td>
<td>29400</td>
<td>13260</td>
<td>11778</td>
<td>2694</td>
<td>2805</td>
<td>3194</td>
<td>0.08</td>
<td>0.94</td>
<td>2.39</td>
</tr>
<tr>
<td>EcoK10I_Offgtel_red</td>
<td>Fail</td>
<td>89525</td>
<td>21112</td>
<td>29339</td>
<td>10910</td>
<td>10483</td>
<td>2177</td>
<td>2265</td>
<td>2565</td>
<td>0.08</td>
<td>0.94</td>
<td>2.73</td>
</tr>
<tr>
<td>Average Sample</td>
<td>Pass</td>
<td>82370</td>
<td>28963</td>
<td>36213</td>
<td>12946</td>
<td>13238</td>
<td>2340</td>
<td>2428</td>
<td>2776</td>
<td>0.08</td>
<td>0.93</td>
<td>1.30</td>
</tr>
</tbody>
</table>

The table contains the following columns pertaining to the Identification result:

- **Sample Name:** Name of each sample.
- **All Attributes:** Pass/Fail flag. If a sample fails ANY attribute (not just in Identification QC), it will be marked as Fail.
- **# MS2:** The number of MS2 scans in each sample.
- **# Identified Feature:** The total number of features with identified spectra.
- **# PSM:** Total number of PSMs identified in the sample.
- **# Peptide:** Total number of distinct peptides identified in the sample.
- **# Sequence:** Total number of distinct peptide backbones identified in the sample.
- **# Protein Group:** Total number of protein groups identified in the sample.
- **# Top Protein:** Total number of top proteins identified in the sample.
- **# All Protein:** Total number of proteins identified in the sample.
- **Missed Cleavage Ratio:** The ratio of identified peptides with missed cleavages versus peptides without any missed cleavages.
- **Enzyme Specificity Ratio:** The ratio of identified peptides with specific digest versus peptides without.
- **# PSM / # Peptide Rate:** The ratio of # PSM and # Peptide for each sample.
- MS1 Error Mean (SD): Shows the MS1 error mean in ppm and the standard deviation
- MS2 Error Mean (SD): Shows the MS/MS error mean in Da and the standard deviation
- ID Rate (%): Calculated as percent of #PSM / # MS2

Row and Column selection works the same as Data QC, where the selected Column will update the histogram to show the distribution of the column for each sample, and the selected Row will update the charts according to how they compare with the control sample. The charts include a venn diagram for protein and peptide overlap, a #PSM/#Peptide distribution, and the MS1 and MS2 error distributions violin plots.

Right-click on the charts to export the image at any time.

### 8.7.4 Quantification Quality Control (QC) - Quantification QC

The Quantification QC tab provides a table with statistics of interest and relevant charts from LFQ results.

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>All Attributes</th>
<th># Identified Feature</th>
<th># Quantified Feature</th>
<th>Missing Value (%)</th>
<th>Feature Correlation</th>
<th># Identified Peptide</th>
<th># Quantified Peptide</th>
<th># Identified Protein</th>
<th># Quantified Protein Group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>Pass</td>
<td>59458</td>
<td>57361</td>
<td>0.56</td>
<td>5.0000</td>
<td>48536</td>
<td>50117</td>
<td>7373</td>
<td>7041</td>
</tr>
<tr>
<td>Sample 2</td>
<td>Pass</td>
<td>51739</td>
<td>57310</td>
<td>0.58</td>
<td>5.0000</td>
<td>48394</td>
<td>50117</td>
<td>7268</td>
<td>7025</td>
</tr>
<tr>
<td>Sample 3</td>
<td>Pass</td>
<td>51689</td>
<td>57345</td>
<td>0.56</td>
<td>5.0000</td>
<td>48547</td>
<td>50117</td>
<td>7240</td>
<td>7026</td>
</tr>
<tr>
<td>Sample 4</td>
<td>Pass</td>
<td>51688</td>
<td>57345</td>
<td>0.56</td>
<td>5.0000</td>
<td>48547</td>
<td>50117</td>
<td>7240</td>
<td>7026</td>
</tr>
<tr>
<td>Sample 5</td>
<td>Pass</td>
<td>50885</td>
<td>57310</td>
<td>0.57</td>
<td>5.0000</td>
<td>48394</td>
<td>50117</td>
<td>7268</td>
<td>7025</td>
</tr>
<tr>
<td>Sample 6</td>
<td>Pass</td>
<td>52002</td>
<td>57345</td>
<td>0.56</td>
<td>5.0000</td>
<td>48547</td>
<td>50117</td>
<td>7240</td>
<td>7026</td>
</tr>
<tr>
<td>Sample 7</td>
<td>Pass</td>
<td>53323</td>
<td>57310</td>
<td>0.58</td>
<td>5.0000</td>
<td>48394</td>
<td>50117</td>
<td>7268</td>
<td>7025</td>
</tr>
<tr>
<td>Average Sample</td>
<td>Pass</td>
<td>51181</td>
<td>57347</td>
<td>0.56</td>
<td>5.0000</td>
<td>48547</td>
<td>50117</td>
<td>7240</td>
<td>7026</td>
</tr>
</tbody>
</table>

The table contains the following columns pertaining to the Quantification result:

- **Sample Name**: Name of each sample.
- **All Attributes**: Pass/Fail flag. If a sample fails ANY attribute (not just in Quantification QC), it will be marked as Fail.
- **# Identified Feature**: The total number of features with identified spectra. Same number as in Identification QC.
- **# Quantified Feature**: The total number of features used for feature vectors from the sample.
- **Missing Value (%):** Displays the missing value after ID Transfer for each sample. This is the same number as in the LFQ result.
• Feature Correlation: Measures the pearson correlation coefficient of the features compared to the control sample. This is the same value as the feature correlation in the LFQ result sample correlation pop-up.
• # Identified Peptide: Total number of distinct peptides identified in the sample. This is the same number as in Identification QC. When selecting this column, the # Quantified Peptide column is also selected.
• # Quantified Peptide: Total number of distinct peptides quantified in the sample. When selecting this column, the # Identified peptide column is also selected.
• # Identified Protein Group: Total number of protein groups identified in the sample. This is the same number as in Identification QC. When selecting this column, the # Quantified protein group column is also selected.
• # Quantified Protein Group: Total number of protein groups quantified in the sample. When selecting this column, the # Identified protein group column is also selected.

Row and Column selection works the same as Data QC, where the selected Column will update the histogram to show the distribution of the column for each sample, and the selected Row will update the charts according to how they compare with the control sample. The distribution chart for Peptides and Protein groups are slightly different in that the Identified and Quantified columns are always selected together and displayed on the same chart. A correlation plot between top protein, peptide, and feature is drawn for the selected sample and control sample. An RT Alignment chart is provided to show the retention time alignment between selected sample and control sample.

Right-click on the charts to export the image at any time.

8.7.5 Quantification Quality Control (QC) - Standard Peptides
When standard peptides were added to QC function, an additional tab is in the QC result node. A table will show which peptides are found in each sample, and the RT and m/z for those standard peptides can be viewed for quick comparison between samples.
Each sample and the standard peptides are listed in the table. The table can be sorted, but the control sample will always be the first index. Click on the columns to select/deselect that standard peptide for viewing in the RT and M/Z reproducibility charts. Multiple columns can be selected, and if none are selected then all standard peptides will be displayed. For those standard peptides of interest, there are checkboxes on the right side of the charts which can be toggled to show missing values. Those samples that are missing will have a vertical bar drawn, so that it is easier to tell which samples are missing the standard peptides.

Right-click on the charts to export the image at any time.

### 8.7.6 Identification Quality Control (QC) - QC Result Exports

As mentioned, all images in the QC result node can be exported by right-clicking on the chart, and selecting Export image.

The Export Chart pop-up will appear, and the image can be exported at different scales.
Besides image exporting, the tables can be exported in CSV files and each QC view can be exported in HTML format. Double-click the Export node to find the QC Export options.

<table>
<thead>
<tr>
<th>QC result Exports</th>
</tr>
</thead>
<tbody>
<tr>
<td>QC result in CSV</td>
</tr>
<tr>
<td>Data QC CSV</td>
</tr>
<tr>
<td>Identification QC CSV</td>
</tr>
<tr>
<td>Quantification QC CSV</td>
</tr>
<tr>
<td>Standard Peptide QC CSV</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Export in HTML</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summary view</td>
</tr>
<tr>
<td>Data QC report</td>
</tr>
<tr>
<td>Identification QC report</td>
</tr>
<tr>
<td>Quantification QC report</td>
</tr>
<tr>
<td>Standard Peptide report</td>
</tr>
</tbody>
</table>
9. Isobaric Labelling Quantification (TMT / iTRAQ)

Isobaric labelling quantification with chemical labels at the MS2 (or MS3) level is one of the three quantification modes that are supported by the optional PEAKS Q module of PEAKS Studio. In contrast to LFQ Quantification, TMT/iTRAQ Quantification is based on the relative intensities of the fragment peaks at fixed m/z values within an MS2 (or MS3) spectrum. To prepare for this, isobaric chemical labels are introduced into samples during the sample preparation stage. The samples are then combined and analyzed together in an LC-MS/MS experiment.

The same peptides from different samples will have the same precursor m/z and retention time, and are fragmented together. In the MS/MS (MS3) scans, labels from different samples will produce different reporter ions, which can then be used to calculate the quantification ratio between samples. Both user-defined and commercial labels (i.e. iTRAQ and TMT) are supported by PEAKS Q.

9.1 Isobaric Labelling Quantification Analysis Workflow and Settings

Select the PEAKS Q workflow in the Workflow Selection step

Click on Data Refine to proceed through the workflow. For Isobaric labelling quantification, it is recommended to **disable** the associate feature with Chimera option in Data Refine step. Proceed to the next step to see the DB Search parameters.
The following parameter settings are provided:

**Precursor Mass Error Tolerance:** The maximum mass difference between the peptide and the monoisotopic mass of the precursor ion.

**Precursor Mass Tolerance unit:** PEAKS generates peptides within the precursor mass error tolerance. Precursor mass error tolerance can be specified in either Daltons or ppm.

**Fragment Mass Error Tolerance:** PEAKS uses this value when scoring peptide sequences. PEAKS Studio considers a fragment ion to be matched if the calculated m/z is within the specified error tolerance. Units are Daltons.

Both precursor and fragment error tolerance parameters need to be set consistently with the mass accuracy of the instrument.

**Enzyme Settings:** PEAKS Studio digests the protein database *in silico* to generate peptide candidates. It is necessary to specify the enzyme for protein digestion from the Enzyme drop-down menu.

**Note:** When the selected dataset is digested with different enzymes, "Specified by each sample" allows samples to be analyzed separately using their respective enzymes specified during project creation.
**Note:** "None" is a special enzyme allowing non-specific cleavage at both ends of the peptide. It is the recommended option when no digestion enzyme was used or when the digestion enzyme exhibits a high degree of non-specificity.

**Missed Cleavages:** This specifies the number of missed cleavage sites allowed in a peptide.

**Digest Mode:** This specifies the type of in silico cleavages allowed at a peptide termini based on the method used for protein digestion. If "Specific" is selected, the specificity of the selected enzyme is strictly enforced at both termini of a peptide. "Semispecific" allows one terminus to disobey the enzyme specificity rules.

**Note:** If the enzyme is specified as "None", then no matter which mode is selected for "Digest Mode", the "None" enzyme digest rule will be applied.

**PTM Settings:** Click “Set PTM” to open the PTM Options dialog and specify the fixed PTMs and a few common variable PTMs expected in the experiment. Highlight PTMs and click on the button to place them as either Fixed or Variable PTM. From there, highlighted PTMs can be removed by clicking the “Remove” button. Clicking “Remove All” will reset the PTM options. Highlighting and clicking “Switch Type” will move the Fixed PTM in into the Variable PTM list and vice-verse. Click on “New” to create a new custom PTM. While a PTM is selected, click on “View” to see more details.

**Note:** Multiple PTMs can be highlighted by holding down the CTRL key or SHIFT key on the keyboard.

**Note:** For a specific residue, only one fixed modification is allowed.

Once the PTMs are selected, the maximum number of variable PTMs per peptide can be specified and a value of less than 4 is recommended.

**Note:** The use of variable modifications increases the size of the computational search space for the de novo sequencing algorithm. It is recommended not to select too many variable modifications in PEAKS de novo. See PEAKS PTM for alternate search options.

**Target Database:** Select a protein sequence database for the search. Select one from the list of databases that have been configured in PEAKS and set the taxonomy by clicking on , if applicable. To configure a new database, click on next to the Target Database dropdown to open the Database Configuration dialog.

**Contaminant Database:** To search for contaminants, select a contaminant database from the list of databases that have been configured in PEAKS. Keep the selection as N/A if no contaminant database is required for the search.

**Peptide Length:** The length for a peptide can be set when creating an analysis, where the default peptide length range is between 6 and 45.

**Deep Learning Boost:** Check this option to use PEAKS enhanced deep learning algorithm to better identify PSMs in the DB Search process.

Click on Quantification to proceed to the next step and select the Reporter Ion Quantification (eg. iTRAQ/TMT) option.
The following parameters are available:

**Select Methods:** From the Select Methods drop-down menu, choose the appropriate quantification method used in the experiment, e.g. TMT-10plex

**Mass Error Tolerance:** This parameter is used to locate the reporter ion peaks in the MS/MS scan. Either Da or ppm can be selected as the mass unit.

**Report Ion Type:** This parameter allows for the selection of MS2 or MS3 as the reporter ion type for quantification.

**Perform Purity Correction:** Select the checkbox to edit correction factors for chemical labels (if applicable). Once the checkbox is selected, click the down arrow to open the dialogue that allows users to setup the purity correction. The dialogue is setup to mimic a Certificate of Analysis, where each factor within the table can be modified by double-clicking the particular cell of interest.
**Quality Control:** The TMT QC function is used to specify replicate channels with a CV threshold filter. Those replicate channels that have CV above the filter will be filtered out and will not be used in the peptide and protein ratio calculation.

Click on the tool icon next to Spectrum Filters to open the Spectrum filter dialog.

![Spectrum Filters](image)

- **Quality:** A higher quality peptide indicates that the peptide is more quantifiable. Factors that affect the quality score include the identification -10LgP score, the noise around the reporter ions, and the mass error of reporter ions.
- **Reporter Ion Intensity:** Peptides with at least one reporter ion above this value will be kept.
- **Number of Channels Present:** Only quantifiable peptides that contain at least the number of labels selected will be used in protein quantification.
- **Reference Channel Present:** Only quantifiable peptides in which the reference label is present will be used in protein quantification. Note: The reference label is set in the experiment setting pop-up. See Experiment Setting section below

Click on the tool icon next to Protein Filters to open the Protein filter dialog

![Protein Filters](image)

- **Significance Method:** ANOVA. Significance ≥ 0.0, Unique peptides ≥ 1, 1 ≤ Fold change ≤ 64.
**Modified Form Exclusion:** Select the modified exclusion checkbox to exclude peptides with variable modifications from protein ratio calculation. Modified peptides have different ionization efficiency than unmodified ones, so this option allows users to exclude them to avoid this from having an effect on your quantification results.

**Significance Method:** ANOVA is the significance calculation that can be applied to the list of protein groups. Significance scores will only be calculated if each group contains at least two replicates. Lg (protein abundance) is used for the significance testing calculation. Please see reference below for further details.

**Note:** For more details on ANOVA, refer to the following paper: "On the comparison of several mean values: an alternative approach ", Biometrika, 1951, 38(3/4): 330-336.

The other filters are the default filters set in the TMT/iTRAQ result Summary View. They can be modified later.

**Significance:** Only protein groups with a significance above this threshold will be listed in the “Protein” view. The significance score is calculated as the -10log of the significance testing p-value (e.g. Significance score of 20 is equivalent to a p-value of 0.05). PEAKS provides the significance testing method: ANOVA.

**FDR:** Adopt the Benjamin-Hochberg method to adjust the p-value to the false discovery rate for all protein groups that have already passed the other filters. Only protein groups with significance scores passing the calculated FDR will be listed in the "Protein" view. Either this or "Significance" can be selected to set a significance threshold.

**Fold Change:** Only protein groups above the lower threshold or below the upper threshold value will be listed in the "Protein" view.

**Unique Peptides:** Only protein groups that were quantified with this number of peptides (or more) will be listed in the "Protein" view.

Click on the tool icon next to Experiment Setting to open the Experiment Setting dialog.
Select Experiment: When running quantification on an identification node that searched data within multiple samples (i.e., replicate experiments within different treatments), this drop-down menu allows for the selection of a single sample or every sample (All Experiments) for analysis. Every label of the sample(s) selected will appear in the screen located below the search bar.

Setting Sample Channel Alias: The name of the Sample Channel can be changed by using the alias name.

When there are two or more samples, the Inter-Experiment Normalization tab is enabled.
Perform Inter-Experiment Normalization: Check the Perform Inter-Experiment Normalization box if a spike label or normalization channel was used in the experiment. This is used to enable quantification between experiments.

Alias/Spike Channel Drop-down Menus: Select the channel in each sample that will be used for normalization. The software assumes that similar peptide features containing this label across experiments will approach a 1:1 ratio. Differences are assumed to be due to systematic error and all labeled channels are normalized to this spike label.

Exclude Spike Label for Significance: Since the spike label is assumed to be in a 1:1 ratio between experiments, it should not be included in the significance calculation.

Click on Intra Sample Normalization to switch the tab for intra-sample normalization.
No Normalization: If there are multiple samples in the quantification run and all samples are selected, No Normalization is chosen as the default option. It will be the only normalization option available for selection unless "Perform Inter Experiment Normalization" is check-marked in Experiment Settings.

Auto normalization: This option will calculate a global ratio from the total intensity of all labels in all quantifiable peptides. Factor values and Expected Ratios between the channels (i.e., the labels) will be presented. Selecting another sample from the Experiment drop-down menu will display the results of other samples.

Manual input: This option performs the same default calculation as Auto normalization but also allows for the manual change of each label’s Expected Ratios. After changing a ratio and hitting “Enter” on the keyboard, PEAKS will recalculate the Factor for the non-referenced label(s). The Experiment drop-down menu can be opened to display the results of another sample.

Use Internal Standard Proteins: This option displays a list of identified proteins that can be selected to create a normalization ratio. Use the search bar to quickly locate specific proteins. Right-clicking in the Protein list will open a pop-up menu to select all highlighted rows or remove all selected proteins instead. The expected ratios can be manually modified. It is assumed that the total protein content of the selected proteins in different labels have the ratios entered in the ‘Expected ratio’ column. If desired, sample names can be changed by double-clicking the chosen channel below the “Name” column and typing the new name.

Click on Experiment Groups to switch the tab and set Experiment Grouping.

Experiment Group: TMT/iTRAQ requires sample grouping. All available channels are listed in the left window. After selecting a channel, it can be added to a new group by clicking the button or to an existing group by
clicking the button and selecting that option from the drop-down menu. If you select multiple channels, you can add each one individually to create multiple new groups by clicking the button. Channels and Groups can be removed by clicking the button next to each element. The names and colours for each group can be edited by clicking on the coloured square dropdown, respectively.

Note: If no experiment groups are set, PEAKS Studio will automatically set each channel in their own group. There will be a warning pop-up to give notice of this.

Click on “Report” to proceed to the final step. Report filters can be set here which will filter the final result.
These filters are provided:

**PSM/Peptide:** this section is a mandatory section to filter out some results under the user-defined filter. Any changes to the filter will create a new analysis with old parameters and update the results and statistics.

**PSM -10lgP:** The PEAKS peptide score (-10lgP) is calculated for every peptide-spectrum match (PSM) reported by PEAKS DB, PEAKS PTM, and SPIDER. The score is derived from the p-value that indicates the statistical significance of the peptide-spectrum match. A peptide may be matched to many spectra, resulting in multiple PSMs. In that case, the peptide’s score is calculated as the maximum among all PSM scores. For details of the scoring algorithm, please refer to the publication, "*PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification*", Mol Cell Proteomics, 2011 Dec 20. A minimum requirement can be set and all identifications must pass this filter.

**PSM FDR (%):** The PSM filter can also be set to use the false discovery rate. FDR is estimated using a "decoy-fusion" approach.

**Peptide FDR (%):** The Peptides can also be set to use the false discovery rate. Peptides are scored based on their top PSM and using a "decoy-fusion" approach, the result is reported at that FDR.

**Note:** Decoy fusion is an enhanced target-decoy method for result validation with FDR. Decoy fusion appends a decoy sequence to each protein as the "negative control" for the search. See BSI's web tutorial (http://www.bioinfor.com/fdr-tutorial/) for more details.
De Novo ALC (%): This filter controls the minimum ALC score required for display in the de novo only result. A default of 50% is set.

Denovo Only Tag Sharing: The number set here corresponds to the length of de novo only tags to be shown in the result Protein coverage. For de novo only peptides that are not fully matched, they must share a tag length with of minimum of this parameter with the protein residue to be displayed.

Denovo Only Fully Matched: Checking this option will show the de novo only peptides in the Protein coverage regardless of what tag sharing parameter is set if the entire de novo only sequence matches the protein residue.

In the Report step, there also is the option to specify an Analysis Name (which can be later renamed) and/or decide to save the Workflow to quickly re-use in future analyses.

9.2 Understanding TMT/iTRAQ Results

Once completed, a TMT/iTRAQ result node will be added to the project tree. Double-click on this node to open the Isoberic Labelling Quantification result that contains three tabs as described below:

Summary: The outline of the isobaric labelling search results with statistics. This is the place to examine the overall performance of the experiment and adjust filters. If there are too many proteins, the heatmap will not show on the summary page.

Proteins: The quantified proteins with a list of supporting peptide features for each protein.

Peptides: The quantified peptides are listed in a table. Extracted ion chromatogram (XIC) and supporting peptide features are also displayed for each peptide.
9.3. TMT/iTRAQ - Summary View

9.3.1 Summary View - Protein Filters

Filter settings used in the screenshot above means that no protein filter is applied, thus giving all quantifiable protein.

Alternately, parameters can be adjusted for additional analysis by updating the protein filters above.

**Significance:** Only protein groups with a significance above this threshold will be listed in the Protein View. The significance score is calculated as the \(-10\log_{10}\) of the significance testing p-value. PEAKS provides ANOVA significance testing methods. A significance score threshold of 20 is recommended, which equals to a significance testing p-value of 0.01. Either this or “FDR (adjusted p-value)” can be selected to set a significance threshold.

**Fold Change:** Only proteins at or above this fold change threshold will be listed in the Protein View.

**Used Peptides:** Only proteins with at least this many number of used peptides will be listed in the Protein View.

Click on **Apply** to apply these filters changes to the result. This will update charts on the Summary View and will update the Protein table accordingly.

9.3.2 Summary View - Notes

Click on **Notes** to open the Notes dialog and save information about the result and saved in section 1. Notes on the Summary page.
Example 1: Notes

1. Notes

Example 1: Notes
9.3.3 Summary View - Heat Map

Figure 1: Protein profile heatmap (Cell colour represents the log2(ratio) to the average abundance across different samples)

The heat map displays the protein groups that passed the filters. The relative protein abundance is represented as a heat map of the representative proteins of each protein group. These representative proteins are clustered if they exhibit a similar expression trend across the samples. The hierarchical clustering is generated using a neighbour-joining algorithm with a Euclidean distance similarity measurement of the log2 ratios of the abundance of each sample relative to the average abundance. Similarly, the conditions in different samples are clustered if they exhibit a similar expression trend across the protein groups. An interactive protein profile heat map is opened in a new window when the button is clicked.

Mouse over the dendogram in a section of the heatmap and click on it to open the Variation Trend chart.
The Variation Trend chart displays the expression level of the selected proteins in the samples. Mouse over points on the variation trend chart to see a tooltip of the protein, sample, and log2ratio. Click on the Export Data button to export the variation trend chart data in a text file format. When Displayed is selected, the log2 ratios will be in the export. When Original is selected, the original ratio values will be in the export instead.
9.3.4 Summary View - Volcano plot

Figure 2. volcano plot combines a statistical test with the magnitude of the change enabling quick visual identification of those data-points that display large-magnitude changes and that are also statistically significant. The volcano plot in Figure 2(a) plots significance versus fold-change of the quantified proteins.

Horizontal broken grey line: The selected significance threshold

Vertical broken grey lines: The selected fold change thresholds

Plotting points in this way results in two regions of interest in the plot: those points that are found towards the top of the plot that are far to either the left or the right side. These represent values that display large magnitude fold changes (hence being left or right of center) as well as high statistical significance (hence being towards the top).

Figure 2. The volcano plot for proteins.

In this figure, the ratio is the group ratio set with respect to the base group. If more than two groups are present, the max ratio is used instead.

The Protein View contains an interactive volcano plot that can be hidden or displayed. Markers for the proteins that are above the set significance threshold will be displayed in colour. Scrolling the cursor over a marker will highlight it and, subsequently, clicking on this marker will highlight the protein in the interface.
9.3.5 Summary View - Filtration parameters and statistics

Tables 1 and 2 list the filtration parameters and filtered results, respectively.

<table>
<thead>
<tr>
<th>Table 1. Result filtration parameters.</th>
<th>Table 2. Statistics of filtered result.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Significance</td>
<td>Peptide-Spectrum Matches 23905</td>
</tr>
<tr>
<td>Min Fold Change</td>
<td>All Peptide sequences 6872</td>
</tr>
<tr>
<td>Max Fold Change</td>
<td>Protein groups 763</td>
</tr>
<tr>
<td>Used peptides</td>
<td></td>
</tr>
<tr>
<td>≥ 50</td>
<td></td>
</tr>
<tr>
<td>≥ 2</td>
<td></td>
</tr>
<tr>
<td>≤ 64</td>
<td></td>
</tr>
<tr>
<td>≥ 1</td>
<td></td>
</tr>
</tbody>
</table>

9.3.6 Summary View - Search parameters

Table 3 lists the parameters as set in the workflow.

<table>
<thead>
<tr>
<th>Table 3. Search Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Search Engine Name: PEAKS</td>
</tr>
<tr>
<td>Reporter Ion Quantification method: TMT16PLEX</td>
</tr>
<tr>
<td>Mass Error Tolerance: 15.00 ppm</td>
</tr>
<tr>
<td>Reporter Ion type: MS2</td>
</tr>
<tr>
<td>Perform Purity Correction: No</td>
</tr>
<tr>
<td>Spectrum Filters:</td>
</tr>
<tr>
<td>Quality ≥ 0</td>
</tr>
<tr>
<td>Reporter Ion Intensity ≥ 0.0</td>
</tr>
<tr>
<td>Number of channels present ≥ 1</td>
</tr>
<tr>
<td>Reference Channel Present: No</td>
</tr>
<tr>
<td>Protein Filters:</td>
</tr>
<tr>
<td>Significance method: ANOVA</td>
</tr>
<tr>
<td>Significance ≥ 0.0</td>
</tr>
<tr>
<td>Fold change between: [1.0 - 64.0]</td>
</tr>
<tr>
<td>Unique peptides ≥ 1</td>
</tr>
<tr>
<td>Select Experiment: All Experiments</td>
</tr>
<tr>
<td>Perform Inter-Experiment Normalization: Yes</td>
</tr>
<tr>
<td>Exclude Spike Channel for Significance: Yes</td>
</tr>
<tr>
<td>Normalization method: No Normalization</td>
</tr>
<tr>
<td>Groups:</td>
</tr>
<tr>
<td>Group 1:</td>
</tr>
<tr>
<td>Sample 1 TMT16-126</td>
</tr>
<tr>
<td>Sample 1 TMT16-127N</td>
</tr>
<tr>
<td>Sample 1 TMT16-127C</td>
</tr>
<tr>
<td>Sample 1 TMT16-128N</td>
</tr>
<tr>
<td>Group 2:</td>
</tr>
<tr>
<td>Sample 1 TMT16-128C</td>
</tr>
<tr>
<td>Sample 1 TMT16-129N</td>
</tr>
<tr>
<td>Sample 1 TMT16-129C</td>
</tr>
<tr>
<td>Sample 1 TMT16-130N</td>
</tr>
<tr>
<td>Group 3:</td>
</tr>
<tr>
<td>Sample 1 TMT16-130C</td>
</tr>
<tr>
<td>Sample 1 TMT16-131N</td>
</tr>
<tr>
<td>Sample 1 TMT16-131C</td>
</tr>
<tr>
<td>Sample 1 TMT16-132N</td>
</tr>
<tr>
<td>Group 4:</td>
</tr>
<tr>
<td>Sample 1 TMT16-132C</td>
</tr>
<tr>
<td>Sample 1 TMT16-133N</td>
</tr>
<tr>
<td>Sample 1 TMT16-133C</td>
</tr>
<tr>
<td>Sample 1 TMT16-134N</td>
</tr>
<tr>
<td>Reference Label: Sample 1 TMT16-126</td>
</tr>
</tbody>
</table>
The Proteins View lists all the quantified proteins present in the sample, characterizes each protein at the amino acid level, and lists the supporting peptide features of each protein. It has six components:

**Protein Table**: Lists all the quantified proteins present in the sample.

**Protein View Filters**: Allows filtering the protein table to select specific proteins.

**Protein Volcano Plot**: Click on a marker that is above the significance threshold (the horizontal dashed line) set in the Summary View. Upon selection, the corresponding protein in the Protein Table will be highlighted.

**PTM Profiling**: Click on to open PTM profiling.

**Coverage**: Characterizes the protein sequence at the amino acid level. All the PTMs occurred on the protein sequence will be displayed together with the MS/MS data supporting the inference.

**Peptides**: Lists the supporting peptides identified from the selected proteins in the protein table.

### 9.4.1 Proteins View - Protein Table controls

At the top of the Proteins View, there are the options to filter, search, and export the protein table.
9.4.1.1 Protein Table controls - Protein View Filters

Click on the button to open the Protein View Filter.

The following options are available for filtering the protein table:

**Show All or Top proteins in each group:** Adjust the protein list based on Protein grouping, top is selected by default. Proteins are grouped based on parsimony.

- **Top:** Show only the top proteins in each group. These proteins are supported by the most unique peptides in the group. Proteins in the group that share a subset of the unique peptides that support the top protein (subproteins) will not be displayed.
- **All:** Show all proteins in each group. Proteins are grouped together if they are supported by the same set or a sub-set of the top protein in the group. If a protein is supported by a peptide not supporting the top protein it will be added to a new group.

**Protein accession/name contains:** Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

**Protein description contains:** Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

**Note:** In these above two filters, multiple entries can be separated by a semi-colon.

**PTM contains:** Filter protein results by the presence of selected PTMs. Proteins that are supported by peptides containing modifications that match the selected PTMs will be included. If multiple PTMs match the modification name, they will all be included in the table. For SPIDER results, the mutations will also be listed in the PTM contains filter.

Click on the button to remove all filters.

Click on to apply the Protein View Filter to the Protein table.
Click on to discard changes and close the dialog.

### 9.4.1.2 Protein Table controls - Protein count
At the top of the protein table shows the count of protein groups and number of proteins in the table based on the current Protein View Filters.

775 protein groups, total 886 proteins

### 9.4.1.3 Protein Table controls - Protein Table Search Function
An easy-to-use search function is available for the protein table. The protein accession or description substring can be searched for, and all search results can be navigated through with the Up and Down arrows.

**Protein accession/name contains:** Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

**Protein description contains:** Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

### 9.4.1.4 Protein Table controls - Protein Table Export
Click on the button to export the contents of the current table. This considers the Protein View Filters, the sorting applied to the table and is consistent with the view. There is the option to export the Protein table and/or the Supporting peptide table in CSV format.

### 9.4.2 Proteins View - Protein Table
Each row in the table represents a group of proteins that are supported by a common set of peptides. To expand the group, click the right arrow button at the left.
The columns in the Protein Table are:

- **Accession**: The accession number of the protein as seen in the FASTA database.
- **Cluster**: A unique identifier that classifies which protein group the protein belongs to.
- **Top**: Indicates whether this protein is a top protein.
- **Significance**: The protein significance.
- **-10lgP**: The $-10\log_{10} P$ score of the top PSM.
- **Sample Profile**: The protein abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the sample channels, abundances, and ratios with respect to the base sample.
- **Group Profile**: The protein abundance among the groups is depicted as a heat map. The Group Profile is determined by calculated the total abundance (area) of supporting peptides within a group. Hold the cursor on a profile to view the group channels, abundances, and ratios compared to the group that contains the base sample.
- **Group Intensity**: The intensity of each group as in the Group Profile tooltip.
- **Group Ratio**: The ratio of each group as in the Group Ratio tooltip.
- **Coverage (%)**: The percentage of the protein sequence that is covered by supporting peptides. The coverage is visualized by a colored bar. Light blue blocks indicate the parts of the sequence covered by low-confidence peptides. Dark blue blocks indicate the parts covered by high-confidence peptides. This is the total coverage including results from all samples.
- **#Peptides**: The number of high-confidence supporting peptides.
- **#Unique**: The number of high-confidence supporting peptides that are mapped to only one protein group. Unique peptides with same sequence but different modifications are only counted once in this number.
- **PTM**: The identified modifications displayed with color-coded icons.
- **Avg. Mass**: The protein mass calculated using the average mass.
- **Description**: The protein's header information as seen in the FASTA database.

**Note**: For #Peptides and #Unique, two peptides with the same starting and ending positions in the protein are counted as one, regardless of their PTM forms. This is to follow the MCP (Molecular & Cellular Proteomics) guidelines.

### 9.4.3 Proteins View - Coverage Pane

The Protein Coverage view visually maps the supporting peptides and de novo tags to the protein selected in the Protein table. It also shows all identified sites with modifications or mutations to assist with protein
characterization at the amino acid level. The "Protein Coverage" view is composed of three major components - Protein Sequence Coverage, Coverage Control Panel, and Protein Tools.

9.4.3.1 Protein Sequence Coverage

This area visualizes the coverage of the protein sequence.

Regions in the protein sequence that are covered by supporting peptides are displayed in bold font with a grey background. Confident modifications and mutations identified in supporting peptides are displayed as icons above the protein sequence. Modifications are represented by colored icons with the first letter of its modification name. If a residue is modified by more than one modification in the same supporting peptide, "**" is used instead of a letter. The header information of the protein is shown on the top of the protein sequence.

The supporting peptides and matched de novo tags can be shown as colored bars under the protein sequence. Placing the cursor over a bar shows a tooltip with detailed information of the peptide.
Left clicking on a bar shows the details from which the supporting peptide is identified.

Right clicking on a peptide bar shows additional options. “Show in spectrum” has the same behaviour as left clicking on the peptide bar which is to open the annotated spectrum chart. “Show in table” has the function to jump to the supporting peptides table.

**9.4.3.2 Coverage Control Panel**

This panel controls what to display in the protein sequence coverage view.
Sequence Display Mode: The protein sequence can be displayed in either "outline" or "coverage" mode.

Outline mode: In outline mode, the protein sequence is displayed without supporting peptides and matched de novo tags. However, the sequence coverage at a specific position can be examined by left-clicking a residue.

Coverage mode: In coverage mode, all the supporting peptides and matched de novo tags are shown.

AAs per line and 10AA gap: Specify the number of amino acids per line and whether to show a gap for every 10 amino acids in the protein sequence.

Sequence Display Option: Select whether to view the protein sequence as Fasta or a specific enzyme used in the project. When a particular enzyme is selected, the protein sequence will be highlighted to emphasize expected peptides generated by the selected enzyme's cleavage specificity.
**Modification minimal ion intensity / Ascore:** A modification is considered confident if there are fragment ions supporting the modified residue with the minimal peak intensity above the specified threshold. Only confident modifications are displayed as icons above the protein sequence.

- **Show confident modification site**
- **Minimal ion intensity:** 5
- **Minimal Ascore:** 20

**Modification table:** The modification table shows the modifications identified in supporting peptides on the protein. For each modification, the number of supporting PSMs with this particular modification is shown. The checkbox on the left controls whether to indicate the modification in the protein coverage.

<table>
<thead>
<tr>
<th>ΔM</th>
<th>PTM</th>
<th>#</th>
</tr>
</thead>
<tbody>
<tr>
<td>T+304.21 ΔM</td>
<td>TMT Triplex</td>
<td>27</td>
</tr>
<tr>
<td>P+75.97 ΔM</td>
<td>Phosphorylation (STY)</td>
<td>27</td>
</tr>
<tr>
<td>O+15.99 ΔM</td>
<td>Oxidation (M)</td>
<td>3</td>
</tr>
</tbody>
</table>

**9.4.3.3 Proteins View - Protein Tools**
The Proteins View toolbar is at the upper-right corner of the protein sequence coverage tab and contains helpful protein tools.

- **Full screen:** When selected, the protein coverage window expands in full screen mode. Click to return from full screen mode.

- **Tool box:** The tool box contains the following tools:
  - Copy protein sequence. This copies the protein sequence of the current selected protein into the system clipboard.
  - Save protein coverage as image. Different scale options are provided to increase the image size.
  - Coverage statistics. This provides useful statistics based on the current selected protein.

**PTM Profiling:** Calculates the difference in abundance between peptides with confidently identified PTMs versus unmodified peptides. Peptide feature areas are used for this comparison.

The PTM Profiling tool provides quantitative information of modified peptides compared with unmodified peptides for the modification sites of the protein across all MS samples.
The inputs for the profiling can be set using the drop-down lists at the top-left corner of the dialog.

- **Proteins**: The accession ID of the selected protein to be analyzed for PTM profiling is shown in the title of the dialog. The selected protein can not be changed within the dialog. To run profiling on another protein, close the dialog, select the protein of interest from the protein table and run the profiling by clicking the button.

- **Modification**: The drop-down list contains the detected and selected confident PTMs for the protein. Select the type of modification to study a specific PTM or choose “All the PTMs” to visualize the profiling information of all PTM sites at together.

  **Note**: The fixed PTMs are not used for profiling. In addition to this, the labeling modifications are not considered either.

- **Peptides**: Data source used for the analysis. “All” considers all the peptides for this protein.

- **Apply protein-level normalization**: By default, the protein-level normalization is not applied. If checked, the displayed modified/unmodified area is obtained by dividing the true modified/unmodified area by the corresponding ratio of the selected protein.
The PTM Profile Table shows the detailed quantitative information for the modification sites. Each row in the table shows the modified and unmodified area for each sample of the selected PTM at each detected modification site. The table can be sorted by any column. The following list describes the contents in each column:

- **(Check box):** Allow the selection of each table row. Only the marked rows are used for the PTM Profile Chart.
- **Protein Position:** The detected modification sites in the protein. The modified amino acid symbol together with the position in the protein sequence is shown.
- **PTM:** The type of PTM being profiled.
- **-10logP:** The best -10logP identification score for the corresponding modified peptides.
- **AScore/ion intensity:** The best PTM AScore / ion intensity associated with the corresponding PTM at the specified site of the modified peptides.
- **Group Modified:** Total area of the features associated with the corresponding modified peptides for each sample.
- **Group Unmodified:** Total area of the features associated with the corresponding unmodified peptides for each sample.

![PTM Profile Heat Map](image-url)
The PTM Profile Heat Map is located at the bottom left corner in the dialog. The relative abundance of modified peptides in all groups for the selected modification sites as a heat map. The color scheme is determined by the log2 ratios of the modified area of each group relative to the average abundance. For more details about the heat map, please refer to section 9.3.3 Summary View - Heat Map.

**Spectrum Annotation View**

The Spectrum Annotation View provides a graphical representation for the best modified peptide and the best unmodified peptide when “All” is selected from the “Peptides” drop-down list. The top spectrum corresponds to the best modified peptide, whereas the bottom spectrum shows the best unmodified peptide.

**Peptides Table**: Peptides table lists the modified peptides that contain the selected PTM at the specified modification site.
All Matches Table: The supporting PSMs of the selected peptide in Peptides Table are listed in the All Matches Table. The headers are self-explanatory. The last column Area is the area of the feature that is associated with the corresponding PSM.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Peptide Position</th>
<th>Ion Intensity(%)</th>
</tr>
</thead>
</table>

XICs: It shows the XICs figure for the selected modified match in the All Matches Table.

PTM Profile Export
The PTM profile can be exported as an image as well as in text format.

**Image export:** To save the profile chart, right click and choose Save to Image... menu item to save the displayed graph as a PNG file.

**Text format export:** Click button at the top-right corner to export the calculated PTM Profile data to a text file in CSV format. The PTM Profile of the selected modification on the selected protein will be exported by default. In addition, profiles on all proteins for the selected modification may also be exported, if “Export ptm profile for all proteins” is checked. The exported file contains all necessary information to construct the PTM profile graph. The PTM profile data are grouped together by protein, then by modification type, and then by the modification site on the protein sequence. The headers of the CSV column are similar to the PTM Profile Table and also self-explanatory. The following items are explained for clarity:
• **Modified Peptide Sequence Window**: protein sequence containing 10 AA ahead of the modification site and 10 AA after the modification site with padding as necessary.

• **Sample Best AScore / Ion Intensity**: best AScore or Ion Intensity for each sample.

**Note**: When "Export PTM profile for all proteins" is not checked, only the profiling data that is marked in the PTM Profile Table will be exported.

### 9.4.3.4 Proteins View - Supporting Peptides

The supporting peptides assigned to the protein are shown in a separate tab beside the Protein Coverage view. The sequence can be clicked and a pop-up window will show up to display the feature details.

<table>
<thead>
<tr>
<th>Coverage</th>
<th>Peptides</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peptide</td>
<td>Quality</td>
</tr>
<tr>
<td>A+384.210E+55+7.9YTPM+20L+9.9</td>
<td>25.64 79.78 1.5</td>
</tr>
<tr>
<td>A+384.210E+55+7.9YTPM+20L+9.9</td>
<td>25.63 79.79 2.2</td>
</tr>
<tr>
<td>A+384.210E+55+7.9YTPM+20L+9.9</td>
<td>27.53 103.70 4.6</td>
</tr>
<tr>
<td>A+384.210E+55+7.9YTPM+20L+9.9</td>
<td>23.82 75.05 1.7</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>25.92 43.81 0.1</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>29.88 69.09 2.4</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>29.89 103.64 5.5</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>28.49 39.09 4.5</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>21.11 56.22 2.2</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>23.20 74.51 4.5</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>29.88 66.79 6.0</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>29.88 74.40 2.4</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>20.61 103.71 4.5</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>23.68 36.01 2.3</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>28.13 38.32 0.8</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>29.54 39.04 6.6</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>20.61 74.40 4.5</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>24.45 80.40 3.6</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>25.25 59.57 1.6</td>
</tr>
<tr>
<td>Cx+384.210E+55+7.9YTPM+20L+9.9</td>
<td>25.25 59.57 1.6</td>
</tr>
</tbody>
</table>

This table contains the following information of the quantifiable support peptides for the selected protein:

- **Peptide**: The amino acid sequence of the peptide, as determined in PEAKS Database or Spectral Library search. A modified residue is followed by a pair of parentheses enclosing the modification mass. All non-unique peptides and those not passing the filter will be excluded automatically.

- **Used**: Shows whether this peptide is used to calculate the protein abundance. The top three peptides with highest abundance are used to calculate the protein abundance.

- **Quality**: This defines how well the peptide can be quantified. It is calculated by taking the average of the quality scores of the feature vectors.

- **-10lgP**: The -10lg of a p-value represents the likelihood that the observed change between conditions is caused by random chance. The peptides are first separated into groups based on similar quality score. Significance is then calculated for each quality score group, or bin, using a two-tail T-test that assumes log normal distribution but does not assume equal variance. The -10log p-value is then calculated from this result and then displayed in the significance column.

- **Avg. ppm**: A peptide can have one or more feature vectors. Each feature vector has its own ppm calculated from the mass error of each feature. The average mass error of a peptide is the average of mass errors of all the feature vectors. This column displays the average mass error in ppm.

- **Sample Profile**: The peptide abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the samples, the peptide areas in samples, and the corresponding ratios with respect to the base sample.

- **Group Profile**: The peptide abundance among the groups is depicted as a heat map. Hold the cursor on a profile to view the groups, the peptide areas in groups, and the corresponding ratios with respect to the group which contains the base sample.
- **Group Intensity**: This column shows the peptide area for a group. It is calculated by summing up the intensities of feature vectors within a group.
- **Max Ratio**: This shows the maximum of the fold change values compared to the reference sample.
- **Start**: This shows the protein position of the first residue of the peptide.
- **End**: This shows the protein position of the last residue (inclusive) of the peptide.
- **PTM**: The types and the numbers of modifications present in the peptide shown using color-coded icons. Scroll over each icon to see the modification name and mass.

### 9.5 TMT/iTRAQ result - Peptides View

Quantified proteins are displayed in the Peptide View along with their detailed information.

### 9.5.1 Peptides View - Peptide Table

The peptides are grouped together based on quantifiable feature vectors with the same sequence. When there are more than 1000 peptides, the table is split into multiple pages.
The view is similar to the support peptide table described in the previous section Section 9.4.3.4 Proteins View - Supporting Peptides. The Peptide table entries may be filtered by using the Peptide View Filters on top of the peptide table. To reiterate, the columns are:

- **Peptide**: The amino acid sequence of the peptide, as determined in PEAKS Database or Spectral Library search. A modified residue is followed by a pair of parentheses enclosing the modification mass. All non-unique peptides and those not passing the filter will be excluded automatically.
- **Quality**: This defines how well the peptide can be quantified. It is calculated by taking the average of the quality scores of the feature vectors.
- **-10lgP**: The -10lg of a p-value represents the likelihood that the observed change between conditions is caused by random chance. The peptides are first separated into groups based on similar quality score. Significance is then calculated for each quality score group, or bin, using a two-tail T-test that assumes log normal distribution but does not assume equal variance. The -10log p-value is then calculated from this result and then displayed in the significance column.
- **Mass**: The theoretical mass of the peptide.
- **ppm**: A peptide can have one or more feature vectors. Each feature vector has its own ppm calculated from the mass error of each feature. The average mass error of a peptide is the average of mass errors of all the feature vectors. This column displays the average mass error in ppm.
- **m/z**: The precursor mass-to-charge ratio.
- **RT**: The retention time (elution time) at which the spectrum was recorded in the data.
- **#Feature**: The number of features associated with this peptide.
- **Sample Profile**: The peptide abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the samples, the peptide areas in samples, and the corresponding ratios with respect to the base sample.
- **Group Profile**: The peptide abundance among the groups is depicted as a heat map. Hold the cursor on a profile to view the groups, the peptide areas in groups, and the corresponding ratios with respect to the group which contains the base sample.
- **Group Intensity**: This column shows the peptide area for a group. It is calculated by summing up the intensities of feature vectors within a group.
- **Group Ratio**: This column shows the peptide ratio for a group, based on the ratio of intensities of feature vectors within a group.
- **Scan**: The Scan of the top PSM and which data file it is from.
- **PTM**: The types and the numbers of modifications present in the peptide shown using color-coded icons. Scroll over each icon to see the modification name and mass.
- **Accession**: The proteins that this peptide was quantified in.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)” . Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

### 9.5.2 Peptide Table controls

The menu above the peptide table provides options to scroll through the table of peptides, filter the table, show optional columns and search for specific entries.
9.5.2.1 Peptide Table controls - Pagination
For easy viewing and to improve performance, the peptide table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.

In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

9.5.2.2 Peptide Table controls - View Filter
Click on the Peptide View Filter button to open the Peptide View Filter which is used to filter the Peptide table. The following are the filter options provided:

- **Peptide sequence contains**: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains for further filtering if necessary.

- **Peptide score (-10lgP)**: Filters out all peptides with -10lgP scores below this threshold.

- **Peptide length**: Filters for peptides of length within this min/max threshold.

- **PTM contains**: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text they will all be included in the table.

  Note: In all filters, multiple entries can be separated by a semi-colon.

Click on the Reset button to reset all filters to their default setting.

Click on OK to apply the filters to the Peptides table.

Click on Cancel to discard the changes in the View Filter.
9.5.2.3 Peptide Table controls - Optional Columns

Click on the button to open the optional columns dropdown. By default, the group intensities and ratios are displayed. The sample intensities and ratios can also be displayed in the peptide table for easy view and sorting. For projects with many samples, it might be useful to hide these optional columns and reduce the width of the peptide table.

9.5.2.4 Peptide Table controls - Search function

On the top right of menu, there is a search function with 4 options. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- **Scan =**: Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction.
- **m/z ≈**: Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit. For example, searching for m/z = 560.21 will find all m/z between [560.20, 560.22]
- **RT ≈**: Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/- 1 of the last significant digit set. For example, searching for RT = 55.5 will find all m/z between [55.4, 55.6]
- **Seq contains**: Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered.
Additionally, since the peptide table shows both I/L and they cannot be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVK”, “IVLK”, and “IVIK”

9.5.3 Feature selection breadcrumb
There is a sample and feature selection breadcrumb which allows you to switch between features associated with the peptide. Click on the Sample in the breadcrumb to reveal other samples and the features in that sample related to the current peptide selection. From here you can select other samples, fractions, or features.

By clicking on the fraction level, all fractions in this sample are revealed as well as the features in each fraction associated with this peptide. From here you can select other fractions and features to view.

Clicking on the feature level will show all features in the currently selected fraction. Features with Feature Id = 0 are known as fake features and are constructed to enhance identification.

Switch between PSMs with the last portion of the breadcrumb.

All other components in the Peptide tab interface will update according to the currently selected PSM.

For timsTOF instruments, Precursor ID is used rather than Scan number.

9.5.4 Protein jump button
Click on  to jump to a protein that contains the currently selected PSM. If there are more than 1 protein, a dropdown is revealed which allows the user to select which protein to jump to. If a protein is filtered out due to a protein summary filter or summary page protein filter, then the option to jump will be grayed out.
9.5.5 Annotated Spectrum Chart

For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

9.5.5.1 Annotated Spectrum Chart - Chart Navigation

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.

Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.

Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.

See the whole spectrum: Double click in the spectrum or click the 1:1 button.

9.5.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings

Click on the button to show the spectrum annotation settings.
The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places**: Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation**: Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated**: Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation**: Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info**: Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity**: Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

### 9.5.5.3 Annotated Spectrum Chart - Additional Chart Controls

Click on  or  to toggle between absolute and percentage intensity values in the Y axis.

Click on  to reset the zoom level of the annotated spectrum chart to the default.

Click on  to zoom into the X axis by a factor of 2.

Click on  to zoom into the Y axis by a factor of 2.
The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

### 9.5.6 Ion Match Table

The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings".

<table>
<thead>
<tr>
<th>Ion Match</th>
<th>Survey</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td>b</td>
</tr>
<tr>
<td>10</td>
<td>1016.498</td>
</tr>
<tr>
<td>11</td>
<td>1285.576</td>
</tr>
<tr>
<td>12</td>
<td>1394.626</td>
</tr>
<tr>
<td>13</td>
<td>1481.659</td>
</tr>
<tr>
<td>14</td>
<td>1596.682</td>
</tr>
<tr>
<td>15</td>
<td>1453.703</td>
</tr>
<tr>
<td>16</td>
<td>1781.773</td>
</tr>
<tr>
<td>9</td>
<td>2122.880</td>
</tr>
</tbody>
</table>

Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

### 9.5.7 Survey Chart

The “Survey” tab displays the precursor ion spectrum. The buttons that appear in this section are identical to those explained above in Section 9.5.5.3 Annotated Spectrum Chart - Additional Chart Controls
9.5.8 Reporter Ion

<table>
<thead>
<tr>
<th>Ion Match</th>
<th>Survey</th>
<th>Reporter Ion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Channel</td>
<td>Intensity</td>
<td>Normalized Intensity</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>1.12E5</td>
<td>1.12E5</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>1.08E5</td>
<td>1.08E5</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>1.1E5</td>
<td>1.1E5</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>1.02E5</td>
<td>1.02E5</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>1.51E5</td>
<td>1.51E5</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>1.48E5</td>
<td>1.48E5</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>1.35E5</td>
<td>1.35E5</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>1.34E5</td>
<td>1.34E5</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>6.9E4</td>
<td>6.9E4</td>
</tr>
<tr>
<td>Sample 1 TMT..</td>
<td>6.86E4</td>
<td>6.86E4</td>
</tr>
</tbody>
</table>

This view provides details on characteristics of the reporter ions used in the experiment. The table provides information on the intensity, the raw intensity, and the ratio of each channel with respect to the set reference channel. Additionally, an associated MS2 or MS3 spectrum is included of the reporter ions in the experiment, determined by the selection made in the Quantification parameter settings. This is an interactive spectrum, enabling zooming in and out using the scroll wheel of the mouse. Double-clicking will zoom out and reveal the entire MS3 spectrum.

9.5.9 LC/MS Snapshot

The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The peptide feature is highlighted in purple and the selected PSM marker is blue. The span of the orange line in the LC-MS Snapshot designates the isolation window.
Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a \( m/z \) RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

### 9.6 TMT/iTRAQ Exporting

In Project View, double-click on to open the Export node.
9.6.1 TMT/iTRAQ result - Exporting - comma separated values (CSV) format
There are several options for exporting the PEAKS Isobaric Labelling Quantification results in CSV format:

- **Proteins CSV**: The list of protein quantifications, filtered by the protein filters in the Summary view and will be saved to proteins.csv.
- **Peptide CSV**: A complete list of all quantified peptides that pass the peptide feature filters set on the "Summary" page and their associated details.
- **Support peptides CSV**: These are the supporting peptides of the quantified proteins and their associated details.
- **PSM (Feature Vector) CSV**: All predicted peptide features and their associated quantification information. Both identified and unidentified peptide features are included. To include unidentified peptide features, set the Peptide Id Count equal to 0 in the Peptide feature filter.
- **Replicate CV CSV**: For TMT results with Quality Control replicate channels specified, a list of calculated CV values is exported. This will give an idea of the distribution of the variance and can be used to further tune TMT quality control settings.

9.6.2 TMT/iTRAQ result - Exporting - HTML format
**Export in HTML**: This will generate single or multiple HTML reports in the specified location.

The following exporting options are available:

- **Summary view**: The “Summary” view page will be saved as a summary.html file in HTML format in the specified location.
- **Protein coverage**: The coverage pane will be saved for each protein.
- **Supporting peptides**: A list of supporting peptides will be saved for each protein.
- **Best unique PSM**: The best unique PSM will be saved for each protein.
- **Export peptide list in html**: creates a separate HTML-containing peptide identifications.

An individual protein will have its own HTML output file, where the corresponding protein coverage, the supporting peptides, and the best unique PSM are gathered.
10. Stable Isotope Labeling by Amino acids in Cell culture (SILAC)

Precursor ion quantification with isotope labels at the MS level is one of the three quantification modes that are supported by the optional PEAKS Q module for PEAKS Studio. Typically, an unlabelled sample is mixed with a sample subjected to stable isotope labelling. The samples are then analyzed together in an LC-MS/MS experiment. The same peptide from different samples is recognized by a set of precursor ion peaks with similar retention time and mass differences equal to the expected difference caused by the stable isotope label. The ratio is calculated from the intensities of those peaks. PEAKS Q supports both user-defined labels and commercial quantification labels. PEAKS supports the analysis of Super-SILA experiments that uses a mixture of SILAC-labeled cells as a spike-in standard for accurate quantification of other unlabeled or labeled samples.

10.1 Precursor Ion Quantification Analysis Workflow and Settings

Select the PEAKS Q workflow in the Workflow Selection step

The following steps are the same as in the PEAKS DB workflow (as in section 6.1 Database Search Workflow and Parameter Settings) and is reiterated below.

Click on Data Refine to proceed through the workflow.

Data Refine Options

- Associate feature with Chimera [DOA]
The following parameter settings are provided:

**Precursor Mass Error Tolerance:** The maximum mass difference between the peptide and the monoisotopic mass of the precursor ion.

**Precursor Mass Tolerance unit:** PEAKS generates peptides within the precursor mass error tolerance. Precursor mass error tolerance can be specified in either Daltons or ppm.

**Fragment Mass Error Tolerance:** PEAKS uses this value when scoring peptide sequences. PEAKS Studio considers a fragment ion to be matched if the calculated m/z is within the specified error tolerance. Units are Daltons.

Both precursor and fragment error tolerance parameters need to be set consistently with the mass accuracy of the instrument.

**Enzyme Settings:** PEAKS Studio digests the protein database *in silico* to generate peptide candidates. It is necessary to specify the enzyme for protein digestion from the Enzyme drop-down menu.

**Note:** When the selected dataset is digested with different enzymes, "Specified by each sample" allows samples to be analyzed separately using their respective enzymes specified during project creation.
**Note:** "None" is a special enzyme allowing non-specific cleavage at both ends of the peptide. It is the recommended option when no digestion enzyme was used or when the digestion enzyme exhibits a high degree of non-specificity.

**Missed Cleavages:** This specifies the number of missed cleavage sites allowed in a peptide.

**Digest Mode:** This specifies the type of in silico cleavages allowed at a peptide termini based on the method used for protein digestion. If "Specific" is selected, the specificity of the selected enzyme is strictly enforced at both termini of a peptide. "Semispecific" allows one terminus to disobey the enzyme specificity rules.

**Note:** If the enzyme is specified as "None", then no matter which mode is selected for "Digest Mode", the "None" enzyme digest rule will be applied.

**PTM Settings:** Click "Set PTM" to open the PTM Options dialog and specify the fixed PTMs and a few common variable PTMs expected in the experiment. Highlight PTMs and click on the button to place them as either Fixed or Variable PTM. From there, highlighted PTMs can be removed by clicking the “Remove” button. Clicking “Remove All” will reset the PTM options. Highlighting and clicking “Switch Type” will move the Fixed PTM in into the Variable PTM list and vice-verse. Click on “New” to create a new custom PTM. While a PTM is selected, click on “View” to see more details.

**Note:** Multiple PTMs can be highlighted by holding down the CTRL key or SHIFT key on the keyboard.

**Note:** For a specific residue, only one fixed modification is allowed.

Once the PTMs are selected, the maximum number of variable PTMs per peptide can be specified and a value of less than 4 is recommended.

**Note:** The use of variable modifications increases the size of the computational search space for the de novo sequencing algorithm. It is recommended not to select too many variable modifications in PEAKS de novo. See PEAKS PTM for alternate search options.

**Target Database:** Select a protein sequence database for the search. Select one from the list of databases that have been configured in PEAKS and set the taxonomy by clicking on [Set View], if applicable. To configure a new database, click on [New] next to the Target Database dropdown to open the Database Configuration dialog.

**Contaminant Database:** To search for contaminants, select a contaminant database from the list of databases that have been configured in PEAKS. Keep the selection as N/A if no contaminant database is required for the search.

**Peptide Length:** The length for a peptide can be set when creating an analysis, where the default peptide length range is between 6 and 45.

**Deep Learning Boost:** Check this option to use PEAKS enhanced deep learning algorithm to better identify PSMs in the DB Search process.

Click on Quantification to proceed to the next step and select the Precursor Ion Quantification (eg. SILAC) option.
The following parameters are available:

**Select Methods:** From the Select Methods drop-down menu, choose the appropriate quantification method used in the experiment, e.g. SILAC-2plex (R10,K8)

**RT Error Range:** The retention time shift tolerance is the maximum elution time difference that is considered for the quantification of an identified peptide after RT alignment.

**Ion Mobility Range(1/k0):** Only available for timsTOF data. The maximum 1/k0 difference that is considered for the quantification of an identified peptide after RT alignment.

**Match Between Runs (MBR):** Enabled for two or more samples. The options are described below.

**Mass Error Tolerance:** Controls the allowable difference in mass between the identified MS/MS scan in one sample versus features in other samples. This can be set similar to the precursor mass error tolerance used for database search.

**Retention Time Shift:** The maximum allowable difference between the retention time of the identified MS/MS scan and the peptide feature considered for MBR.
Auto Detect: Let PEAKS Studio detect the optimal Retention Time Shift Tolerance for your data. An initial quantification run will be performed to determine the retention time shift trend. The optimal error tolerance will then be used for the full search.

Feature Intensity: Set a value for the minimum intensity a peptide feature must exhibit to be considered for MBR. By default, the integrated area under the curve of the peptide feature will be used to represent feature intensity.

CCS Error Tolerance (1/k0): This parameter is only enabled when the analysis contains timsTOF data. The maximum collisional cross section (CCS) differential between an identified MS/MS scan and peptide features in other samples to allow MBR.

Sample Groups: SILAC requires sample grouping. All available samples are listed in the left window. After selecting a sample, it can be added to a new group by clicking the button or to an existing group by clicking the button and selecting that option from the drop-down menu. If you select multiple channels, you can add each one individually to create multiple new groups by clicking the button. Samples and Groups can be removed by clicking the button next to each element. The names and colours for each group can be edited by clicking on the coloured square dropdown, respectively.

Conditions: The number of conditions depends on the quantification method selected. For example, SILAC-2plex(R10,K8) has 2 conditions and SILAC-3plex(R10,K8|R6,K4) has 3 conditions to study. Each label in a sample represents one unique condition. To remove a label from analysis, uncheck the corresponding condition. Condition names can be renamed.

Reference Condition: The condition which is the denominator of the calculated ratios. The reference condition can be chosen from the selected conditions only.

Click on the tool icon next to Peptide Feature Filters to open the Peptide Feature Filters dialog.
**Average Area:** The minimum average area of all peptide features associated with the peptide to be included in the result. Features with higher abundance have been shown to be more reproducible across replicates.

**Charge between:** Only peptides with feature vectors that all fall within this range will be used in protein quantification.

**Quality:** A quality value will be applied to each quantified peptide to represent the likely reproducibility of the quantification result. Set the minimum quality score cut-off. Factors that affect the quality score include m/z difference, RT difference, XIC shape similarity, and the feature intensities.

**Number of labels with ID present:** Only peptides with an identified MS/MS scan in the number of given labels will be included in the peptide table.

**Number of labels with feature present:** Only peptides with a predicted LC-MS peptide features in the number of given labels will be included in the peptide table.

**Reference label present:** Only quantifiable peptides in which the reference label peptide feature is present will be included in the peptide table if this is checked.

Click on the tool icon next to Protein Filters to open the Protein Filters dialog.

---

Significance Method: ANOVA, Significance ≥ 0.0, Unique peptides ≥ 1, Fold change ≥ 0.4.
**Significance:** Only protein groups with a significance above this threshold will be listed in the “Protein” view. The significance score is calculated as the -$10\lg$ of the significance testing p-value. Significance is calculated using welch’s ANOVA.

**FDR% (adjusted p-value):** Adopt the multiple hypothesis testing Benjamin-Hochberg method to adjust the p-value to the given false discovery rate. Only protein groups with significance scores passing the calculated FDR will be listed in the "Protein" view. Either this or "Significance" can be selected to set a significance threshold.

**Fold Change:** The required magnitude of change of abundance between conditions. The minimum value is one because the fold change is calculated as the magnitude of change where the lower value is always the denominator.

**Unique Peptides:** Only protein groups with unique peptides at or above this number will be listed in the "Protein" view. Peptides are considered unique if they are only matched to one protein group.

**Modified Form Exclusion:** When checked, peptides that are identified as both a modified and unmodified form will be excluded from the protein ration calculation. This does not apply to the quantification labels. When a peptide is split between an unmodified and modified form, the ratio of both forms may differ from the protein ratio. Therefore, including modified peptides in the protein ratio calculation might lead to incorrect results. By checking this box, if a peptide has modified and unmodified forms, both will not be used as candidates, if a peptide only has one modified form, it is still a candidate, but if a peptide has more than one modified form, all forms will not be used as candidates.

Click on the tool icon next to Normalization to open the Normalization dialog.

**Normalization Method:** No Normalization.
Use TIC: Using the total intensity of all quantifiable peptides from each condition, a global ratio is created. The global ratio is used as a normalization factor for all proteins and peptides.

Manual input: This option performs the same default calculation as auto normalization ("Use TIC") but also allows for the manual change of each sample's expected ratios. This should be used whenever the expected ratio between conditions isn’t 1:1.

Use internal standard proteins: This option displays a list of identified proteins, which can be selected to create a normalization factor. A search bar is included to enable the quick location of select proteins.

No Normalization: All samples are automatically assigned Factors of 1.0.

Click on “Report” to proceed to the final step. Report filters can be set here which will filter the final result.
These filters are provided:

**PSM/Peptide**: this section is a mandatory section to filter out some results under the user-defined filter. Any changes to the filter will create a new analysis with old parameters and update the results and statistics.

**PSM -10lgP**: The PEAKS peptide score (-10lgP) is calculated for every peptide-spectrum match (PSM) reported by PEAKS DB, PEAKS PTM, and SPIDER. The score is derived from the p-value that indicates the statistical significance of the peptide-spectrum match. A peptide may be matched to many spectra, resulting in multiple PSMs. In that case, the peptide’s score is calculated as the maximum among all PSM scores. For details of the scoring algorithm, please refer to the publication, *"PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification ", Mol Cell Proteomics, 2011 Dec 20*. A minimum requirement can be set and all identifications must pass this filter.

**PSM FDR (%)**: The PSM filter can also be set to use the false discovery rate. FDR is estimated using a "decoy-fusion" approach.

**Peptide FDR (%)**: The Peptides can also be set to use the false discovery rate. Peptides are scored based on their top PSM and using a "decoy-fusion" approach, the result is reported at that FDR.

**Note**: Decoy fusion is an enhanced target-decoy method for result validation with FDR. Decoy fusion appends a decoy sequence to each protein as the "negative control" for the search. See BSI’s web tutorial (http://www.bioinfor.com/fdr-tutorial/) for more details.
**De Novo ALC (%):** This filter controls the minimum ALC score required for display in the de novo only result. A default of 50% is set.

**De novo Only Tag Sharing:** The number set here corresponds to the length of de novo only tags to be shown in the result Protein coverage. For de novo only peptides that are not fully matched, they must share a tag length with of minimum of this parameter with the protein residue to be displayed.

**De novo Only Fully Matched:** Checking this option will show the de novo only peptides in the Protein coverage regardless of what tag sharing parameter is set if the entire de novo only sequence matches the protein residue.

In the Report step, there also is the option to specify an Analysis Name (which can be later renamed) and/or decide to save the Workflow to quickly re-use in future analyses.

### 10.2 Understanding SILAC Results

Once completed, a SILAC result node will be added to the project tree. Double-click on this node to open the Precursor Ion Quantification result that contains three tabs as described below:

**Summary:** The outline of the isobaric labelling search results with statistics. This is the place to examine the overall performance of the experiment and adjust filters. If there are too many proteins, the heatmap will not show on the summary page.

**Proteins:** The quantified proteins with a list of supporting peptide features for each protein.

**Peptides:** The quantified peptides are listed in a table. Extracted ion chromatogram (XIC) and supporting peptide features are also displayed for each peptide.
10.3. SILAC - Summary View

1. Notes
2. Result Statistics

Table 1: Summary Statistics

<table>
<thead>
<tr>
<th>Sample Name</th>
<th>SILAC Peptide</th>
<th>Precursor</th>
<th>SILAC Peptide</th>
<th>pID = 1 (%)</th>
<th>pID = 2 (%)</th>
<th>pID = 3 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>123</td>
<td>0</td>
<td>121</td>
<td>107 (85.1)</td>
<td>10 (1.4)</td>
<td>6 (0.3)</td>
</tr>
<tr>
<td>Sample 2</td>
<td>367</td>
<td>0</td>
<td>320</td>
<td>254 (92.2)</td>
<td>39 (1.3)</td>
<td>34 (9.3)</td>
</tr>
<tr>
<td>Sample 3</td>
<td>371</td>
<td>0</td>
<td>342</td>
<td>362 (59.6)</td>
<td>75 (20.2)</td>
<td>34 (9.2)</td>
</tr>
</tbody>
</table>

Figure 1: Protein profile heatmap (Cell color represents the log2(ratio) to the average abundance across different samples)

10.3.1 Summary View - Protein Filters

Filter settings used in the screenshot above means that no protein filter is applied, thus giving all quantifiable protein.

Alternately, parameters can be adjusted for additional analysis by updating the protein filters above.

**Significance:** Only protein groups with a significance above this threshold will be listed in the Protein View. The significance score is calculated as the -10log10 of the significance testing p-value. PEAKS provides ANOVA significance testing methods. A significance score threshold of 20 is recommended, which equals to a significance testing p value of 0.01. Either this or “FDR (adjusted p-value)” can be selected to set a significance threshold.

**Fold Change:** Only proteins at or above this fold change threshold will be listed in the Protein View.

**Used Peptides:** Only proteins with at least this many number of used peptides will be listed in the Protein View.

Click on Apply to apply these filters changes to the result. This will update charts on the Summary View and will update the Protein table accordingly.

10.3.2 Summary View - Notes

Click on Notes to open the Notes dialog and save information about the result and saved in section 1. Notes on the Summary page.
1. Notes

Example 1: Notes
The heat map displays the protein groups that passed the filters. The relative protein abundance is represented as a heat map of the representative proteins of each protein group. These representative proteins are clustered if they exhibit a similar expression trend across the samples. The hierarchical clustering is generated using a neighbour-joining algorithm with a Euclidean distance similarity measurement of the log2 ratios of the abundance of each sample relative to the average abundance. Similarly, the conditions in different samples are clustered if they exhibit a similar expression trend across the protein groups. An interactive protein profile heat map is opened in a new window when the button is clicked options.

Mouse over the dendogram in a section of the heatmap and click on it to open the Variation Trend chart.
The Variation Trend chart displays the expression level of the selected proteins in the samples.

Mouse over points on the variation trend chart to see a tooltip of the protein, sample, and log2ratio.

Click on the Export Data button to export the variation trend chart data in a text file format. When Displayed is selected, the log2 ratios will be in the export. When Original is selected, the original ratio values will be in the export instead.

10.3.4 Summary View - Volcano plot

Figure 2. Volcano plot combines a statistical test with the magnitude of the change enabling quick visual identification of those data-points that display large-magnitude changes and that are also statistically significant. A separate volcano plot will be shown for each condition relative to the reference condition.

Horizontal broken grey line: The selected significance threshold
Vertical grey lines: The selected fold change thresholds

Plotting points in this way results in two regions of interest in the plot: those points that are found towards the top of the plot that are far to either the left or the right side. These represent values that display large magnitude fold changes (hence being left or right of center) as well as high statistical significance (hence being towards the top).
In this figure, the ratio is the group ratio set with respect to the base group. If more than two groups are present, the max ratio is used instead.

The Protein View contains an interactive volcano plot that can be hidden or displayed. Markers for the proteins that are above the set significance threshold will be displayed in colour. Scrolling the cursor over a marker will highlight it and, subsequently, clicking on this marker will highlight the protein in the interface.

10.3.5 Summary View - Ratio Distribution Histogram

Click on the button in the Summary header to open the Ratio Distribution Histogram.

Select from the sample dropdown to view the ratio distribution of each sample.

For every condition, the distribution of ratios relative to the reference condition is plotted. Right-click on the image to export the chart.
10.3.6 Summary View - Filtration parameters and statistics

Tables 2 and 3 list the filtration parameters and filtered results, respectively.

<table>
<thead>
<tr>
<th>Table 2. Result filtration parameters.</th>
<th>Table 3. Statistics of filtered result.</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDR (adjusted p-value)</td>
<td>SILAC Peptide</td>
</tr>
<tr>
<td></td>
<td>6478</td>
</tr>
<tr>
<td>Min Fold Change</td>
<td>All Peptide sequences</td>
</tr>
<tr>
<td>≥ 1</td>
<td>10618</td>
</tr>
<tr>
<td>Max Fold Change</td>
<td>Features</td>
</tr>
<tr>
<td>≤ 64</td>
<td>52487</td>
</tr>
<tr>
<td>Used peptides</td>
<td>SILAC Precursor</td>
</tr>
<tr>
<td>≥ 1</td>
<td>17779</td>
</tr>
<tr>
<td></td>
<td>Protein groups</td>
</tr>
<tr>
<td></td>
<td>448</td>
</tr>
</tbody>
</table>

10.3.7 Summary View - Search parameters

Table 4 lists the parameters as set in the workflow.

<table>
<thead>
<tr>
<th>Table 4. Search Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Search Engine Name: PEAKS</td>
</tr>
<tr>
<td>Quantification type: ICAT/SILAC</td>
</tr>
<tr>
<td>Silac Method: SILAC-3plex (R10,K8</td>
</tr>
<tr>
<td>Retention Time Tolerance: 1.00</td>
</tr>
<tr>
<td>Enable Match Between Run: No</td>
</tr>
<tr>
<td>5 samples in 1 group</td>
</tr>
<tr>
<td>Group 1:</td>
</tr>
<tr>
<td>Sample 1:</td>
</tr>
<tr>
<td>Light: Condition 1</td>
</tr>
<tr>
<td>Medium: Condition 2</td>
</tr>
<tr>
<td>Heavy: Condition 3</td>
</tr>
<tr>
<td>Sample 2:</td>
</tr>
<tr>
<td>Light: Condition 1</td>
</tr>
<tr>
<td>Medium: Condition 2</td>
</tr>
<tr>
<td>Heavy: Condition 3</td>
</tr>
<tr>
<td>Sample 3:</td>
</tr>
<tr>
<td>Light: Condition 1</td>
</tr>
<tr>
<td>Medium: Condition 2</td>
</tr>
<tr>
<td>Heavy: Condition 3</td>
</tr>
<tr>
<td>Sample 4:</td>
</tr>
<tr>
<td>Light: Condition 1</td>
</tr>
<tr>
<td>Medium: Condition 2</td>
</tr>
<tr>
<td>Heavy: Condition 3</td>
</tr>
<tr>
<td>Sample 5:</td>
</tr>
<tr>
<td>Light: Condition 1</td>
</tr>
<tr>
<td>Medium: Condition 2</td>
</tr>
<tr>
<td>Heavy: Condition 3</td>
</tr>
<tr>
<td>Conditions: Condition 1 Condition 2 Condition 3</td>
</tr>
<tr>
<td>Reference Condition: Condition 1</td>
</tr>
<tr>
<td>Peptide Feature Filter:</td>
</tr>
<tr>
<td>Avg. area ≥ 0.0</td>
</tr>
<tr>
<td>Qualty ≥ 1.0</td>
</tr>
<tr>
<td>Charge between: [1 - 10]</td>
</tr>
<tr>
<td>Number of labels with ID present ≥ 1</td>
</tr>
<tr>
<td>Number of labels with feature present ≥ 1</td>
</tr>
<tr>
<td>Protein Filter:</td>
</tr>
<tr>
<td>Significance method ANOVA</td>
</tr>
<tr>
<td>FDR ≤ 0.01</td>
</tr>
<tr>
<td>Fold change between: [0.0 - 64.0]</td>
</tr>
<tr>
<td>Unique peptides ≥ 1</td>
</tr>
<tr>
<td>Normalization method: No Normalization</td>
</tr>
</tbody>
</table>
10.4 SILAC - Protein View

The **Proteins View** lists all the quantified proteins present in the sample, characterizes each protein at the amino acid level, and lists the supporting peptide features of each protein. It has six components:

**Protein Table**: Lists all the quantified proteins present in the sample.

**Protein View Filters**: Allows filtering the protein table to select specific proteins.

**Protein Volcano Plot**: Click on a marker that is above the significance threshold (the horizontal dashed line) set in the Summary View. Upon selection, the corresponding protein in the Protein Table will be highlighted.

**PTM Profiling**: Click on  to open PTM profiling.

**Coverage**: Characterizes the protein sequence at the amino acid level. All the PTMs occurred on the protein sequence will be displayed together with the MS/MS data supporting the inference.

**Peptides**: Lists the supporting peptides identified from the selected proteins in the protein table.

10.4.1 Proteins View - Protein Table controls

At the top of the Proteins View, there are the options to filter, search, and export the protein table.
10.4.1.1 Protein Table controls - Protein View Filters

Click on the [Protein View Filter] button to open the Protein View Filter.

The following options are available for filtering the protein table:

**Show All or Top proteins in each group**: Adjust the protein list based on Protein grouping, top is selected by default. Proteins are grouped based on parsimony.

- **Top**: Show only the top proteins in each group. These proteins are supported by the most unique peptides in the group. Proteins in the group that share a subset of the unique peptides that support the top protein (subproteins) will not be displayed.
- **All**: Show all proteins in each group. Proteins are grouped together if they are supported by the same set or a sub-set of the top protein in the group. If a protein is supported by a peptide not supporting the top protein it will be added to a new group.

**Protein accession/name contains**: Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

**Protein description contains**: Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

**Note**: In these above two filters, multiple entries can be separated by a semi-colon.

**PTM contains**: Filter protein results by the presence of selected PTMs. Proteins that are supported by peptides containing modifications that match the selected PTMs will be included. If multiple PTMs match the modification name, they will all be included in the table.

Click on the [Reset] button to remove all filters.

Click on [OK] to apply the Protein View Filter to the Protein table.
Click on [Cancel] to discard changes and close the dialog.

10.4.1.2 Protein Table controls - Protein count
At the top of the protein table shows the count of protein groups and number of proteins in the table based on the current Protein View Filters.

775 protein groups, total 886 proteins

10.4.1.3 Protein Table controls - Protein Table Search Function
An easy-to-use search function is available for the protein table. The protein accession or description substring can be searched for, and all search results can be navigated through with the Up and Down arrows.

Protein accession/name contains: Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

Protein description contains: Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

10.4.1.4 Protein Table controls - Protein Table Export
Click on the [ ] button to export the contents of the current table. This considers the Protein View Filters, the sorting applied to the table and is consistent with the view. There is the option to export the Protein table and/or the Supporting peptide table in CSV format.

10.4.2 Proteins View - Protein Table
Each row in the table represents a group of proteins that are supported by a common set of peptides. To expand the group, click the right arrow button at the left.
The columns in the Protein Table are:

- **Accession**: The accession number of the protein as seen in the FASTA database.
- **Cluster**: A unique identifier that classifies which protein group the protein belongs to.
- **Top**: Indicates whether this protein is a top protein.
- **Condition Significance**: The significance for each condition
- **-10lgP**: The -10lgP score of the top PSM.
- **Coverage (%)**: The percentage of the protein sequence that is covered by supporting peptides. The coverage is visualized by a colored bar. Light blue blocks indicate the parts of the sequence covered by low-confidence peptides. Dark blue blocks indicate the parts covered by high-confidence peptides. This is the total coverage including results from all samples.
- **#Peptides**: The number of high-confidence supporting peptides.
- **#Unique**: The number of high-confidence supporting peptides that are mapped to only one protein group. Unique peptides with same sequence but different modifications are only counted once in this number.
- **PTM**: The identified modifications displayed with color-coded icons.
- **Condition Profile**: The protein abundance among the samples is depicted as a heat map. Hold the cursor on a profile to view the sample channels, abundances, and ratios with respect to the base sample.
- **Condition Ratio**: The ratio columns show the peptide abundance in each condition for all groups. A peptide's abundance in each condition is the sum of all supporting feature vectors abundance within the filters. Then the peptide abundance in each condition is compared to the reference condition to obtain the peptide condition ratio in each sample separately. The peptide condition ratio of a group is computed by calculating the median from the peptide condition ratios in the samples in that group.
- **Avg. Mass**: The protein mass calculated using the average mass.
- **Description**: The protein's header information as seen in the FASTA database.

Note: For #Peptides and #Unique, two peptides with the same starting and ending positions in the protein are counted as one, regardless of their PTM forms. This is to follow the MCP (Molecular & Cellular Proteomics) guidelines.

### 10.4.3 Proteins View - Coverage Pane

The Protein Coverage view visually maps the supporting peptides and de novo tags to the protein selected in the Protein table. It also shows all identified sites with modifications or mutations to assist with protein characterization at the amino acid level. The "Protein Coverage" view is composed of three major components - Protein Sequence Coverage, Coverage Control Panel, and Protein Tools.
### 10.4.3.1 Protein Sequence Coverage

This area visualizes the coverage of the protein sequence.

Regions in the protein sequence that are covered by supporting peptides are displayed in bold font with a grey background. Confident modifications and mutations identified in supporting peptides are displayed as icons above the protein sequence. Modifications are represented by colored icons with the first letter of its modification name. If a residue is modified by more than one modification in the same supporting peptide, "**" is used instead of a letter. The header information of the protein is shown on the top of the protein sequence.

The supporting peptides and matched de novo tags can be shown as colored bars under the protein sequence.

Placing the cursor over a bar shows a tooltip with detailed information of the peptide.
Left clicking on a bar shows the details from which the supporting peptide is identified.

Right clicking on a peptide bar shows additional options. “Show in spectrum” has the same behaviour as left clicking on the peptide bar which is to open the annotated spectrum chart. “Show in table” has the function to jump to the supporting peptides table.

10.4.3.2 Coverage Control Panel
This panel controls what to display in the protein sequence coverage view.
Sequence Display Mode: The protein sequence can be displayed in either "outline" or "coverage" mode.

Outline mode: In outline mode, the protein sequence is displayed without supporting peptides and matched de novo tags. However, the sequence coverage at a specific position can be examined by left-clicking a residue.

Coverage mode: In coverage mode, all the supporting peptides and matched de novo tags are shown.

AAs per line and 10AA gap: Specify the number of amino acids per line and whether to show a gap for every 10 amino acids in the protein sequence.

Sequence Display Option: Select whether to view the protein sequence as Fasta or a specific enzyme used in the project. When a particular enzyme is selected, the protein sequence will be highlighted to emphasize expected peptides generated by the selected enzyme's cleavage specificity.
Modification minimal ion intensity / Ascore: A modification is considered confident if there are fragment ions supporting the modified residue with the minimal peak intensity above the specified threshold. Only confident modifications are displayed as icons above the protein sequence.

Modification table: The modification table shows the modifications identified in supporting peptides on the protein. For each modification, the number of supporting PSMs with this particular modification is shown. The checkbox on the left controls whether to indicate the modification in the protein coverage.

10.4.3.3 Proteins View - Protein Tools
The Proteins View toolbar is at the upper-right corner of the protein sequence coverage tab and contains helpful protein tools.

Full screen: When selected, the protein coverage window expands in full screen mode. Click to return from full screen mode.

Tool box: The tool box contains the following tools:
- Copy protein sequence. This copies the protein sequence of the current selected protein into the system clipboard.
- Save protein coverage as image. Different scale options are provided to increase the image size.
- Coverage statistics. This provides useful statistics based on the current selected protein.

PTM Profiling: Calculates the difference in abundance between peptides with confidently identified PTMs versus unmodified peptides. Peptide feature areas are used for this comparison.

The PTM Profiling tool provides quantitative information of modified peptides compared with unmodified peptides for the modification sites of the protein across all MS samples.
The inputs for the profiling can be set using the drop-down lists at the top-left corner of the dialog.

- **Proteins**: The accession ID of the selected protein to be analyzed for PTM profiling is shown in the title of the dialog. The selected protein can not be changed within the dialog. To run profiling on another protein, close the dialog, select the protein of interest from the protein table and run the profiling by clicking the button.

- **Modification**: The drop-down list contains the detected and selected confident PTMs for the protein. Select the type of modification to study a specific PTM or choose “All the PTMs” to visualize the profiling information of all PTM sites at together.

  **Note**: The fixed PTMs are not used for profiling. In addition to this, the labeling modifications are not considered either.

- **Peptides**: Data source used for the analysis. “All” considers all the peptides for this protein.

- **Apply protein-level normalization**: By default, the protein-level normalization is not applied. If checked, the displayed modified/unmodified area is obtained by dividing the true modified/unmodified area by the corresponding ratio of the selected protein.

**PTM Profile Table**
The PTM Profile Table shows the detailed quantitative information for the modification sites. Each row in the table shows the modified and unmodified area for each sample of the selected PTM at each detected modification site. The table can be sorted by any column. The following list describes the contents in each column:

- **(Check box):** Allow the selection of each table row. Only the marked rows are used for the PTM Profile Chart.
- **Protein Position:** The detected modification sites in the protein. The modified amino acid symbol together with the position in the protein sequence is shown.
- **PTM:** The type of PTM being profiled.
- **-10logP:** The best -10logP identification score for the corresponding modified peptides.
- **AScore/ion intensity:** The best PTM AScore / ion intensity associated with the corresponding PTM at the specified site of the modified peptides.
- **Condition Sample Ratio:** Displays the ratio for each Condition versus the reference Condition for each sample.

PTM Profile Heat Map
The PTM Profile Heat Map is located at the bottom left corner in the dialog. The relative abundance of modified peptides in all groups for the selected modification sites as a heat map. The color scheme is determined by the log2 ratios of the modified area of each group relative to the average abundance. For more details about the heat map, please refer to section 10.3.3 Summary View - Heat Map.

**Spectrum Annotation View**

The Spectrum Annotation View provides a graphical representation for the best modified peptide and the best unmodified peptide when “All” is selected from the “Peptides” drop-down list. The top spectrum corresponds to the best modified peptide, whereas the bottom spectrum shows the best unmodified peptide.
Peptides Table: Peptides table lists the modified peptides that contain the selected PTM at the specified modification site.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Peptide Position</th>
<th>Ion Intensity (%)</th>
<th>Condition 1:Sample 1 Ratio</th>
<th>Condition 2:Sample 2 Ratio</th>
<th>Condition 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>M(+58.01)EDSM(+15.98)DM(+15.09)</td>
<td>1</td>
<td>0.31</td>
<td>-</td>
<td>0.33</td>
<td>0.49</td>
</tr>
<tr>
<td>M(+58.01)EDS(+15.98)DM(+15.09)</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

All Matches Table: The supporting PSMs of the selected peptide in Peptides Table are listed in the All Matches Table. The headers are self-explanatory. The last column Area is the area of the feature that is associated with the corresponding PSM.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Fraction</th>
<th>z</th>
<th>Quality</th>
<th>Condition 1 Area</th>
<th>Condition 2 Area</th>
<th>Condition 3 Area</th>
<th>Ion Intensity(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample 1</td>
<td>k5_ac_17042017_phos.raw</td>
<td>3</td>
<td>74.74</td>
<td>1.0434E5</td>
<td>3.1908E4</td>
<td>3.0446E4</td>
<td>0</td>
</tr>
</tbody>
</table>

XICs: It shows the XICs figure for the selected modified match in the All Matches Table. Each condition is plotted as a separate curve.
PTM Profile Export

The PTM profile can be exported as an image as well as in text format.

**Image export:** To save the profile chart, right click and choose Save to Image... menu item to save the displayed graph as a PNG file.

**Text format export:** Click the [Save to Text Format] button at the top-right corner to export the calculated PTM Profile data to a text file in CSV format. The PTM Profile of the selected modification on the selected protein will be exported by default. In addition, profiles on all proteins for the selected modification may also be exported, if “Export ptm profile for all proteins” is checked. The exported file contains all necessary information to construct the PTM profile graph. The PTM profile data are grouped together by protein, then by modification type, and then by the modification site on the protein sequence. The headers of the CSV column are similar to the PTM Profile Table and also self-explanatory. The following items are explained for clarity:

- **Modified Peptide Sequence Window:** protein sequence containing 10 AA ahead of the modification site and 10 AA after the modification site with padding as necessary.
- **Sample Best AScore / Ion Intensity:** best AScore or Ion Intensity for each sample.

**Note:** When “Export PTM profile for all proteins” is not checked, only the profiling data that is marked in the PTM Profile Table will be exported.

10.4.3.4 Proteins View - Supporting Peptides

The supporting peptides assigned to the protein are shown in a separate tab beside the Protein Coverage view. The sequence can be clicked and a pop-up window will show up to display the feature details.
This table contains the following information of the quantifiable support peptides for the selected protein:

- **Peptide**: The amino acid sequence of the peptide, as determined in PEAKS Database or Spectral Library search. A modified residue is followed by a pair of parentheses enclosing the modification mass. All non-unique peptides and those not passing the filter will be excluded automatically.

- **Used**: Shows whether this peptide is used to calculate the protein abundance. The top three peptides with highest abundance are used to calculate the protein abundance.

- **-10lgP**: The -10lg of a p-value represents the likelihood that the observed change between conditions is caused by random chance. The peptides are first separated into groups based on similar quality score. Significance is then calculated for each quality score group, or bin, using a two-tail T-test that assumes log normal distribution but does not assume equal variance. The -10log p-value is then calculated from this result and then displayed in the significance column.

- **Average Area**: Each peptide feature associated with the peptide has its own area under the XIC curve. The average area of is the average area of all the features associated with the peptide.

- **Condition Profile**: The relative peptide abundance for each condition compared to the reference condition among the samples is depicted as a heat map. Hold the cursor on a profile to view the samples, the areas, and the ratios with respect to the reference condition.

- **Condition Ratio**: The ratio columns show the relative peptide abundance for each condition with respect to the reference condition for all groups. A peptide's abundance in each condition is the sum of its supporting peptide feature abundances within the peptide filters. Then the peptide abundance in each condition is compared to the reference condition to obtain the peptide condition ratio in each sample separately. The peptide condition ratio of a group is computed by calculating the median from the peptide condition ratios of the samples in that group.

- **#Vector**: This shows the number of quantifiable feature vectors of a peptide. A feature vector is a full set of peptide features used to calculate a ratio.

- **Start**: This shows the protein position of the first residue of the peptide.

- **End**: This shows the protein position of the last residue (inclusive) of the peptide.

- **PTM**: The types and the numbers of modifications present in the peptide shown using color-coded icons. Scroll over each icon to see the modification name and mass.

### Table: Quantifiable Support Peptides for Selected Protein

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Used</th>
<th>-10lgP</th>
<th>Avg.Area</th>
<th>Condition 1: Ratio Profile</th>
<th>Condition 2: Ratio Profile</th>
<th>Condition 3: Ratio Profile</th>
<th>Group 1: Condition 2 Ratio</th>
<th>Group 1: Condition 3 Ratio</th>
<th>#Vector</th>
<th>Start</th>
<th>End</th>
<th>PTM</th>
</tr>
</thead>
<tbody>
<tr>
<td>AEK...</td>
<td>&lt;27.77</td>
<td>1.2664</td>
<td>0.99</td>
<td>&lt;27.77</td>
<td>0.32</td>
<td>2</td>
<td>122</td>
<td>134</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>C115...</td>
<td>&lt;17.52</td>
<td>2.9765</td>
<td>0.85</td>
<td>&lt;17.52</td>
<td>0.70</td>
<td>1</td>
<td>104</td>
<td>134</td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>C115...</td>
<td>&lt;35.05</td>
<td>4.3109</td>
<td>0.83</td>
<td>&lt;35.05</td>
<td>0.68</td>
<td>3</td>
<td>104</td>
<td>134</td>
<td></td>
<td></td>
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<tr>
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<td>&lt;29.23</td>
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<td>&lt;29.23</td>
<td>0.85</td>
<td>3</td>
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<td>134</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>DELN...</td>
<td>&lt;40.33</td>
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<td>0.83</td>
<td>&lt;40.33</td>
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<td></td>
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<tr>
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<td>6.0905</td>
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<td>&lt;19.09</td>
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<td>53</td>
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<td>&lt;64.03</td>
<td>0.60</td>
<td>18</td>
<td>55</td>
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<tr>
<td>GPG...</td>
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<td>&lt;33.54</td>
<td>0.34</td>
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<td>0.74</td>
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<tr>
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<tr>
<td>IV...</td>
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<td>0.72</td>
<td>&lt;56.11</td>
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<td>113</td>
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<td></td>
<td></td>
</tr>
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<td>IV...</td>
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<td>3.935</td>
<td>0.69</td>
<td>&lt;36.41</td>
<td>0.55</td>
<td>3</td>
<td>111</td>
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<td></td>
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</tr>
<tr>
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<td>2.2502</td>
<td>0.30</td>
<td>&lt;21.10</td>
<td>0.86</td>
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<td>157</td>
<td>169</td>
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<td>0.86</td>
<td>&lt;31.94</td>
<td>0.63</td>
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</tr>
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<td>LA...</td>
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<td>&lt;40.55</td>
<td>0.57</td>
<td>9</td>
<td>158</td>
<td>169</td>
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<td></td>
</tr>
<tr>
<td>LA...</td>
<td>&lt;32.53</td>
<td>1.2187</td>
<td>0.70</td>
<td>&lt;32.53</td>
<td>0.80</td>
<td>9</td>
<td>158</td>
<td>169</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LA...</td>
<td>&lt;59.57</td>
<td>1.2195</td>
<td>0.73</td>
<td>&lt;59.57</td>
<td>0.61</td>
<td>1</td>
<td>113</td>
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<td></td>
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</tr>
<tr>
<td>MA...</td>
<td>&lt;22.47</td>
<td>1.2402</td>
<td>0.70</td>
<td>&lt;22.47</td>
<td>0.41</td>
<td>5</td>
<td>1</td>
<td>23</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

10.5 SILAC result - Peptides View

Quantified proteins are displayed in the Peptide View along with their detailed information.
10.5.1 Peptides View - Peptide Table

The peptides are grouped together based on quantifiable feature vectors with the same sequence. When there are more than 1000 peptides, the table is split into multiple pages.

The view is similar to the support peptide table described in the previous section Section 10.4.3.4 Proteins View - Supporting Peptides. The Peptide table entries may be filtered by using the Peptide View Filters on top of the peptide table. To reiterate, the columns are (with two extra columns Quality score and Accession):

- **Peptide**: The amino acid sequence of the peptide, as determined in PEAKS Database or Spectral Library search. A modified residue is followed by a pair of parentheses enclosing the modification mass. All non-unique peptides and those not passing the filter will be excluded automatically.

- **-10lgP**: The -10lg of a p-value represents the likelihood that the observed change between conditions is caused by random chance. The peptides are first separated into groups based on similar quality score. Significance is then calculated for each quality score group, or bin, using a two-tail T-test that assumes log normal distribution but does not assume equal variance. The -10log p-value is then calculated from this result and then displayed in the significance column.

- **Average Area**: Each peptide feature associated with the peptide has its own area under the XIC curve. The average area of is the average area of all the features associated with the peptide.
### Quality Score
The quality score represents the predicted reproducibility of the quantification result. Factors that affect the quality score include m/z difference, RT difference, XIC shape similarity, and the feature intensities.

### Condition Profile
The relative peptide abundance for each condition compared to the reference condition among the samples is depicted as a heat map. Hold the cursor on a profile to view the samples, the areas, and the ratios with respect to the reference condition.

### Condition Ratio
The ratio columns show the relative peptide abundance for each condition with respect to the reference condition for all groups. A peptide's abundance in each condition is the sum of its supporting peptide feature abundances within the peptide filters. Then the peptide abundance in each condition is compared to the reference condition to obtain the peptide condition ratio in each sample separately. The peptide condition ratio of a group is computed by calculating the median from the peptide condition ratios of the samples in that group.

### #Vector
This shows the number of quantifiable feature vectors of a peptide. A feature vector is a full set of peptide features used to calculate a ratio.

### Start
This shows the protein position of the first residue of the peptide.

### End
This shows the protein position of the last residue (inclusive) of the peptide.

### PTM
The types and the numbers of modifications present in the peptide shown using color-coded icons. Scroll over each icon to see the modification name and mass.

### Accession
The proteins supported by the peptide, separated by ',', if the peptide supports multiple proteins. If none of the proteins that the peptide supports pass the protein filters, the accession field will be empty.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

### 10.5.2 Peptide Table controls
The menu above the peptide table provides options to scroll through the table of peptides, filter the table, and search for specific entries.

#### 10.5.2.1 Peptide Table controls - Pagination
For easy viewing and to improve performance, the peptide table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.

In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

#### 10.5.2.2 Peptide Table controls - View Filter
Click on the **Peptide View Filter** button to open the Peptide View Filter which is used to filter the Peptide table. The following are the filter options provided:
- **Peptide sequence contains**: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains for further filtering if necessary.

- **Peptide score (-10lgP)**: Filters out all peptides with -10lgP scores below this threshold.

- **Peptide length**: Filters for peptides of length within this min/max threshold.

- **Protein Accession Contains**: Enter a peptide accession, only peptides that contain that accession will be displayed in the peptide table.

- **Show Transferred IDs Only**: Only display peptides where match between runs was used to include at least one peptide feature in the result.

- **Ratio Filter**: Only peptides where the ratio for the selected group and condition fall within the specified range will be displayed in the peptide table.

- **PTM contains**: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text they will all be included in the table.

  Note: In all filters, multiple entries can be separated by a semi-colon.

Click on the Reset button to reset all filters to their default setting.

Click on OK to apply the filters to the Peptides table.

Click on Cancel to discard the changes in the View Filter.
10.5.2.3 Peptide Table controls - Search function

On the top right of menu, there is a search function with 1 option. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- **Seq contains:** Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered. Additionally, since the peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”

10.5.3 Feature vector selection breadcrumb

There is a sample and feature selection breadcrumb which allows you to switch between features associated with the peptide. Click on the Sample in the breadcrumb to reveal other samples and the features in that sample related to the current peptide selection. From here you can select other samples, fractions, or features.

By clicking on the fraction level, all fractions in this sample are revealed as well as the features in each fraction associated with this peptide. From here you can select other fractions and features to view.

Clicking on the feature vector level will show all features vectors in the currently selected fraction.

Switch between Conditions with the last portion of the breadcrumb.

All other components in the Peptide tab interface will update according to the currently selected Condition.
10.5.4 Protein jump button

Click on the button to jump to a protein that contains the currently selected PSM. If there are more than 1 protein, a dropdown is revealed which allows the user to select which protein to jump to. If a protein is filtered out due to a protein summary filter or summary page protein filter, then the option to jump will be grayed out.

10.5.5 Annotated Spectrum Chart

For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

10.5.5.1 Annotated Spectrum Chart - Chart Navigation

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.

Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.

Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.

See the whole spectrum: Double click in the spectrum or click the 1:1 button.

10.5.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings

Click on the button to show the spectrum annotation settings.
The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places:** Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation:** Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated:** Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation:** Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info:** Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity:** Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

### 10.5.5.3 Annotated Spectrum Chart - Additional Chart Controls

Click on  or  to toggle between absolute and percentage intensity values in the Y axis.

Click on  to reset the zoom level of the annotated spectrum chart to the default.

Click on  to zoom into the X axis by a factor of 2.

Click on  to zoom into the Y axis by a factor of 2.
The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

10.5.6 MS1 Spectrum

This view provides an MS1 spectrum zoomed in on the reporter ions in the experiment. This is an interactive spectrum, enabling zooming in and out using the scroll wheel of the mouse. Double-clicking will zoom in on an area of interest. Clicking and dragging will slide the spectrum to the left or right.

10.5.7 eXtracted Ion Chromatogram (XIC) chart

This map presents the changes in feature intensity over the retention time range of the peaks for all labels of the selected peptide. Additionally, the coloured isotopic distribution plot - M0, M+1, M+2, etc. - of each label is included, which shows the relative composition of each label.
10.5.8 LC/MS Snapshot

This LC/MS screen is zoomed into the features that correspond to the isotopic labels, e.g. “Light” or “Heavy”. The m/z between the markers for these features should equate to the mass difference of the two labels in this sample.
Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a m/z RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

10.6 SILAC Exporting

In Project View, double-click on to open the Export node.

10.6.1 SILAC - Exporting - comma separated values (CSV) format

There are several options for exporting the PEAKS Precursor Ion Quantification results in CSV format:

- **Proteins CSV**: The list of protein quantifications, filtered by the protein filters in the Summary view and will be saved to proteins.csv.
- **Peptide CSV**: A complete list of all quantified peptides that pass the peptide feature filters set on the “Summary” page and their associated details.
- **Support peptides CSV**: These are the supporting peptides of the quantified proteins and their associated details.
• **Feature Vector CSV**: All predicted peptide features and their associated quantification information. Both identified and unidentified peptide features are included. To include unidentified peptide features, set the Peptide Id Count equal to 0 in the Peptide feature filter.

10.6.2 SILAC - Exporting - HTML format

**Export in HTML**: This will generate single or multiple HTML reports in the specified location.

The following exporting options are available:

- **Summary view**: The “Summary” view page will be saved as a summary.html file in HTML format in the specified location.
- **Protein coverage**: The coverage pane will be saved for each protein.
- **Supporting peptides**: A list of supporting peptides will be saved for each protein.
- **Best unique PSM**: The best unique PSM will be saved for each protein.
- **Export peptide list in html**: creates a separate HTML-containing peptide identifications.

An individual protein will have its own HTML output file, where the corresponding protein coverage, the supporting peptides, and the best unique PSM are gathered.
11. Glycan Search

PEAKS Glycan is an add-on module for PEAKS Studio that provides a unique solution for in-depth Glycoproteomic analysis. PEAKS Glycan module utilises a glycopeptide-based approach to profile the glycoproteins in your sample using LC-MS/MS data. As part of the PEAKS Studio platform, PEAKS Glycan module presents a refined database search and newly developed algorithms to facilitate the identification and characterisation of both N- and O-linked glycans. Glycan profiling is done at specific sites within a protein (positional profiling) and compares glycopeptide abundances across samples by label-free quantification. Currently, it supports ThermoFisher and Bruker instruments and includes an easy-to-use project wizard for workflow and analysis parameter selection that will be familiar to all PEAKS users.

Glycosylation, the attachment of sugar moieties or glycans to organic molecules, is one of the most common post-translation modifications (PTM) and plays a critical function in a wide range of biological processes. Glycans are the most complex and heterogeneous class of carbohydrate-based structures due to their non-template-driven biosynthetic process. They can be linear or branched, and they are attached to the side chain of amino acids within glycoproteins. The glycans in glycoproteins can be categorised as being N-linked, glycans bound to nitrogen atom of asparagine (N), or O-linked, glycans bound to the oxygen of serine (S) or threonine (T) side chains. Importantly, glycosylation can affect a protein's structure, stability, and function, and is important for physiological and pathological functions.

Glycoproteomics involves identifying glycans and the associated glycosylation sites within proteins across the proteome of a given cell or tissue. At the protein level, amino acid sequences, or individual amino acids, are used to predict glycosylation sites. Fragmentation data from MS/MS spectra are used to identify these sites and the attached N- or O-linked glycan structures. In the past, LC-MS/MS-based glycoproteomics was challenged by complexity of structures present in a sample, resulting in an extensive search space consisting of all potential peptide sequences, glycans' compositions and structures, in addition to other PTMs. PEAKS Glycan module was designed to handle these challenges of mass spectrometric glycoproteome analysis, and as a comprehensive data analysis tool it provides a highly sensitive and accurate glycoproteomics software solution.

11.1 Glycan Search Analysis Workflow and Setting

Select the PEAKS Search workflow in the Workflow Selection step.

The following steps are similar to the PEAKS DB workflow (as in section 6.1 Database Search Workflow and Parameter Settings) and is reiterated below. There are additional Glycan Option parameters for Glycan Search.

Click on Data Refine to proceed through the workflow.
The following parameter settings are provided:

**Precursor Mass Error Tolerance:** The maximum mass difference between the peptide and the monoisotopic mass of the precursor ion.

**Precursor Mass Tolerance unit:** PEAKS generates peptides within the precursor mass error tolerance. Precursor mass error tolerance can be specified in either Daltons or ppm.

**Fragment Mass Error Tolerance:** PEAKS uses this value when scoring peptide sequences. PEAKS Studio considers a fragment ion to be matched if the calculated m/z is within the specified error tolerance. Units are Daltons.

**Glycan Fragment ion:** This enables the selection of a fragment ion m/z tolerance for Glycans. Glycan Search uses this value when scoring glycan peptides. PEAKS considers that a fragment ion is matched if the calculated m/z is within the tolerance. Glycan Fragment ion mass tolerance can be specified in either Da or ppm.

**Ion Mobility Tolerance (1/k0):** This tolerance can only be set for timsTOF instruments. This setting is the allowable maximum shift in mean 1/k0 of an identified peptide between different runs.

Both precursor and fragment error tolerance parameters need to be set consistently with the mass accuracy of the instrument.
**Enzyme Settings:** PEAKS Studio digests the protein database *in silico* to generate peptide candidates. It is necessary to specify the enzyme for protein digestion from the Enzyme drop-down menu.

**Note:** When the selected dataset is digested with different enzymes, "Specified by each sample" allows samples to be analyzed separately using their respective enzymes specified during project creation.

**Note:** "None" is a special enzyme allowing non-specific cleavage at both ends of the peptide. It is the recommended option when no digestion enzyme was used or when the digestion enzyme exhibits a high degree of non-specificity.

**Missed Cleavages:** This specifies the number of missed cleavage sites allowed in a peptide.

**Digest Mode:** This specifies the type of in silico cleavages allowed at peptide termini based on the method used for protein digestion. If "Specific" is selected, the specificity of the selected enzyme is strictly enforced at both termini of a peptide. "Semispecific" allows one terminus to disobey the enzyme specificity rules.

**Note:** If the enzyme is specified as "None", then no matter which mode is selected for "Digest Mode", the "None" enzyme digest rule will be applied.

**PTM Settings:** Click “Set PTM” to open the PTM Options dialog and specify the fixed PTMs and a few common variable PTMs expected in the experiment. Highlight PTMs and click on the button to place them as either Fixed or Variable PTM. From there, highlighted PTMs can be removed by clicking the “Remove” button. Clicking “Remove All” will reset the PTM options. Highlighting and clicking “Switch Type” will move the Fixed PTM in into the Variable PTM list and vice-versa. Click on “New” to create a new custom PTM. While a PTM is selected, click on “View” to see more details.

**Note:** Multiple PTMs can be highlighted by holding down the CTRL key or SHIFT key on the keyboard.

**Note:** For a specific residue, only one fixed modification is allowed.

Once the PTMs are selected, the maximum number of variable PTMs per peptide can be specified and a value of less than 4 is recommended.

**Note:** The use of variable modifications increases the size of the computational search space for the de novo sequencing algorithm. It is recommended not to select too many variable modifications in PEAKS de novo. See PEAKS PTM for alternate search options.

**Target Database:** Select a protein sequence database for the search. Select one from the list of databases that have been configured in PEAKS and set the taxonomy by clicking on , if applicable. To configure a new database, click on next to the Target Database dropdown to open the Database Configuration dialog.

**Glycan Option:**

Glycan Search can be selected based on the glycan linkage nature: N-Link, O-Link, or N-Link + O-Link Glycan Search. N-glycans are bound to Asparagine (N), while O-glycans are bound to Serine (S) / Threonine (T). To run N-link followed by O-link, check both checkboxes next to the Glycan Database selection.
### Enrichment Data
The Enrichment Data checkbox specifies whether the data selected in the project is considered to be enriched data. Enriched data will search the whole protein database regardless of the size of the database. Also, if there are <500 entries in the database, the whole database is used for searching glycan peptides as well. If Enrichment Data checkbox is unchecked and the database is greater than 500 entries, then the Database search result set of proteins is used for searching for glycan peptides. Setting this option may drastically increase computing time for large databases.

### Search Glycopeptide Only
Reports only glycopeptides and does not report non-glycopeptide results from database search. This option is only available if the previous parameter “Enrichment Data” is selected.

### NH4, Na, K
Select these to search for NH4, Na, or K adducts respectively.

### Maximum Adduct Per Peptide
Sets the limit to the number of adducts on each glycopeptide.

### Maximum Fucose
Sets the limit to the number of Fucose molecules for each glycan. O-link may have two glycans.

### Composition or Structure
If selecting Composition based, Glycan PSMs with the same composition and different structures are merged and reported as one glycopeptide. If selecting Structure, Glycan PSMs with the same composition but different structures are reported as separate glycopeptides.

Click on “Report” to proceed to the final step. Report filters can be set here which will filter the final result.
These filters are provided:

**Score Filter**: this section is a mandatory section to filter out some results under the user-defined filter. Any changes to the filter will create a new analysis with old parameters and update the results and statistics.

For each of N-link and O-link glycan search (if enabled), there is a Peptide Score and Glycan Score to be set.

**Peptide score**: The PEAKS peptide score (-10lgP) is calculated for every peptide-spectrum match (PSM) reported by PEAKS DB, PEAKS PTM, and SPIDER. The score is derived from the p-value that indicates the statistical significance of the peptide-spectrum match. A peptide may be matched to many spectra, resulting in multiple PSMs. In that case, the peptide’s score is calculated as the maximum among all PSM scores. For details of the scoring algorithm, please refer to the publication, "PEAKS DB: De Novo sequencing assisted database search for sensitive and accurate peptide identification", Mol Cell Proteomics, 2011 Dec 20. A minimum requirement can be set and all identifications must pass this filter.

**Peptide FDR (%)**: The Peptides can also be set to use the false discovery rate. Peptides are scored based on their top PSM and using a “decoy-fusion” approach, the result is reported at that FDR.

**Note**: Decoy fusion is an enhanced target-decoy method for result validation with FDR. Decoy fusion appends a decoy sequence to each protein as the "negative control" for the search. See BSI's web tutorial (http://www.bioinfor.com/fdr-tutorial/) for more details.

**Glycan Score**: Glycopeptides above this score threshold will remain in the glycopeptide list.
**Glycan Score FDR (%):** Similar to Peptide FDR, the glycopeptides above the false discovery rate calculated for the whole set of glycopeptides will be reported.

**Protein -10lgP:** Proteins above the -10lgP (a combination of Peptide score and Glycan score) filter will be reported in the Protein table. This is an initial filter set that can be adjusted in the Summary page.

**Unique Peptides:** Proteins with this minimum # of unique peptides will be reported in the Protein table. This is an initial filter set that can be adjusted in the Summary page.

In the Report step, there also is the option to specify an Analysis Name (which can be later renamed) and/or decide to save the Workflow to quickly re-use in future analyses.

Click Finish to start running the project or analysis.

### 11.2 Understanding Glycan Search results

The Glycan Search result is the main result view. To open the Glycan Search result, double-click on the **Glycan Search** node in the Project View under any Analysis.

The Glycan Search view comes with 6 main tabs for result visualisation.

- Summary View
- Proteins View
- Glycopeptide View
- Peptide View
- Feature View
- Partial View

### 11.3 Glycan Search Summary View

The Summary View is selected by default when a Glycan Search result is opened. The ‘Summary View’ reports key statistics in table format as an overview of the result. Protein filters and Notes about the experiment can be set here. To select the Summary View, click on the ‘Summary’ tab.
1. Notes

2. Result Statistics

Table 1. Statistics of data.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Lipid-MS</th>
<th>Refined</th>
<th>GlycoID</th>
<th>Identified</th>
<th>PSMs</th>
<th>Peptides</th>
<th>Sequences</th>
<th>Proteins</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>7982</td>
<td>41781</td>
<td>41781</td>
<td>20305</td>
<td>39079</td>
<td>927</td>
<td>3019</td>
<td>1923</td>
</tr>
<tr>
<td>Sample 1</td>
<td>3905</td>
<td>19709</td>
<td>19709</td>
<td>6945</td>
<td>43114</td>
<td>375</td>
<td>1101</td>
<td>1865</td>
</tr>
<tr>
<td>Sample 2</td>
<td>4072</td>
<td>22072</td>
<td>22072</td>
<td>13360</td>
<td>45565</td>
<td>552</td>
<td>1518</td>
<td>2018</td>
</tr>
</tbody>
</table>

Figure 1. Precursor mass error of Non-Glycopeptide-spectrum matches (PSM) in filtered result. (a) Distribution of precursor mass error in ppm; (b) Scatterplot of precursor m/z versus precursor mass error in ppm.

Figure 2. Precursor mass error of Glycopeptide-spectrum matches (PSM) in filtered result. (a) Distribution of precursor mass error in ppm; (b) Scatterplot of precursor m/z versus precursor mass error in ppm.

11.3.1 Summary View - Notes

1. Notes

This is an example EXPERIMENT 1

This is an example EXPERIMENT 1
Click on the Notes button in the top bar to open the Note Entry editor. In the header of the report, information such as a description of the experiment can be entered. Lots of formatting options are provided to give the user flexibility on how they want to show this information. Clicking OK will save the notes into the result, which will remain with the project.

11.3.2 Summary View - Protein Filters
Protein Filters are provided in the Summary View.

The Protein score (-10lgP) is calculated as the weighted sum of the -10lgP scores of the protein’s supporting peptides. After removing any redundant peptides, the supporting peptides are sorted by -10lgP scores in descending order, and the k-th ranked peptide contributes to the weighted sum with a weight of 1/k.

The unique peptides filter is based on the number of unique supporting peptides for the protein.

Changing the -10lgP filter or unique peptides filter and clicking the red ‘Apply’ button will affect the Protein table in the Protein tab, and all proteins below the thresholds will be filtered out. This will also affect jumping functions from other tables (such as jumping from the Glycan table and Peptide table) as that protein is no longer accessible. The protein table exports will also be affected accordingly.

11.3.3 Summary View - Statistics of Data
2. Result Statistics

Table 1. Statistics of data.

<table>
<thead>
<tr>
<th></th>
<th>MS1</th>
<th>MS/MS</th>
<th>#Refined</th>
<th>Glycan</th>
<th>#Features</th>
<th>Identified GlycP</th>
<th>GlycPSMs</th>
<th>#PSMs</th>
<th>#Scans</th>
<th>#Features</th>
<th>Identified Peptides</th>
<th>#Peptides</th>
<th>#Sequences</th>
<th>#Proteins</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total</td>
<td>7982</td>
<td>41781</td>
<td>41781</td>
<td>20365</td>
<td>9079</td>
<td>927</td>
<td>3019</td>
<td>3923</td>
<td>1135</td>
<td>479</td>
<td>460</td>
<td>513</td>
<td>77</td>
<td>124</td>
</tr>
<tr>
<td>Sample 1</td>
<td>1065</td>
<td>19709</td>
<td>19709</td>
<td>6945</td>
<td>43514</td>
<td>375</td>
<td>1581</td>
<td>1865</td>
<td>478</td>
<td>245</td>
<td>328</td>
<td>359</td>
<td>59</td>
<td>93</td>
</tr>
<tr>
<td>Sample 2</td>
<td>2017</td>
<td>22072</td>
<td>22072</td>
<td>13360</td>
<td>45565</td>
<td>552</td>
<td>1518</td>
<td>2058</td>
<td>657</td>
<td>326</td>
<td>378</td>
<td>429</td>
<td>72</td>
<td>119</td>
</tr>
</tbody>
</table>

Table 1 Statistics of Data tabulates MS1 and MS2 information, as well as identification in the peptide and protein level. This table in the “Summary” view provides a condensed statistical information for the analysis.

For timsTOF Instruments, the statistics table has been modified to show key information about the #Frames and #Precursors.

11.3.4 Summary View - Non-glycopeptide Precursor mass error distribution
Figure 1. plots the distribution of non-glycopeptide PSMs in the filtered result. Figure 1. (a) shows the number of PSMs for each mass error in ppm. Figure 1. (b) shows the scatterplot of mass error in ppm across the m/z range
11.3.5 Summary View - Glycopeptide Precursor mass error distribution

Figure 2. plots the distribution of glycopeptide PSMs in the filtered result. Figure 2. (a) shows the number of PSMs for each mass error in ppm. Figure 2. (b) shows the scatterplot of mass error in ppm across the m/z range.

Figure 1. Precursor mass error of Non-Glycopeptide-spectrum matches (PSM) in filtered result. (a) Distribution of precursor mass error in ppm. (b) Scatterplot of precursor m/z versus precursor mass error in ppm.
11.3.6 Summary View - Number of Glycosylation Sites (top 20)

Table 2. Number of glycosylation sites (top 20).

<table>
<thead>
<tr>
<th>Glycan Moiety</th>
<th>Structure</th>
<th>Type</th>
<th>Sample 1</th>
<th>Sample 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>(HexNAC)4(Hex)5(NeuAc)2</td>
<td></td>
<td>N-Link</td>
<td>32</td>
<td>39</td>
</tr>
<tr>
<td>(HexNAC)4(Hex)5(NeuAc)1</td>
<td></td>
<td>N-Link</td>
<td>10</td>
<td>16</td>
</tr>
<tr>
<td>(HexNAC)5(Hex)6(NeuAc)3</td>
<td></td>
<td>N-Link</td>
<td>10</td>
<td>14</td>
</tr>
<tr>
<td>(HexNAC)1</td>
<td></td>
<td>O-Link</td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td>(HexNAC)5(Hex)6(NeuAc)2</td>
<td></td>
<td>N-Link</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>(HexNAC)4(Hex)4(Fuc)2(NeuAc)2</td>
<td></td>
<td>N-Link</td>
<td>3</td>
<td>10</td>
</tr>
<tr>
<td>(HexNAC)1(Hex)1(NeuAc)1</td>
<td></td>
<td>O-Link</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td>(HexNAC)4(Hex)4(NeuAc)2</td>
<td></td>
<td>N-Link</td>
<td>9</td>
<td>2</td>
</tr>
<tr>
<td>(Hex)1</td>
<td></td>
<td>O-Link</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>(HexNAC)4(Hex)4</td>
<td></td>
<td>N-Link</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>(HexNAC)4(Hex)5</td>
<td></td>
<td>N-Link</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>(HexNAC)2(Hex)5</td>
<td></td>
<td>N-Link</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>(HexNAC)4(Hex)2(Fuc)2(NeuAc)1</td>
<td></td>
<td>O-Link</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>(HexNAC)5(Hex)5(NeuAc)1</td>
<td></td>
<td>N-Link</td>
<td>3</td>
<td>6</td>
</tr>
<tr>
<td>(HexNAC)4(Hex)5(Fuc)1(NeuAc)2</td>
<td></td>
<td>N-Link</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>(HexNAC)5(Hex)5(Fuc)1(NeuAc)1</td>
<td></td>
<td>N-Link</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>(HexNAC)5(Hex)5(NeuAc)1</td>
<td></td>
<td>O-Link</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>(HexNAC)4(Hex)5(Fuc)1(NeuAc)1</td>
<td></td>
<td>N-Link</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td>(HexNAC)5(Hex)5(NeuAc)2</td>
<td></td>
<td>N-Link</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>(HexNAC)3(Hex)4(NeuAc)1</td>
<td></td>
<td>N-Link</td>
<td>2</td>
<td>6</td>
</tr>
</tbody>
</table>

Table 2 Number of Glycosylation Sites (top 20) shows the 20 most abundant glycans found in the result. It will list both N-linked and O-linked Glycans for combined N-link and O-linked Glycan searches. For every position in proteins in the protein table, if the glycan is found at that position it will be counted as a Glycosylation Site in this table. For the full Glycosylation Sites table, use the Glycosylation Sites CSV export from the Export node.

Note: If running with Glycan Composition only (instead of Structure based) in the Glycan Search workflow parameters, this table will not display the structure.

11.3.7 Summary View - Result Filtration Parameters.

Table 3 Result Filtration Parameters simply states the protein filters that are currently set in the Summary page. If the Summary page is used in a report, the protein filters set for that analysis report can be obtained here.

Table 3. Result filtration parameters.

| Protein -10lgP | ≥15 |
| Proteins unique peptides | ≥1 |
11.4 Glycan Proteins View

The ‘Proteins View’ lists protein identifications that have been filtered by the current settings in "Summary" view. It also visualizes the protein sequence coverage for identified proteins. The view has four main components:

- **Protein table:** This lists protein identifications that have been filtered by the current settings in the Summary View and the “Protein View Filters”.

- **Coverage:** This visualizes the protein sequence coverage of the selected protein, showing the mapping of support peptides and de novo tags to the protein sequence.

- **Peptides:** This lists supporting peptides mapped to the selected protein.

- **Glycopeptides:** This lists supporting glycan peptides mapped to the selected protein.

### 11.4.1 Protein Table

Each row in the table represents a group of proteins that are supported by a common set of peptides. A dark blue node indicates that the group has multiple proteins. To expand the group, click the "→" button at the left. To collapse the group, click the "↓" button.

**Note:** A supporting peptide is considered highly confident if it has a peptide -10lgP score above the set threshold.
• **Accession:** The accession number of the protein as seen in the FASTA database.

• **Cluster:** This is a unique number to identify which protein group a protein belongs to. Those with the same number will be grouped together.

• **Top:** This column is provided to indicate if the protein is considered a top protein.

• **-10lgP:** The protein confidence score.

• **Coverage(%)** The percentage of the protein sequence that is covered by supporting peptides. The coverage is visualized by a colored bar. Purple blocks indicate the parts of the sequence covered by glycan peptides (either N-link or O-link). Blue blocks indicate the parts covered by high-confidence native peptides. This is the total coverage including results from all samples.

• **#Peptides:** The number of high-confidence supporting peptides (both glyco peptides and native peptides).

• **#Unique:** The number of high-confidence supporting peptides (both glyco peptides and native peptides) that are mapped to only one protein group. Unique peptides with same sequence but different modifications are only counted once in this number.

• **PTM:** The identified modifications displayed with color-coded icons. Mouse over the cell to see more details on which PTMs are included in the protein.

• **Glycan:** The identified glycans displayed with color-coded icons. Mouse over the cell to see more details on which glycans are included in the protein.

• **Glyco Sites:** The sites in the protein with glycans are listed here.

• **Avg. Mass:** The protein mass calculated using the average mass.

• **Description:** The protein’s header information as seen in the FASTA database.

**Note:** For #Peptides and #Unique, two peptides with the same starting and ending positions in the protein are counted as one, regardless of their PTM forms. This is to follow the MCP (Molecular & Cellular Proteomics) guidelines.

Any columns in the protein table can be reorganized and shuffled. Most columns are able to be sorted.

11.4.2 Protein View Filters

Click on the [Protein View Filter] button to access the Protein View Filters.
Show All or Top proteins in each group: Adjust the protein list based on Protein grouping. All is selected by default. Proteins are grouped based on parsimony.

- **All**: Show all proteins in each group. Proteins are grouped together if they are supported by the same set or a sub-set of the top protein in the group. If a protein is supported by a peptide not supporting the top protein it will be added to a new group.

- **Top**: Show only the top proteins in each group. These proteins are supported by the most unique peptides in the group. Proteins in the group that share a subset of the unique peptides that support the top protein (sub-proteins) will not be displayed.

Show Proteins checkboxes: Checkboxes for Glycoproteins and Non-Glycoproteins are provided. If Glycoproteins is checked, then proteins with supporting glycan peptides will show in the Protein table. If Non-Glycoproteins proteins is checked, then proteins with supporting native peptides will show in the Protein table. If neither are checked, the protein table would be empty.

Protein accession/name contains: Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.

Protein description contains: Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

Protein sample area: Filter proteins based on total protein area found in each sample. Sample areas are calculated using the total of all peptide features from unique supporting peptides. If a protein is identified and has areas from multiple samples, only one of the samples must pass this filter to be included.
**PTM contains:** Filter protein results based on the presence of selected PTMs. Proteins that are supported by peptides containing modifications that match the selected PTMs will be included. If multiple PTMs match the modification name, they will all be included in the table.

**Glycan contains:** Filter protein results by the presence of selected Glycans. Proteins that are supported by glycan peptides containing the selected glycans will be included. If multiple glycans match the modification name, they will all be included in the table.

**Note:** In all filters, multiple entries can be separated by a semi-colon.

Click on the **Reset** button to reset all filters to their defaults.

Click on the **OK** button to apply the changes.

Otherwise click on the **Cancel** button to cancel the changes and keep the previous settings.

### 11.4.3 Protein Optional Columns

Click on the **** button to show the optional columns dropdown to show/hide additional columns in the protein table.

- **Sample Coverage**
  - Coverage(%) Sample 1
  - Coverage(%) Sample 2

- **Sample Spectra Counts**
  - #Spec Sample 1
  - #Spec Sample 2

- **Sample Areas**
  - Area Sample 1
  - Area Sample 2

The options provided are sample level coverage, spectra counts, and areas.

Since there can be many samples in an analysis, all the sample level optional columns are not selected by default.

### 11.4.4 Protein Table Search Function

An easy to use search function is available for the protein table. The protein accession or description substring can be searched for and all search results can be navigated through with the Up and Down arrows.
11.4.5 Protein Coverage

The "Protein Coverage" view visually maps the supporting peptides and de novo tags to the protein selected in the Protein table. It also shows all identified sites with modifications or mutations to assist with protein characterization at the amino acid level. The "Protein Coverage" view is composed of three major components - Protein Sequence Coverage, Coverage Control Panel, and Protein Tools.

11.4.5.1 Protein Sequence Coverage

This area visualizes the coverage of the protein sequence.

Regions in the protein sequence that are covered by supporting peptides are displayed in bold font with a grey background. Confident modifications identified in supporting peptides are displayed as icons above the protein sequence. Modifications are represented by colored icons with the first letter of its modification name.

The supporting native peptides and glycan peptides are shown as colored bars under the protein sequence. Non-glycopeptides are blue coloured, N-linked peptides are purple coloured, and O-linked peptides are orange coloured.

Placing the cursor over a bar shows a tooltip with detailed information of the peptide. Left clicking on a bar shows the annotated spectrum from which the supporting peptide is identified.
Right clicking on a peptide bar shows additional options. “Show in spectrum” has the same behaviour as left clicking on the peptide bar which is to open the annotated spectrum chart. “Show in table” has the function to jump to either the supporting glycopeptides table or supporting non-glycopeptides table.

Clicking on the button will show the full screen view. Click on to return to the original view.

Click on the button reveals three more options for the protein coverage.

- Copy protein sequence: This will simply copy the protein sequence along with the header into the clipboard so that it can be easily pasted.
- Save image as: The image of the protein coverage can be exported with different sizes.
- Coverage statistics: Additional statistics about the coverage are shown here.

11.4.5.2 Glycan Profile

Clicking on will open the Glycan Profile.

The Glycan Profile table will show all Glycan Moiety at every Protein Position. By default, all Glycosylation sites, Glycan Moiety, and Samples are selected. Click on the dropdowns above the table to select specific options to show in the table.

When selecting a row in the Glycan Profile table, the top right spectrum chart and the bottom right XIC chart is updated with the selection.
The bottom left chart has two options. The Glycan XIC chart shows the combined XIC of glycans for each of the selected samples. Mouse drag in the chart area to zoom in. The Glycan Site Profiles shows a bar chart of the area distribution between samples of each glycan moiety for this protein, pictured below.

There is a radio button above the chart to switch between Percentage and Absolute area.

Right click on the chart area to export an image of the glycan profile chart.

On the top right there is the option to Save to Text Format. Click on the button to export the Glycan Profile table in CSV format.

**11.4.5.3 Glycan Pie Chart**

In the protein coverage, a purple coloured amino acid in the residue indicates a glycosylation site. Click on it to open the Glycan Pie Chart.

The Glycan Pie Chart shows the relative area of each glycan at that site. The top 7 glycans will be shown as slices in the pie chart and the rest will be categorized as ‘Other’.

An accompanying table is provided for the actual area values detected for each sample at this protein site. Mouse over the Glycan moiety column to see the cartoon structure of that moiety. Mouse over the glycan profile or group profile to see a pop-up with the ratios for each sample compared to the base sample.

Some of the columns are sortable where necessary and can be rearranged if needed.
11.4.5.4 Coverage Control Panel

This panel controls what to display in the protein sequence coverage view.
**Sequence Display Mode:** The protein sequence can be displayed in either "outline" or "coverage" mode.

"coverage" mode is the default mode to see the details of supporting glycan peptides and native peptides by viewing their coverage bars.

In “outline” mode, the protein sequence is displayed without supporting peptide bars. However, the sequence coverage at a specific position can be examined by left-clicking on a residue.
### AAs per line and 10AA gap:
Specify the number of amino acids per line and whether to show a gap for every 10 amino acids in the protein sequence.

### Sequence Display Option:
Select whether to view the protein sequence as Fasta or a specific enzyme used in the project. When a particular enzyme is selected, the protein sequence will be highlighted to emphasize expected peptides generated by the selected enzyme's cleavage specificity.

### Modification Table:
Also known as the PTM Table, specific glycans or PTMs can be deselected to hide from the protein coverage view. The # column indicates the number of instances that the modification is found in supporting peptides of this protein. Reminder that peptide bars are combined if they share the same start and end positions and have the same modifications.

### 11.4.6 Supporting Glycopeptides Table
The "Glyco Peptides" tab displays a table of supporting glycan peptides for the protein of interest. This table is similar to the Glycan Table in the "Glycan" view, except that "Accession" is excluded and the columns "(Checkbox)", "Unique", "Start", and "End" are included. Descriptions of extra columns are given below. See 11.5.1 Glycan Table for all descriptions.

<table>
<thead>
<tr>
<th>Coverage</th>
<th>Glycopeptides</th>
<th>Peptides</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peptide</td>
<td>Glycan Type</td>
<td>Glycan Type</td>
</tr>
<tr>
<td>DA00162.200000009-1.1</td>
<td>true</td>
<td>PAAPAP</td>
</tr>
<tr>
<td>2</td>
<td>true</td>
<td>0.00</td>
</tr>
<tr>
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<td>true</td>
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<td>12</td>
<td>true</td>
<td>0.00</td>
</tr>
<tr>
<td>13</td>
<td>true</td>
<td>0.00</td>
</tr>
<tr>
<td>14</td>
<td>true</td>
<td>0.00</td>
</tr>
<tr>
<td>15</td>
<td>true</td>
<td>0.00</td>
</tr>
<tr>
<td>16</td>
<td>true</td>
<td>0.00</td>
</tr>
<tr>
<td>17</td>
<td>true</td>
<td>0.00</td>
</tr>
<tr>
<td>18</td>
<td>true</td>
<td>0.00</td>
</tr>
<tr>
<td>19</td>
<td>true</td>
<td>0.00</td>
</tr>
<tr>
<td>20</td>
<td>true</td>
<td>0.00</td>
</tr>
</tbody>
</table>

#### Unique:
This shows whether the glycol peptide is a unique supporting glyco peptide to the protein group.

#### Start:
This shows the peptide's starting position in the protein.

#### End:
This shows the peptide's ending position (inclusive) in the protein.

Most columns are sortable and can be arranged where necessary.

### 11.4.7 Supporting Native Peptides Table
The Supporting Native Peptides table is also known as the "Peptides" tab and it displays a table of supporting native peptides for the protein of interest. This table is similar to the Peptide table in the "Peptide" view, except that "Accession" is excluded and the columns "(Checkbox)", "Unique", "Start", and "End" are included. Descriptions of extra columns are given below. See 11.6.1 Peptide Table for all descriptions.
<table>
<thead>
<tr>
<th>Peptide</th>
<th>Unique</th>
<th>Mass</th>
<th>Length</th>
<th>ppm</th>
<th>m/z</th>
<th>RT</th>
<th>Scan</th>
<th>FF</th>
<th>Score</th>
<th>Start</th>
<th>End</th>
<th>PFM</th>
<th>AScore</th>
</tr>
</thead>
<tbody>
<tr>
<td>KVFLI[S/L/C]-ST[D/Q]GNY[Q/G]W[Q/G]</td>
<td>true</td>
<td>120.66</td>
<td>44</td>
<td>0.9</td>
<td>1195.3483</td>
<td>65.44</td>
<td>1026.1</td>
<td>0</td>
<td>51</td>
<td>51</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RLNP[Q/G]DSD[Q/G]Y[Q/G]TSQLTIF</td>
<td>true</td>
<td>100.61</td>
<td>3167.5031</td>
<td>30</td>
<td>1.2</td>
<td>1056.8446</td>
<td>41.40</td>
<td>1166.6</td>
<td>5</td>
<td>52</td>
<td>81</td>
<td></td>
<td></td>
</tr>
<tr>
<td>K[Q/D]T[F/G]S[C/L]-ST[D/Q]GNY[Q/G]W[Q/G]</td>
<td>true</td>
<td>73.00</td>
<td>4770.5980</td>
<td>44</td>
<td>1.1</td>
<td>1195.3974</td>
<td>79.71</td>
<td>2149.3</td>
<td>0</td>
<td>8</td>
<td>51</td>
<td></td>
<td></td>
</tr>
<tr>
<td>K[Q/D]T[F/G]S[C/L]-ST[D/Q]GNY[Q/G]W[Q/G]</td>
<td>true</td>
<td>70.09</td>
<td>2337.1133</td>
<td>21</td>
<td>0.3</td>
<td>835.2910</td>
<td>33.89</td>
<td>816.2</td>
<td>0</td>
<td>307</td>
<td>327</td>
<td></td>
<td></td>
</tr>
<tr>
<td>KVFLI[S/L/C]-ST[D/Q]GNY[Q/G]W[Q/G]</td>
<td>true</td>
<td>59.14</td>
<td>4779.3200</td>
<td>44</td>
<td>1.6</td>
<td>1195.8488</td>
<td>85.24</td>
<td>2528.2</td>
<td>1</td>
<td>8</td>
<td>51</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Unique:** This shows whether the native peptide is a unique supporting native peptide to the protein group.

**Start:** This shows the peptide's starting position in the protein.

**End:** This shows the peptide's ending position (inclusive) in the protein.

Most columns are sortable and can be arranged where necessary.

### 11.5 Glycan Search Glycopeptide View

The "Glycopeptide View" displays the resulting glycan peptide identifications. The table on the top shows the details of each peptide identification. The bottom section provides additional information about the glycan-peptide spectrum matches for the peptide selected in the table.
### 11.5.1 Glycan Table

The Glycan Table shows the filtered glycopeptide identification results. Each row in the table is a glycopeptide identification represented by its highest-scoring PSM. When there are more than 1000 peptides, the table is split into multiple pages.

Note: If there are multiple PSMs matched to a peptide, the table displays the information for the top-scoring PSM. Peptides with the same sequence but different modifications are considered different and shown in different rows.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

The following list describes the contents in each of the default columns:

- **Peptide**: The amino acid sequence of the glycan peptide, as determined by the Glycan Search workflow. A modified residue is followed by a pair of parentheses enclosing the modification or glycan mass.

- **Glycan**: The Glycan moiety on this glycopeptide. For N-link results, there will always be 1. For O-link results, there could be more than 1.

---

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Glycan</th>
<th>Glycan Type</th>
<th>TMS</th>
<th>Glycan Score</th>
<th>Peptide Score</th>
<th>5-Score</th>
<th>Mass</th>
<th>Length</th>
<th>ppm</th>
<th>m/z</th>
<th>RT</th>
<th>Area Sample 1</th>
<th>Area Sample 2</th>
<th>Scan</th>
<th>Feature</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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<td></td>
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<td>2</td>
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<tr>
<td>3</td>
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<tr>
<td>4</td>
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<tr>
<td>5</td>
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<tr>
<td>6</td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

---

---

**To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.**
- **Glycan Type:** Simply states whether the glycan moiety is N-link or O-link. This column is particularly useful for N-Link + O-link search workflows.

- **-10lgP:** The peptide -10lgP score. The score indicates the scoring significance of a peptide-precursor spectrum match.

- **Glycan Score:** The normalized sum of ions log intensity, based on glycan fragment ion, peptide ion, and glycopeptide ion (Y ion).

- **Peptide Score:** Classifies glycan peptides with the normalized sum of ions log intensity, based on peptide fragment ions, and those peptide fragment ions with glycan. Different fragmentation type considers different peptide fragment ion:
  - CID and HCD considers b,y ion without glycan and b,y ion with at most 1 glycan node from root
  - ETD considers b,y,c,z,z' ion without glycan and b,y,c,z,z' ion with whole glycan from root
  - ETHCD considers b,y,c,z,z' ion without glycan, b,y,c,z,z' ion with 1 glycan from root, and b,y,c,z,z' ion with whole glycan from root

- **S-Score (%):** Measures the confidence of the top glycopeptide candidate as opposed to other candidates for the spectrum. For glycan candidates with the same composition and same peptide, the candidate is sorted by matched glycan y-ion count. $S$-Score = (most y-ion count - 2nd most y-ion count)/(most y-ion count). The higher the score the better. 100% indicates only 1 result, and it is the best match. 0% indicates that the top1 and top2 result are very similar, and we cannot confidently say the result is the best match.

- **Mass:** The monoisotopic mass of the glycan peptide.

- **Length:** The number of amino acids in the sequence backbone.

- **ppm:** The precursor mass error, calculated as $10^6 \times \frac{\text{precursor mass} - \text{theoretical peptide mass}}{\text{theoretical peptide mass}}$.

- **m/z:** The precursor mass-to-charge ratio.

- **RT:** The retention time (elution time) of the spectrum as recorded in the data.

- **Area per Sample:** The combined area for all features in that sample associated with this peptide.

- **Scan:** The scan number of the spectrum that matches the peptide sequence with the highest -10lgP.

- **#Feature:** For single sample analysis, shows number of features assigned to the peptide sequence from identification. For multiple samples, if peptide has feature vectors, it will show number of features used (area > 0) in all feature vectors. If not, it shows the number of features from identification.

- **Id Count:** Number of identifications in total.

- **Adduct:** Indicates whether this glycopeptide has an adduct and lists which adduct.

- **Isotopic Shift:** Indicates the isotopic mass shift.

- **Accession:** The accessions of proteins that contain this glycan peptide.

- **PTM:** The types and the numbers of modifications present in the peptide shown in color-coded icons.
• **A Score**: Localization score assigned to modifications on the peptide. It is the \(-10\) log of a p-value. In this case, the p-value is the probability that the modification occurs at the reported position compared to other possible positions. A \(-10\)lgP of 20 is equal to a p-value of 0.01. Scores are listed in the format AminoAcid, peptide position, modification, A Score. For example, serine at amino acid 3 is phosphorylated with an A Score of 54.01 will be reported as: S3 P 54.01.

11.5.2 Glycan Table controls
The menu above the peptide table provides options to scroll through the table of glycan peptides, filter the table, show optional columns and search for specific entries.

11.5.2.1 Glycan Table controls - Pagination
For easy viewing and to improve performance, the glycan table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.

In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

11.5.2.2 Glycan Table controls - View Filter
Click on the button to open the Glycan View Filter which is used to filter the Glycan table. The following are the filter options provided:
• **Peptide sequence contains**: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs or Glycans in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains or Glycan contains for further filtering if necessary.

• **Peptide sample area**: Filters based on total peptide area found in each sample. Sample areas are calculated using the total of all peptide features from all spectra that identified the glycan peptide. If the peptide is found in multiple samples, only one sample must pass this filter for the peptide to be included.

• **Peptide score (-10lgP)**: Filters out all peptides with -10lgP scores below this threshold.

• **Peptide length**: Filters for peptides of length within this min/max threshold.

• **PTM contains**: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text, they will all be included in the table.

• **Glycan contains**: Type the names of Glycans you would like to include. Peptides containing Glycans that match the text will be included. If multiple Glycans match the text, they will all be included in the table.

• **N-link Peptide and O-link Peptide checkboxes**: These checkboxes are only provided for N-link + O-link Glycan Search workflow. This is to select/deselect showing either type of Glycan in the table.
• **Note:** In all filters, multiple entries can be separated by a semi-colon.

Click on the Reset button to reset all filters to their default setting.

Click on OK to apply the filters to the Glycan table.

Click on Cancel to discard the changes in the View Filter.

### 11.5.2.3 Glycan Table controls - Optional Columns

Click on the button to open the optional columns dropdown. By default the sample area columns are displayed. Sample Feature count columns can also be shown. For projects with many samples, it might be useful to hide these optional columns and reduce the width of the glycan table.

### 11.5.2.4 Glycan Table controls - Search function

On the top right of menu, there is a search function with 4 options. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- **Scan =:** Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction.

- **m/z ≈:** Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit.

For example, searching for \( \text{m/z} \approx 580.8 \) will find all m/z between \([580.7, 580.9]\)

- **RT ≈:** Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/- 1 of the last significant digit set.
For example, searching for RT = 64 will find all m/z between [63, 65]

- **Seq contains**: Search for the sequence in the table. If specifying a sequence without glycans or PTMs, the results will include that sequence and any sequences with glycans or PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered.

Additionally, since the glycan peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”

11.5.3 Feature selection breadcrumb

There is a sample and feature selection breadcrumb which allows you to switch between features associated with the peptide. Click on the Sample in the breadcrumb to reveal other samples and the features in that sample related to the current glycan peptide selection. From here you can select other samples, fractions, or features.

By clicking on the fraction level, all fractions in this sample are revealed as well as the features in each fraction associated with this peptide. From here you can select other fractions and features to view.
Lastly, clicking on the feature level will show all features in the currently selected fraction. Features with Feature Id = 0 are known as fake features and are constructed to enhance identification.

11.5.4 PSM selection
Underneath the feature breadcrumb will shown the currently selected PSM. Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.

All other components in the Glycan tab interface will update according to the currently selected PSM.

For timsTOF instruments, Precursor ID is used rather than Scan number.

11.5.4.1 Protein jump button
Click on to jump to a protein that contains the currently selected glycan PSM. If there are more than 1 protein, a dropdown is revealed which allows the user to select which protein to jump to. If a protein is filtered out due to a protein summary filter or summary page protein filter, then the option to jump will be grayed out.

11.5.4.2 Show spectrum in Data View
Click on to open the Data Refine MS/MS tab (if not already opened) and jump to the corresponding selected Scan.
11.5.4.3 Show spectrum in LC/MS View

Click on [image] to open the Data Refine LC/MS tab (if not already opened) and jump to the corresponding selected Scan, highlighting the Glycopeptide MS/MS marker and the feature if present.

11.5.4.4 Show Raw Spectrum View

Click on [image] to show the raw spectrum chart of the currently selected Scan.

11.5.5 Annotated Spectrum Chart

For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

11.5.5.1 Annotated Spectrum Chart - Chart Navigation

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

- **Zoom to a m/z region:** Drag horizontally from the start m/z and to the end m/z with left mouse button.
- **Zoom in/out smoothly:** Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.
**Increase/Decrease peak intensity:** Place the cursor in the spectrum and scroll the mouse wheel.

**See the whole spectrum:** Double click in the spectrum or click the 1:1 button.

### 11.5.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings

Click on the button to show the spectrum annotation settings.

![Annotated Spectrum Chart](image)

The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places:** Select the number of decimal places that appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation:** Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated:** Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation:** Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.
**In Place Ion Info:** Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity:** Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

**11.5.3 Annotated Spectrum Chart - Additional Chart Controls**

Click on the dropdown to show annotation on Glycan ions, Peptide ions, or both.

<table>
<thead>
<tr>
<th>All</th>
<th>Glycan</th>
<th>Peptide</th>
</tr>
</thead>
</table>

Click on # or % to toggle between absolute and percentage intensity values in the Y axis.

Click on 1:1 to reset the zoom level of the annotated spectrum chart to the default.

Click on X to zoom into the X axis by a factor of 2.

Click on Y to zoom into the Y axis by a factor of 2.

The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

**11.5.6 Glycan Ion Match Table**

The Glycan Ion Match Table shows the theoretical m/z and observed m/z for each moiety of glycan ions matched in the spectrum. These columns are sortable and can be rearranged.

<table>
<thead>
<tr>
<th>Glycan Ion Match</th>
<th>Ion Match</th>
<th>Survey</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td>Theoretical m/z</td>
<td>Observed m/z</td>
</tr>
<tr>
<td>------------------</td>
<td>------------</td>
<td>-----------</td>
</tr>
<tr>
<td>1</td>
<td>163.060</td>
<td>163.069</td>
</tr>
<tr>
<td>2</td>
<td>204.087</td>
<td>204.087</td>
</tr>
<tr>
<td>3</td>
<td>366.190</td>
<td>366.139</td>
</tr>
<tr>
<td>4</td>
<td>528.192</td>
<td>528.192</td>
</tr>
<tr>
<td>5</td>
<td>690.245</td>
<td>690.247</td>
</tr>
<tr>
<td>6</td>
<td>690.245</td>
<td>690.247</td>
</tr>
<tr>
<td>7</td>
<td>731.272</td>
<td>731.272</td>
</tr>
<tr>
<td>8</td>
<td>1558.712</td>
<td>1556.710</td>
</tr>
<tr>
<td>9</td>
<td>1879.795</td>
<td>1879.206</td>
</tr>
<tr>
<td>10</td>
<td>1989.817</td>
<td>1986.804</td>
</tr>
</tbody>
</table>
11.5.7 Ion Match Table

The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings" (see 11.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings).

Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

11.5.8 Survey Chart

The “Survey” tab displays the precursor ion spectrum. The buttons that appear in this section are identical to those explained above in 11.5.3 Annotated Spectrum Chart - Additional Chart Controls.

11.5.9 LC/MS Snapshot

The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The glycan peptide feature is highlighted in purple and the selected PSM marker is yellow for O-linked and purple for N-linked. The span of the orange line in the LC-MS Snapshot designates the isolation window.
Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a \( m/z \) RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

11.5.10 Precursor Profile

The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with
the retention time along the x-axis and the intensity along the y-axis.

11.6 Glycan Search Peptide View
The "Peptide View" is very similar to the Glycopeptide View and it displays the resulting non-glycopeptide identifications. The table on the top shows the details of each peptide identification. The bottom section provides additional information about the peptide-spectrum matches for the native peptide selected in the table.

Note: If you are already familiar with the Glycopeptide Tab Section 11.5, you can skip this section as the interface is largely the same.

11.6.1 Peptide Table
The Peptide Table shows the filtered native peptide identification results. Each row in the table is a native peptide identification represented by its highest-scoring PSM. When there are more than 1000 peptides, the table is split into multiple pages.
Note: If there are multiple PSMs matched to a peptide, the table displays the information for the top-scoring PSM. Peptides with the same sequence but different modifications are considered different and shown in different rows.

To copy the sequence, right click on a row (or multiple rows) in the table and select “Copy selected peptide sequence(s)”. Another option is to mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells.

The following list describes the contents in each of the default columns:

- **Peptide**: The amino acid sequence of the peptide, as determined by the Glycan Search workflow. A modified residue is followed by a pair of parentheses enclosing the modification.
- **-10lgP**: The peptide -10lgP score. The score indicates the scoring significance of a peptide-precursor spectrum match.
- **Mass**: The monoisotopic mass of the peptide.
- **Length**: The number of amino acids in the sequence backbone.
- **ppm**: The precursor mass error, calculated as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$.
- **m/z**: The precursor mass-to-charge ratio.
- **RT**: The retention time (elution time) of the spectrum as recorded in the data.
• **Area per Sample**: The combined area for all features in that sample associated with this peptide.

• **Sample Profile**: A column only shown for Glycan Search results with two or more samples. Shows a profile view of the ratio between areas in each sample.

• **Group Profile**: A column only shown for Glycan Search results with two or more samples. Shows a profile view of the ratio between areas in each group.

• **#Vector**: A column only shown for Glycan Search results with two or more samples. Counts the number of feature vectors associated with this peptide.

• **Scan**: The scan number of the spectrum that matches the peptide sequence with the highest $-\log_{10}P$.

• **#Feature**: For single sample analysis, shows number of features assigned to the peptide sequence from identification. For multiple samples, if peptide has feature vectors, it will show number of features used (area > 0) in all feature vectors. If not, it shows the number of features from identification.

• **Id Count**: Number of identifications in total.

• **Accession**: The accessions of proteins that contain this native peptide.

• **PTM**: The types and the numbers of modifications present in the peptide shown in color-coded icons.

• **AScore**: Localization score assigned to modifications on the peptide. It is the $-\log$ of a $p$-value. In this case, the $p$-value is the probability that the modification occurs at the reported position compared to other possible positions. A $-\log_{10}P$ of 20 is equal to a $p$-value of 0.01. Scores are listed in the format AminoAcid, peptide position, modification, AScore. For example, serine at amino acid 3 is phosphorylated with an AScore of 54.01 will be reported as: S3 P 54.01.

11.6.2 Peptide Table controls
The menu above the peptide table provides options to scroll through the table of native peptides, filter the table, show optional columns and search for specific entries.

11.6.2.1 Peptide Table controls - Pagination
For easy viewing and to improve performance, the peptide table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.

In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

11.6.2.2 Glycan Table controls - View Filter
Click on the button to open the Peptide View Filter which is used to filter the Peptide table. The following are the filter options provided:
• Peptide sequence contains: Enter an amino acid sequence, only peptides that contain that sequence will be displayed in the peptide table. Adding PTMs in brackets in the sequence search is not handled, meaning only the sequence backbone is considered in the filter. Use the PTM contains for further filtering if necessary.

• Peptide score (-10lgP): Filters out all peptides with -10lgP scores below this threshold.

• Peptide length: Filters for peptides of length within this min/max threshold.

• Peptide sample area: Filters based on total peptide area found in each sample. Sample areas are calculated using the total of all peptide features from all spectra that identified the native peptide. If the peptide is found in multiple samples, only one sample must pass this filter for the peptide to be included.

• PTM contains: Type the names of PTMs you would like to include. Peptides containing PTMs that match the text will be included. If multiple PTMs match the text, they will all be included in the table.

Note: In all filters, multiple entries can be separated by a semi-colon.

Click on the Reset button to reset all filters to their default setting.

Click on OK to apply the filters to the Peptide table.

Click on Cancel to discard the changes in the View Filter.

11.6.2.3 Peptide Table controls - Optional Columns

Click on the button to open the optional columns dropdown. By default the sample area columns are displayed. Sample Feature count columns can also be shown. For projects with many samples, it might be useful to hide these optional columns and reduce the width of the peptide table.
11.6.2.4 Glycan Table controls - Search function

On the top right of menu, there is a search function with 4 options. Search results can be navigated between using the Up or Down arrows.

![Image of search function]

Options for searching include:

- **Scan =**: Search for the Scan number associated with each peptide. One scan can be associated to multiple peptides. For projects with multiple data files, the scan number is searched on every fraction.
- **m/z ≈**: Search for the approximate m/z. The search will consider a range of m/z equal to the search value $\pm 1$ of the last significant digit. For example, searching for m/z $\approx 560.21$ will find all m/z between [560.20, 560.22]
- **RT ≈**: Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value $\pm 1$ of the last significant digit set. For example, searching for RT $\approx 55.5$ will find all m/z between [55.4, 55.6]
- **Seq contains**: Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequences that have a substring with an exact match will be considered. Additionally, since the peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. searching for “LVIK” is the same as searching for “LVK”, “IVLK”, and “IVIK”

11.6.3 Feature selection breadcrumb

There is a sample and feature selection breadcrumb which allows you to switch between features associated with the peptide. Click on the Sample in the breadcrumb to reveal other samples and the features in that sample related to the current peptide selection. From here you can select other samples, fractions, or features.
By clicking on the fraction level, all fractions in this sample are revealed as well as the features in each fraction associated with this peptide. From here you can select other fractions and features to view.

Lastly, clicking on the feature level will show all features in the currently selected fraction. Features with Feature Id = 0 are known as fake features and are constructed to enhance identification.

11.6.4 PSM selection
Underneath the feature breadcrumb will shown the currently selected PSM. Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.

All other components in the Peptide tab interface will update according to the currently selected PSM.

For timsTOF instruments, Precursor ID is used rather than Scan number.

11.6.4.1 Protein jump button
Click on to jump to a protein that contains the currently selected PSM. If there are more than 1 protein, a dropdown is revealed which allows the user to select which protein to jump to. If a protein is filtered out due to a protein summary filter or summary page protein filter, then the option to jump will be grayed out.
11.6.4.2 Show spectrum in Data View

Click on to open the Data Refine MS/MS tab (if not already opened) and jump to the corresponding selected Scan.

11.6.4.3 Show spectrum in LC/MS View

Click on to open the Data Refine LC/MS tab (if not already opened) and jump to the corresponding selected Scan, highlighting the Glycopeptide MS/MS marker and the feature if present.

11.6.4.4 Show Raw Spectrum View

Click on to show the raw spectrum chart of the currently selected Scan.

11.6.5 Annotated Spectrum Chart

For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

11.6.5.1 Annotated Spectrum Chart - Chart Navigation

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.

Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.
Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.

See the whole spectrum: Double click in the spectrum or click the 1:1 button.

**11.6.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings**

Click on the button to show the spectrum annotation settings.

The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

Show Decimal Places: Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

m/z on Fragmentation: Select this to display the m/z value on top of the annotated ions.

m/z on Unannotated: Select this to display the m/z value on top of the peaks without ion matches.

Sequence Fragmentation: Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.
In Place Ion Info: Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

Intensity: Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

### 11.6.5.3 Annotated Spectrum Chart - Additional Chart Controls

Click on \( \# \) or \( % \) to toggle between absolute and percentage intensity values in the Y axis.

Click on \( 1:1 \) to reset the zoom level of the annotated spectrum chart to the default.

Click on \( 2x \) to zoom into the X axis by a factor of 2.

Click on \( 2y \) to zoom into the Y axis by a factor of 2.

The “alignment” check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

### 11.6.6 Ion Match Table

The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in “Spectrum Annotation Settings”.

<table>
<thead>
<tr>
<th>Ion Match</th>
<th>Survey</th>
</tr>
</thead>
<tbody>
<tr>
<td>#</td>
<td>b</td>
</tr>
<tr>
<td>1</td>
<td>1079.50</td>
</tr>
<tr>
<td>11</td>
<td>1265.57</td>
</tr>
<tr>
<td>12</td>
<td>1394.62</td>
</tr>
<tr>
<td>13</td>
<td>1481.69</td>
</tr>
<tr>
<td>14</td>
<td>1596.68</td>
</tr>
<tr>
<td>15</td>
<td>1653.70</td>
</tr>
<tr>
<td>16</td>
<td>1763.77</td>
</tr>
<tr>
<td>17</td>
<td>1870.81</td>
</tr>
<tr>
<td>19</td>
<td>2123.88</td>
</tr>
</tbody>
</table>

Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

### 11.6.7 Survey Chart

The “Survey” tab displays the precursor ion spectrum. The buttons that appear in this section are identical to those explained above in Section 11.6.5.3 Annotated Spectrum Chart - Additional Chart Controls.
11.6.8 LC/MS Snapshot

The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The peptide feature is highlighted in purple and the selected PSM marker is blue. The span of the orange line in the LC-MS Snapshot designates the isolation window.

Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a m/z RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.
11.6.9 Precursor Profile

The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with the retention time along the x-axis and the intensity along the y-axis.

11.7 Glycan Search Feature View

The Glycan Search Feature View shows all real features associated with peptides.
11.7.1 Feature Table
The Feature Table consists of the following columns.

- **Fraction**: The data file from which this feature is detected.
- **Feature ID**: A unique identifier of the feature from the data file.
- **Peptide**: The peptide sequence including PTMs and Glycans.
- **Glycan Type**: For glycan peptides, classifies the glycan as an N-link or O-link glycan.
- **Glycan**: Contains the glycan moiety information for the glycan peptide if applicable.
- **m/z**: Each feature in the feature vector has its own precursor mass-to-charge ratio. This column displays the average precursor mass-to-charge ratio of all of the features in the feature vector.
- **z**: The precursor charge of the feature or feature vector.
- **RT**: Simple the middle point of the retention time range of the feature or features in the feature vector. It is equal to \((\text{RT Begin} + \text{RT End}) / 2\)
- **RT Begin**: The lower bound of the retention time range of the feature or features in the feature vectors.
- **RT End**: The higher bound of the retention time range of the feature or features in the feature vectors.
- **1/k0**: Column shown for timsTOF instruments only, measuring the \(1/k0\) ion mobility range for the feature or feature vector.
- **Area**: Each feature in the feature vector has its own intensity area under the XIC curve. This column displays the average area of all the features in the feature vector.
- **ppm**: The mass error of a feature. For multiple samples, the mass error of the feature vector is calculated from the mass error of each feature in that vector.
- **Id Count**: The total number of identifications found for the peptide in fractions.
- **PTM**: Any PTMs in the peptide are shown here. Mouse over for a description of the PTM.
- **Accession**: A list of proteins that contain this peptide.

11.7.2 Feature Table controls
The menu above the feature table provides options to scroll through the table of features or feature vectors, filter the table, show optional columns and search for specific entries.

11.7.2.1 Feature Table controls - Pagination
For easy viewing and to improve performance, the glycan table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.
In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

11.7.2.2 Feature Table controls - Sample selection
The feature table only shows results from one sample at a time. Use the dropdown to switch between samples and update the Feature table accordingly.

11.7.2.3 Feature Table controls - Search function
On the top right of menu, there is a search function with 2 options. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- \( m/z = \): Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit.

For example, searching for \( m/z = 650 \) will find all m/z between \([649, 651]\)

- **Seq contains**: Search for the sequence in the table. If specifying a sequence without glycans or PTMs, the results will include that sequence and any sequences with glycans or PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered.

Additionally, since the glycan peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVIK” is the same as searching for “LVIK”, “Ivlk”, and “IVIK”

11.7.3 Feature Details
The Feature Details tab contains the eXtracted Ion Chromatogram (XIC) chart that displays the shape of the selected peptide feature vector over the retention time range where they were identified. The table at the top-left corner displays both the area and the ratio of the features in the feature vector. This table also works as a control to select the features that will be used to draw the XIC plot and other plots in the “Sample Features” and “RT Alignment” tabs.

11.7.4 PSM selection
Underneath the feature breadcrumb will shown the currently selected PSM. Click on the Left and Right arrow buttons to switch between PSMs related to the currently selected feature.
All other components in the Peptide tab interface will update according to the currently selected PSM.

For timsTOF instruments, Precursor ID is used rather than Scan number.

### 11.7.4.1 Show spectrum in Data View

Click on ![Data View icon](image) to open the Data Refine MS/MS tab (if not already opened) and jump to the corresponding selected Scan.

### 11.7.4.2 Show spectrum in LC/MS View

Click on ![LC/MS View icon](image) to open the Data Refine LC/MS tab (if not already opened) and jump to the corresponding selected Scan, highlighting the Glycopeptide MS/MS marker and the feature if present.

### 11.7.4.3 Show Raw Spectrum View

Click on ![Raw Spectrum View icon](image) to show the raw spectrum chart of the currently selected Scan.

### 11.7.5 Annotated Spectrum Chart

For the currently selected PSM, a graphical representation is shown in the form of an annotated spectrum chart. There are several controls provided for this view which will be explored in the following sections.

#### 11.7.5.1 Annotated Spectrum Chart - Chart Navigation

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.
Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.

Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.

Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.

See the whole spectrum: Double click in the spectrum or click the 1:1 button.

11.7.5.2 Annotated Spectrum Chart - Spectrum Annotation Settings

Click on the button to show the spectrum annotation settings.

The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. The selected ion types will be displayed in the “Ion Match” table, as well. It is possible to annotate
the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

Show Decimal Places: Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

m/z on Fragmentation: Select this to display the m/z value on top of the annotated ions.

m/z on Unannotated: Select this to display the m/z value on top of the peaks without ion matches.

Sequence Fragmentation: Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

In Place Ion Info: Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

Intensity: Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

11.7.5.3 Annotated Spectrum Chart - Additional Chart Controls

Click on or to toggle between absolute and percentage intensity values in the Y axis.

Click on to reset the zoom level of the annotated spectrum chart to the default.

Click on to zoom into the X axis by a factor of 2.

Click on to zoom into the Y axis by a factor of 2.

The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

The “error map” check box is used to show/hide the error map underneath the chart.

11.7.6 Ion Match Table

The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings".
Note: Uncheck the alignment checkbox in the spectrum annotation view to see the highlighted fragment ions more easily.

11.7.7 Survey Chart

The “Survey” tab displays the precursor ion spectrum. The buttons that appear in this section are identical to those explained above in Section 11.7.5.3 Annotated Spectrum Chart - Additional Chart Controls.

11.7.8 LC/MS Snapshot

The LC-MS Snapshot displays a zoomed-in view of the precursor scan in the LC-MS heat map 2-D view. The peptide feature is highlighted in purple and the selected PSM marker is blue. The span of the orange line in the LC-MS Snapshot designates the isolation window.
Users can change the view from the 2-D heat map to a 3-D intensity view by clicking the intensity view button in the upper right-hand corner of the LC-MS Snapshot. In addition, users can export the LC-MS heat map by clicking the export image button in the upper right-hand corner of the LC-MS Snapshot.

Expand the options dropdown by clicking to reveal a search option to specify a m/z RT (ensure there is a space between these two numbers) location to zoom into. There is also a Peak slider filter to filter out low intensity peaks and a feature area filter to filter out low area features.

11.7.9 Precursor Profile

The Precursor Profile, or eXtracted Ion Chromatogram (XIC) of the precursor ion, graphs the elution profile of the selected peptide feature vector over the retention time range where it was identified. The graph is plotted with the retention time along the x-axis and the intensity along the y-axis.
11.8 Glycan Search Partial View

The "Partial View" displays glycopeptides that either have no identified glycan or no identified peptide (if it has both it will show in the Glycan table). For those with no identified glycan, only the spectrum information is saved. For those with no identified peptide but with identified glycan, both spectrum information and glycan information are saved.

### 11.8.1 Partial Table

The Partial Table shows the filtered glycan peptide identification results. Each row in the table is a glycopeptide identification represented by its highest-scoring PSM. When there are more than 1000 peptides, the table is split into multiple pages.

To copy cell contents, mouse over any cell in the table and hitting Ctrl + C on your keyboard to copy the contents of those selected cells. This also works when selecting multiple rows and hitting Ctrl+C.

The following list describes the contents in each of the columns in the Unidentified table:

- **Index**: Simply the row count of the table. If the table is sorted, the Index remains the same order while the other columns sort.

- **Scan Number**: The scan number of the spectrum.
• **Peptide:** The peptide sequence or mass with partial identification.

• **Peptide Score:** The score assigned to identification of native peptide.

• **Glycan Moiety:** The best glycan moiety on this spectrum. For N-link results, there will always be 1. For O-link results, there could be more than 1.

• **Glycan ID:** The unique ID number we assign to that Glycan structure.

• **Glycan Mass:** The mass of the Glycan.

• **Glycan Type:** Identified whether the glycan is N-link or O-link.

• **Glycan Score:** The normalized sum of ions matched and unmatched. The score can be negative if the sum of unmatched ions intensity exceeds the matched ions.

• **m/z:** The precursor mass-to-charge ratio.

• **z:** The charge of this spectrum.

• **Adduct:** Indicates if an adduct is present and which adducts

• **Isotopic Shift:** Indicates if there is an isotopic shift

• **Fraction:** The fraction that this spectrum belongs to.

11.8.2 Partial Table controls
The menu above the peptide table provides options to scroll through the table of glycan peptides and search for specific entries.

11.8.2.1 Partial Table controls - Pagination
For easy viewing and to improve performance, the glycan table is limited to showing 1000 entries at once. Pagination is used to manage which set of 1000 entries will be viewed.

In the top left, there is a dropdown to select the rows for viewing. Click the left arrow or right arrow to select the previous/next 1000 rows. If the table is sorted or filtered, it may update the pagination set of rows that is currently in view.

11.8.2.2 Partial Table controls - Search function
On the top right of menu, there is a search function with 3 options. Search results can be navigated between using the Up or Down arrows.
Options for searching include:

- **Scan or Precursor Id =**: Search for the Scan number (or Precursor Id for timsTOF instruments) for the unidentified spectrum. For projects with multiple data files, the scan number is searched on every fraction.

- **m/z ≈**: Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit.
  
  For example, searching for m/z = 800.80 will find all m/z between [800.79, 800.81]

- **Seq contains**: Search for the sequence in the table. If specifying a sequence without glycans or PTMs, the results will include that sequence and any sequences with glycans or PTMs with the same backbone. If specifying a sequence with bracket and some mass value, then only sequence that have a substring with an exact match will be considered.

11.8.3 Partial Spectrum Chart

For the currently selected spectrum, a graphical representation is shown in the form of an annotated spectrum chart. Similar to the other annotated spectrum charts, there are several controls provided for this view which will be explored in the following sections. Since there are no peptide identification, several of the provided options that are shared have no use in this chart.

11.8.3.1 Partial Spectrum Chart - Chart Navigation

**Zoom to a m/z region**: Drag horizontally from the start m/z and to the end m/z with left mouse button.

**Zoom in/out smoothly**: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.

**Increase/Decrease peak intensity**: Place the cursor in the spectrum and scroll the mouse wheel.

**See the whole spectrum**: Double click in the spectrum or click the 1:1 button.

11.8.3.2 Partial Spectrum Chart - Spectrum Annotation Settings

Click on the button to show the spectrum annotation settings.
It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places**: Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation**: This option has no use in this chart.

**m/z on Unannotated**: Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation**: This option has no use in this chart.

**In Place Ion Info**: Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity**: Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

**11.8.3.3 Partial Spectrum Chart - Additional Chart Controls**

Click on # or % to toggle between absolute and percentage intensity values in the Y axis.

Click on[](image) to reset the zoom level of the annotated spectrum chart to the default.

Click on 2X to zoom into the X axis by a factor of 2.
Click on 🕵️ to zoom into the Y axis by a factor of 2.

The "alignment" check box ✅ alignment is used to show the peptide alignment.

The “error map” check box ✅ error map is used to show/hide the error map underneath the chart.

11.8.4 Partial Glycan Ion Match Table

The Glycan Ion Match Table shows the theoretical m/z and observed m/z for each moiety of glycan ions matched in the spectrum. These columns are sortable and can be rearranged.

11.8.5 Partial Ion Match Table

The “Ion Match” tab in the bottom pane contains a table with possible fragment ions for the selected MS/MS scan. Each ion in the table shows the calculated mass, where if the fragment ion is present in the spectrum, its mass value is displayed in color. N-terminal ions are shown in blue and C-terminal ions are shown in red. A fragment ion is found when there is a matching peak within the mass error tolerance, as defined in the de novo sequencing parameters, and when relative intensity of the matching peak is at least 2%. The ion types displayed in the table can be configured in "Spectrum Annotation Settings".

11.8.6 Partial Survey Chart

The “Survey” tab displays the precursor ion spectrum. For Orbitrap instruments, the controls are the same that appear in this section are identical to those explained above in 11.8.3 Partial Spectrum Chart - Additional Chart Controls.
11.9 Glycan Search Exporting

In each of the Glycan Search tables, there is an export button. Clicking on the export button will open the Export dialog which will contain options to export the table in CSV format. This export considers all table view filters and sorting applied to the table. This export is meant to be the same as what is shown in the table.

For the total exporting options, the Export node can be opened by double-clicking on the Export Node in the Project View. The Export node contains all the text and HTML exports for the relevant tables provided in the Glycan Search interface.

Click on Select All to select all export options. Click on Clear All to deselect all export options.

Click on any individual checkbox to enable that file for exporting or click on a parent checkbox to select all the exports underneath that group. For example, selecting “Glycan Search result Exports” will check all the CSV exports underneath it.

Click on the Browse button to select a location to save the exports into.

Click on Export to begin exporting all the files. Once the export is complete, a pop-up window will show.

In the following sections we will go over each of the export contents.
Selecting “Glycan Search table exports in CSV” will select all exports that are in .csv format and the exports that are formed from the various tables provided in the Glycan Search result.

**Note:** Unless otherwise specified, the exports consider the summary page filters as set in the analysis. The View Filters are not considered in the Exports as those filters are intended to only filter the view for easy interpretation.

### 11.9.1 Export - Glycan Search Protein CSV
Exports the Protein table into *Glycan Search.proteins.csv* file. The columns in the export consists of those that are explained in the Protein table with optional columns checked and includes a few additional columns and differences:

- **Protein Group:** This is the same as the Cluster column in the proteins table and it is the protein cluster ID.
- **#Glycans:** The count of the unique glycan types identified in this protein.
- **Glycan ID:** A unique ID assigned to each Glycan structure to distinguish between them.
- **Glycan Found:** A TRUE/FALSE flag for easy filtering the .csv which indicates whether there are glycopeptides in this protein.
- **#Glycan Sites:** Indicates the number of sites in the protein residue that has glycosylation.

### 11.9.2 Export - Glycan Search Support Peptide CSV
Exports the Supporting Native Peptides table into *Glycan Search.protein-peptides.csv* file. The columns in the export consists of those that are explained in the Protein supporting glycopeptides table with some useful additional columns:

- **Protein Group:** Since the protein table is not present, a protein group number is provided.
- **Accession:** Likewise, to know which protein this supporting native peptide belongs to, the protein accession is saved in the export.
- **Top:** Indicates whether this is protein is a top protein in the protein group.
- **z:** The charge of the native peptide. Selected based on the top PSM.
- **Area per sample:** The area for each sample associated with this supporting native peptide.

### 11.9.3 Export - Glycan Search Support Glycan Peptide CSV
Exports the Supporting Glycan Peptides table into *Glycan Search.glycoprotein-peptides.csv* file. The columns in the export consists of those that are explained in the Protein supporting non-glycopeptides table with some useful additional columns:

- **Protein Group:** Since the protein table is not present, a protein group number is provided.
- **Accession:** Likewise, to know which protein this supporting native peptide belongs to, the protein accession is saved in the export.
- **Top:** Indicates whether this is protein is a top protein in the protein group.
• **Glycan ID:** A unique ID assigned to each Glycan structure to distinguish between them.

• **Glycan Mass:** The mass of the glycan(s) in this glycan peptide. Provided for easy sorting.

• **Area per sample:** The area for each sample associated with this supporting glycopeptide.

• **Source File:** The source file for which this Scan belongs to. The Scan and Source File of the highest -10lgP score PSM is the one shown.

• **#Feature per sample:** The number of features in each sample associated with this supporting glycopeptide.

• **Glycosylation position in Peptide:** Indicates which amino acid in the peptide chain has the glycosylation.

• **Glycosylation position in Protein:** Indicates which site in the protein residue has the glycosylation that has been identified in this glycopeptide.

### 11.9.4 Export - Glycan Search Peptide CSV
Exports the Native Peptides table into *Glycan Search.peptides.csv* file.

The columns in the export consists of those that are explained in the Peptide table section **11.6.1 Peptide Table**.

### 11.9.5 Export - Glycan Search Glycan Peptide CSV
Exports the Glycan Peptides table into *Glycan Search.glycopeptides.csv* file.

The columns in the export consists of those that are explained in Glycopeptide table section with some useful additional columns:

- **Glycan ID:** A unique ID assigned to each Glycan structure to distinguish between them.

- **Glycan Mass:** The mass of the glycan(s) in this glycan peptide. Provided for easy sorting.

- **z:** The charge of the glycan peptide. Selected based on the top PSM.

- **Glycosylation position in peptide:** Indicates which amino acid in the peptide chain has the glycosylation.

### 11.9.6 Export - Glycan Search PSM CSV
This Peptide Spectrum Match export is separated for each sample and is named in the format *Glycan Search.psms.[sample name].csv*.

The following columns are exported:

- **Peptide:** The amino acid sequence of the peptide, as determined by the Glycan Search search workflow. A modified residue is followed by a pair of parentheses enclosing the modification.

- **-10lgP:** The PSM -10lgP score. The score indicates the scoring significance of a peptide-precursor spectrum match.

- **Mass:** The monoisotopic mass of the peptide.
• **Length**: The number of amino acids in the sequence backbone.

• **ppm**: The precursor mass error, calculated as $10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass}$.

• **m/z**: The precursor mass-to-charge ratio.

• **z**: The charge of the PSM.

• **RT**: The retention time (elution time) of the spectrum as recorded in the data.

• **Scan**: The scan number of the PSM.

• **Area**: The area of the feature associated with this PSM.

• **Source File**: The fraction/data file that this spectrum belongs to.

• **Accession**: The accessions of proteins that contain this native PSM.

• **PTM**: The types and the numbers of modifications present in the peptide shown in color-coded icons.

• **AScore**: Localization score assigned to modifications on the peptide. It is the $-10 \log$ of a p-value. In this case, the p-value is the probability that the modification occurs at the reported position compared to other possible positions. A $-10\lgP$ of 20 is equal to a p-value of 0.01. Scores are listed in the format `AminoAcid, peptide position, modification, Ascore`. For example, serine at amino acid 3 is phosphorylated with an AScore of 54.01 will be reported as: S3 P 54.01.

**11.9.7 Export - Glycan Search Glycan PSM CSV**

This Glycan Peptide Spectrum Match export is separated for each sample and is named in the format `Glycan Search.glycopsms.[sample name].csv`.

The following columns are exported:

• **Peptide**: The amino acid sequence of the glycan peptide, as determined by the Glycan Search search workflow. A modified residue is followed by a pair of parentheses enclosing the modification or glycan mass.

• **Glycan**: The Glycan moiety on this glycopeptide. For N-link results, there will always be 1. For O-link results, there could be more than 1.

• **Glycan ID**: A unique ID assigned to each Glycan structure to distinguish between them. A negative Glycan ID indicates a decoy glycan.

• **Glycan Type**: Simply states whether the glycan moiety is N-link or O-link. This column is particularly useful for N-Link + O-link search workflows.

• **-10lgP**: The peptide $-10\lgP$ score. The score indicates the scoring significance of a peptide-precursor spectrum match.

• **Glycan Score**: The normalized sum of ions log intensity, based on glycan fragment ion, peptide ion, and glycopeptide ion (Y ion).

• **Peptide Score**: Classifies glycan peptides with the normalized sum of ions log intensity, based on peptide fragment ions, and those peptide fragment ions with glycan. Different fragmentation type considers different peptide fragment ion:
- CID and HCD considers b,y ion without glycan and b,y ion with at most 1 glycan node from root
- ETD considers b,y,c,z,z' ion without glycan and b,y,c,z,z' ion with whole glycan from root
- ETHCD considers b,y,c,z,z' ion without glycan, b,y,c,z,z' ion with 1 glycan from root, and b,y,c,z,z' ion with whole glycan from root

- **Glycan Rank**: Each spectrum has up to 10 glycan candidates. The candidates are ordered by their glycan score and the rank means the rank of the glycan score which has the best final score. e.g. Rank 3 means the candidate has the 3rd best glycan score but has the best final score.

- **S-Score (%):** Measures the confidence of the top glycan peptide candidate as opposed to other candidates for the spectrum. S-Score = \( \frac{\text{finalScore top1candidate} - \text{finalScore top2candidate}}{\text{finalScore top1candidate}} \). The higher the score the better. 100% indicates only 1 result, and it is the best match. 0% indicates that the top1 and top2 result are very similar, and we cannot confidently say the result is the best match.

- **Mass**: The monoisotopic mass of the glycan peptide.
- **Glycan Mass**: The mass of the glycan(s) in this glycan peptide. Provided for easy sorting.
- **Length**: The number of amino acids in the sequence backbone.
- **ppm**: The precursor mass error, calculated as \( 10^6 \times (\text{precursor mass} - \text{peptide mass}) / \text{peptide mass} \).
- **m/z**: The precursor mass-to-charge ratio.
- **z**: The charge.
- **RT**: The retention time (elution time) of the spectrum as recorded in the data.
- **Area**: The area of the feature associated with this glycan PSM.
- **Scan**: The scan number of the glycan PSM.
- **Source File**: The fraction/data file that the spectrum belongs to.
- **Accession**: The accessions of proteins that contain this glycan PSM.
- **PTM**: The types and the numbers of modifications present in the peptide shown in color-coded icons.
- **AAScore**: Localization score assigned to modifications on the peptide. It is the -10 log of a p-value. In this case, the p-value is the probability that the modification occurs at the reported position compared to other possible positions. A -10lgP of 20 is equal to a p-value of 0.01. Scores are listed in the format AminoAcid, peptide position, modification, Ascore. For example, serine at amino acid 3 is phosphorylated with an AAScore of 54.01 will be reported as: S3 P 54.01.

11.9.8 Export - Glycan Search Feature CSV

The feature export is named in the format *Glycan Search.features.csv*

The columns shown in Features CSV export are consistent with the Feature with the columns rearranged and a few additional columns:
- **Glycan ID:** A unique ID assigned to each Glycan structure to distinguish between them.

- **-10lgP:** The top PSM -10lgP score. The score indicates the scoring significance of a peptide-precursor spectrum match.

- **Glycan Scores:** The normalized sum of ions log intensity, based on glycan fragment ion, peptide ion, and glycopeptide ion (Y ion).

- **Peptide Scores:** Classifies glycan peptides with the normalized sum of ions log intensity, based on peptide fragment ions, and those peptide fragment ions with glycan. Different fragmentation type considers different peptide fragment ion:
  - CID and HCD considers b,y ion without glycan and b,y ion with at most 1 glycan node from root
  - ETD considers b,y,c,z,z' ion without glycan and b,y,c,z,z' ion with whole glycan from root
  - ETHCD considers b,y,c,z,z' ion without glycan, b,y,c,z,z' ion with 1 glycan from root, and b,y,c,z,z' ion with whole glycan from root

- **Mass:** The monoisotopic mass.

- **Glycan Mass:** The mass of the glycan if present.

- **Length:** The length of the sequence backbone.

- **Glycosylation position in peptide:** Indicates which amino acid in the peptide chain has the glycosylation if it is present.

The Feature Vectors CSV export resembles the Features export except the m/z, RT mean, and Area is provided for each sample in separate columns.

### 11.9.9 Export - Glycan Search Glycosylation Sites CSV

This export is the full glycosylation sites statistics table as shown in the Summary page.

The export is named in the format `Glycan Search.glycosylationSites.csv`

For all samples it will list both N-linked and O-linked Glycans for combined N-link and O-linked Glycan searches. For every position in proteins in the protein table, if the glycan is found at that position it will be counted as a Glycosylation Site in this table.

### 11.9.10 Export - Glycan Search Glycan Partial Spec CSV

This export is for the Partial table and is consistent with the columns in section 11.8.1 Partial Table. There is a separate export for each sample in the result and the exports are named in the format `Glycan Search.glycoPartialIdentifiedSpec.[sample name].csv`

A few extra columns are provided:

- **Glycan Mass:** The mass of the glycan if present.

- **RT:** The Retention Time of the spectrum.

- **PepMass:** The mass of the peptide portion of the unidentified spectrum.
11.9.11 Export - Glycan Search All Spec CSV
This export is for all spectrum in the result. It includes glycopeptides, native peptides, and the unidentified. There is a separate export for each sample in the result and the exports are named in the format Glycan Search.spec.[sample name].csv

The columns included in the export are as follows:

- raw m/z
- raw z
- m/z: The refined m/z.
- z: The charge after refinement (charge correction).
- Scan: The scan number of the spectrum.
- Glycan: Y or N flag indicating whether there is a glycan match.
- Native: Y or N flag indicating whether there is a native peptide match.
- Source File: The fraction or data file this spectrum belongs to.

11.9.12 Export - Glycan Search HTML

Export in HTML: This will generate single or multiple HTML reports in the specified location.

The following exporting options are available:

- Summary view: The “Summary” view page will be saved as a summary.html file in HTML format in the specified location.
- Protein coverage: The coverage pane will be saved for each protein.
- Supporting peptides: A list of supporting peptides will be saved for each protein.
- Supporting glycol peptides: A list of supporting glycopeptides will be saved for each protein.
- Best unique PSM: The best unique PSM will be saved for each protein.
- Export peptide list in html: creates a separate HTML-containing peptide identifications.
- Export glycol peptide list in html: creates a separate HTML-containing glycopeptide identifications.

An individual protein will have its own HTML output file, where the corresponding protein coverage, the supporting peptides, and the best unique PSM are gathered.
12. Tools

PEAKS Studio 11 comes with a few tools to help the user in analyzing results. These are the mass calculator, spectral library viewer, and compare result. There are two other useful tools in setting and monitoring performance which are the performance configuration tool and worker monitor tool.

12.1 Mass Calculator

The Mass Calculator is a handy tool used to quickly calculate mass or m/z based on sequence, modification, b/y-ion, and charge. Click on \[ \text{in the Tools tab to open the calculator.} \]

Copy/Type in a sequence (with modifications) and click on the ‘Calculate’ button to quickly find out the total residue mass and m/z mass.

12.2 Spectral Library Viewer

Click on \[ \text{in the Tools menu to open the Spectral Library Viewer.} \]

The Spectral Library Viewer is very useful in analyzing the contents of PEAKS Studio compatible libraries. It shows library info, distribution charts, library table content, and a spectrum chart of the top 20 ions.

PEAKS Studio compatible libraries are:

- Spectral libraries exported from PEAKS Studio 11 (.tsv).
- Spectral libraries generated from PEAKS Online (.tsv).
- Spectral libraries exported from PEAKS Studio X+ and Xpro (.lib).
- OpenMS generated libraries (.tsv).
- Spectronaut 14 libraries and Spectronaut libraries with the same format and column headers (.xls).
To open a library, click on and select a PEAKS compatible library. Spectral Library Viewer supports PEAKS text format libraries generated by PEAKS Studio, PEAKS Online, and even PEAKS Studio Xpro (the older version of PEAKS Studio). In the Open dialog box, select the .tsv file (for Online and PEAKS Studio 11 spectral libraries) or the .lib file (for PEAKS Studio Xpro spectral libraries). The viewer will detect the library file format and also search for the .info (metainfo) file automatically. The library information and details will be shown in a separate tab in the main window of the viewer. Several libraries can be opened at once as tabs and can be switched between.

Already configured libraries can also be opened by clicking on the icon below for convenience. All configured libraries will be listed in a dropdown.

Click on to export a copy of this library. This library will be saved in PEAKS Studio 11 format. This means that this will convert PEAKS Studio Xpro libraries into the PEAKS Studio 11 format.

Click on to directly configure this library into PEAKS Studio to be used in Library Search.
12.2.1 Library Viewer - Meta Information

Libraries generated by PEAKS products will come with a .info file which displays various information relevant to spectral library and its generation procedure. This file is not required and will not be necessary to configure the library or view in library viewer, but is recommended to accompany each library file. The .info file is automatically detected and loaded, as long as it is the only .info file in the same folder as the spectral library file, otherwise the file name should be the same.

<table>
<thead>
<tr>
<th>Library Info</th>
</tr>
</thead>
<tbody>
<tr>
<td>Library Name: library</td>
</tr>
<tr>
<td>Library Path: D:\share\SpectralLibraries_Desktop\ETHcD_2022-09-23\Analysis 1\library.tsv</td>
</tr>
<tr>
<td>Project Name: ETHcD_2022-09-23</td>
</tr>
<tr>
<td>Result Name: Analysis 1</td>
</tr>
<tr>
<td>Ion Mobility: SL</td>
</tr>
<tr>
<td>Fixed Modifications: Carbamidomethylation 57.0215</td>
</tr>
<tr>
<td>Variable Modifications: Deamidation (NQ) 0.9940 Oxidation (M) 15.9949</td>
</tr>
<tr>
<td>Precursor Tolerance: 10.00 ppm</td>
</tr>
<tr>
<td>Fragment Tolerance: 0.02 Da</td>
</tr>
<tr>
<td>Database: uniprot_sprot_June-2-2020.fasta</td>
</tr>
<tr>
<td>Contaminant Database: no loss -NH3,-H2O etc</td>
</tr>
<tr>
<td>Use RT or iRT: RT</td>
</tr>
<tr>
<td>Acquisition Method: DDA</td>
</tr>
<tr>
<td>Enzyme: Specified by each sample</td>
</tr>
<tr>
<td>Entry Count: 3575</td>
</tr>
</tbody>
</table>

In the top left of the Library Viewer, the Library Info is extracted from the .info file. The following fields are displayed:

- Library Name: The name given to the library file. By default this would be library.tsv.
- Library Path: The location of the library file.
- Project Name: The name of the project used to generate the library.
- Result Name: The name of the analysis which the Database Search result used to generate the library.
- Ion Mobility: Shows as SL if there is no ion mobility in the library. Otherwise it may indicate that there is timesTOF 1/k0 or FAIMS CV values in the library.
- Fixed modifications: Fixed modifications used in the identification search used to generate the spectral library.
- Variable modifications: Variable modifications used in the identification search used to generate the spectral library.
- Precursor mass error tolerance: Precursor mass error tolerance used in the identification search used to generate the spectral library.
- Fragment mass error tolerance: Fragment ion mass error tolerance used in the identification search used to generate the spectral library.
- Database: Sequence database used for the database search used to generate the library.
- Contaminant Database: Contaminant database used for the database search used to generate the library.
- Fragmentation Annotation: Fragmentation annotation of the raw data.
- Use RT or iRT: Libraries can be saved with original RT or normalized iRT values. This indicates which of those options was used in generating this library.
- Acquisition Method: Shows if the library came from DDA or DIA data.
- Enzyme: Enzyme digest selected in the identification search used to generate the spectral library.
- Entry Count: The total number of entries in the library.

Note: A spectral library might not include all of the meta information listed above.
12.2.2 Library Viewer - Statistical Distribution

The spectral Library Viewer is equipped with several statistical analysis of the library for quality assessment and validation. Different statistical plots can be selected from a drop-down list and the corresponding graph is displayed.

The following statistical plots are available for validation purposes:

**Precursor m/z distribution**

**Precursor charge distribution**
Peptide length distribution

Peptide modification overview
Matched ion distribution

Right-click on any of the charts to export the image.
12.2.3 Library Viewer - Library Table
The spectral library entries are listed in the library table. Each row of the table corresponds to an ionized peptide entry in the library.
The library table contains the following columns:

- **Peptide**: Peptide base sequence with no modifications
- **PTM**: Modifications in the peptide, color coded for different modifications. Hovering over a PTM icon shows the detailed modification information such as the modified residue, modification site and modification mass.
- **m/z**: Theoretical mass-over-charge of the peptide.
- **z**: The charge of the peptide.
- **iRT**: Normalized retention time of the peptide if library was generated using iRT. This column will display un-normalized values if the library was general with original RT.
• **timsTOF 1/k0**: If the library is from timsTOF data. There will be 1/k0 values saved in the library which will be used in library search.

• **FAIMS CV**: If the library is from FAIMS data. The CV (compensation voltage) values are saved and will be used in library search.

**Note**: Even if a library does not have timsTOF 1/k0 or FAIMS CV, it can still be used to run library search for those respective data. If the library does have timsTOF 1/k0 or FAIMS CV, it can also be used to run library search for standard data and those column values will be disregarded.

All columns are sortable and can be rearranged.

### 12.2.3.1 Library Table - Pagination

By default, 1000 entries are shown in the table at any one time. This is to ensure performance so that only 1000 entries will be displayed and refreshed at one time. Click on the dropdown or the left/right arrows to switch between the 1000 entry segments.

### 12.2.3.2 Library Table - Search function

On the top menu, there is a search function with 3 options. Search results can be navigated between using the Up or Down arrows.

Options for searching include:

- **m/z ≈**: Search for the approximate m/z. The search will consider a range of m/z equal to the search value +/- 1 of the last significant digit. For example, searching for m/z ≈ 560.21 will find all m/z between [560.20, 560.22]

- **RT ≈**: Search for the approximate RT. Likewise, the search will consider a range of RT equal to the search value +/- 1 of the last significant digit set. For example, searching for RT ≈ 55.5 will find all m/z between [55.4, 55.6]

- **Seq contains**: Search for the sequence in the table. If specifying a sequence without PTMs, the results will include that sequence and any sequences with PTMs with the same backbone. If specifying a sequence with bracket and some mass value (like in the above image), then only sequence that have a substring with an exact match will be considered. Additionally, since the peptide table shows both I/L and they can not be distinguished, then searching for I in the sequence is equivalent to searching for L in the sequence. i.e. Searching for “LVLK” is the same as searching for “LVIK”, “IVLK”, and “IVIK”
12.2.3.3 Library Table - Spectrum Chart

Each library peptide will have a spectrum chart which displays the top 20 ions saved in the spectral library.

Moving the cursor over the peptide sequence in the spectrum will show the mass transitions for a particular amino acid residue.

Zoom to a m/z region: Drag horizontally from the start m/z and to the end m/z with left mouse button.
Zoom in/out smoothly: Place the cursor at a particular m/z value (right below the x-axis), zoom in/out by scrolling the mouse wheel.
Increase/Decrease peak intensity: Place the cursor in the spectrum and scroll the mouse wheel.
See the whole spectrum: Double click in the spectrum or click the 1:1 button.

Click on the button to show the spectrum annotation settings.
The spectra in the PEAKS results can be annotated by selecting ion types from the thorough collection of ions that PEAKS offers. It is possible to annotate the spectrum with various ions for CID, EThcD and ETD. Each of these fragmentation modes can be set with different settings which will be shared with other results with the same fragmentation type.

To reset the defaults, click on the “Reset default” button.

**Show Decimal Places**: Select the number of decimal places that will appear in the ion table and the spectrum view. The default is set to two decimal places.

**m/z on Fragmentation**: Select this to display the m/z value on top of the annotated ions.

**m/z on Unannotated**: Select this to display the m/z value on top of the peaks without ion matches.

**Sequence Fragmentation**: Select this to display the sequence fragmentation on the top-left corner of the “Spectrum Annotation” view.

**In Place Ion Info**: Ion information, m/z value, and relative intensity are all displayed in a pop-up within the “Spectrum Annotation” view when this option is checked and the cursor is placed on a peak.

**Intensity**: Set the intensity threshold for spectrum annotation to low (2%), medium (5%), or high (10%).

Click on 🡇 or 🡧 to toggle between absolute and percentage intensity values in the Y axis.

Click on 🡢 to reset the zoom level of the annotated spectrum chart to the default.

Click on 🡤 to zoom into the X axis by a factor of 2.

Click on 🡥 to zoom into the Y axis by a factor of 2.
The "alignment" check box allows user to display/hide how the fragment ions generated from the peptide align with the spectrum. N-terminal ions are shown in blue and C-terminal ions are shown in red.

12.2.3.4 Library Table - Peptide Details
For each selected peptide, more details are displayed.

<table>
<thead>
<tr>
<th>Peptide Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequence: HTM/QARWK</td>
</tr>
<tr>
<td>Modifications: 3 Oxidation (M) 15.9949</td>
</tr>
<tr>
<td>Mass: 1203.5641</td>
</tr>
<tr>
<td>Pre m/z: 402.1953</td>
</tr>
<tr>
<td>Charge: 3</td>
</tr>
<tr>
<td>iRT: 16.93</td>
</tr>
<tr>
<td># of peaks: 8</td>
</tr>
</tbody>
</table>

- **Sequence**: The peptide backbone sequence without modifications
- **Modifications**: Lists the modifications for this peptide, the mass value and the position in peptide
- **Mass**: The peptide mass
- **Pre m/z**: The precursor m/z
- **Charge**: Charge of the peptide
- **iRT**: This could be original RT or normalized iRT.
- **# of peaks**: The number of peaks in this library entry. Equal to the number of ions in the spectrum chart.

12.2.3.5 Library Table - Ion Table
List of fragment ions. The ion type, m/z and relative intensity of each fragment ion are listed in the table.

<table>
<thead>
<tr>
<th>#</th>
<th>Ion</th>
<th>m/z</th>
<th>intensity(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>z1</td>
<td>158.09</td>
<td>30.86</td>
</tr>
<tr>
<td>2</td>
<td>y1</td>
<td>175.12</td>
<td>100.00</td>
</tr>
<tr>
<td>3</td>
<td>b[2+]</td>
<td>181.08</td>
<td>3.83</td>
</tr>
<tr>
<td>5</td>
<td>c3H2O</td>
<td>229.13</td>
<td>6.00</td>
</tr>
<tr>
<td>6</td>
<td>c3-H</td>
<td>246.13</td>
<td>11.65</td>
</tr>
<tr>
<td>7</td>
<td>c4-H</td>
<td>377.17</td>
<td>4.80</td>
</tr>
<tr>
<td>8</td>
<td>y4</td>
<td>528.38</td>
<td>21.05</td>
</tr>
<tr>
<td>9</td>
<td>y5</td>
<td>543.35</td>
<td>15.03</td>
</tr>
<tr>
<td>10</td>
<td>y6</td>
<td>730.38</td>
<td>18.77</td>
</tr>
<tr>
<td>11</td>
<td>c8H2O</td>
<td>805.35</td>
<td>19.77</td>
</tr>
<tr>
<td>12</td>
<td>b8</td>
<td>806.34</td>
<td>9.25</td>
</tr>
<tr>
<td>13</td>
<td>y7</td>
<td>845.41</td>
<td>53.26</td>
</tr>
<tr>
<td>14</td>
<td>b9</td>
<td>903.39</td>
<td>4.13</td>
</tr>
<tr>
<td>15</td>
<td>z8</td>
<td>950.43</td>
<td>15.46</td>
</tr>
<tr>
<td>16</td>
<td>c10-H</td>
<td>1047.47</td>
<td>30.37</td>
</tr>
<tr>
<td>17</td>
<td>z10</td>
<td>1117.49</td>
<td>4.24</td>
</tr>
</tbody>
</table>
12.3 Compare Result

Click on the Projects menu to open the Compare Result dialog.

In compare result, 2 to 4 identification results can be selected to do a comparison. Important to note that only results in the same category can be used for comparison. There are four categories (which are split based on which workflow they are from):

- PEAKS DB search results: DB Search, PEAKS PTM, SPIDER
- DeepNovo Peptidome results: Peptide Search
- Glycan Search results: Glycan Search
- DIA Identification results: Spectral Library, DB Search

This means that if a DB Search result is selected for comparison, a Peptide Search or Glycan Search result cannot be selected, but another DB Search, a PEAKS PTM, or SPIDER result can be selected.

After selecting the results, a compare result dialog is shown. It has a Proteins tab and Peptide tab with table comparison, venn diagram, and score distribution. There is an additional Glycan tab for Glycan comparisons only.

12.3.1 Compare Result - Protein Compare Table
For results with Proteins, the set of proteins are compared between the different results. The key used for comparison is the Protein ID, which is the protein accession.
The following columns are in the Protein compare table:

- **Protein Description**: The description of the protein.
- **Cover Map**: Indicates which results contain this Protein ID. The cover map order is the same as the other of the results in the table.
- **Score**: PEAKS protein -10lgP score.
- **#Spec**: The number of PSMs matched to supporting peptides for the protein.
- **#Pep**: The number of supporting peptides for the protein.
- **#Uniq**: The number of unique supporting peptides for the protein.
- **%Cov**: The sequence coverage of the protein by supporting peptides.

### 12.3.1 Protein Compare Table - Pagination

By default, 1000 entries are shown in the table at any one time. This is to ensure performance so that only 1000 entries will be displayed and refreshed at one time. Click on the dropdown or the left/right arrows to switch between the 1000 entry segments.
12.3.1.2 Protein Compare Table - Protein Result Filter

Click on the button to open the Protein Result Filter.

Score threshold: Sets a protein score threshold for each of the up to four results in the comparison.

Show protein: Only displays proteins that meet the criteria of the cover map below:

- All: All proteins are listed.
- Common: Only proteins found in every result are listed.
- Unique in X: Only proteins found in the selected result but not other results are listed.

Click on the button to remove all filters.

Click on to apply the Protein View Filter to the Protein table.

Click on to discard changes and close the dialog.

12.3.1.3 Protein Compare Table - Search Function

An easy-to-use search function is available for the protein table. The protein accession or description substring can be searched for, and all search results can be navigated through with the Up and Down arrows.

Protein accession/name contains: Restrict the protein table to only display protein IDs which contain the entered text in its accession/name.
Protein description contains: Only proteins with descriptions that contain the entered text will be included in the protein table if text is entered in this field.

12.3.1.4 Protein Compare Table - Exporting

Click on to open the Export pop-up and export the contents of the Protein table.

![Export Protein Table](image)

The export content is the same as what is displayed in the table, which consider the result filters.

12.3.1.5. Protein Compare Table - Venn Diagram

A Protein Number Venn Diagram is generated which shows the overlap of Proteins between the selected results. The type of venn diagram depends on the number of results nodes selected. As an example, the venn diagram below is comparing 4 result nodes. Different venn diagrams are also available for two sample and three sample comparisons.

The venn diagram considers the result filter and will update accordingly. Right-click on the venn diagram to get the option to export the image.

![Protein Number Venn Diagram](image)

12.3.1.6. Protein Compare Table - Score Distribution

The Protein Score Distribution plot shows the number of proteins at a certain score. The higher the rank, the higher the protein score. This gives a quick overview of how many proteins are at certain score thresholds.

The score distribution chart considers the result filter and will update accordingly. Right-click on the score distribution chart to get the option to export the image.
12.3.2 Compare Result - Peptide Compare Table

The set of peptides are compared between the different results. The key used for comparison is the Peptide sequence.

<table>
<thead>
<tr>
<th>Peptide</th>
<th>PTM</th>
<th>Score Map</th>
<th>Analysis 1 Spectral Library</th>
<th>Analysis 1 DB DB</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>GALKSYCSTLDEETALVSESEGK</td>
<td></td>
<td>45.91 1034.4888 2 34.95 1113.52 1.322 Yes</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>UIYLR</td>
<td></td>
<td>41.85 533.3231 2 36.68 0.8673 - 0.0673 No</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>VPHLR</td>
<td></td>
<td>36.37 935.338 2 35.15 0.8471 - 0.0723 No</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>STVHLDLWPLYQSD</td>
<td></td>
<td>30.83 380.6979 2 30.89 1.0961 - 1.1183 No</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>QAQEREL</td>
<td></td>
<td>36.37 622.8098 2 15.85 0.0831 - 0.0642 No</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>ELLRQDLEGRL</td>
<td></td>
<td>47.58 701.049 2 28.49 0.0915 - 1.0151 No</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>TQDCFNSWYDN</td>
<td></td>
<td>32.25 367.0351 2 30.89 0.8997 - 0.0597 No</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>8DGPRNLCSTDEGQSK</td>
<td></td>
<td>47.58 619.3962 2 28.49 0.0831 - 1.0151 No</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>LIKDNQDCSTDEGQVLKSLQYN</td>
<td></td>
<td>40.10 398.0983 3 23.89 0.7947 - 0.0457 No</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>YCSTDEGQVLKSLQYN</td>
<td></td>
<td>47.63 754.3215 2 17.87 0.8969 - 1.0291 No</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>VPTRH</td>
<td></td>
<td>44.38 519.3281 2 22.82 0.0793 - 0.0957 No</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>DAVMTYCLSTDEGQNFLRLERD</td>
<td></td>
<td>45.28 1431.5772 2 28.49 1.0857 - 1.0875 Yes</td>
<td></td>
</tr>
<tr>
<td>13</td>
<td>PILPRDQSCSTDEGQSLRN</td>
<td></td>
<td>44.39 694.6888 2 33.51 0.7978 - 1.0151 No</td>
<td></td>
</tr>
<tr>
<td>14</td>
<td>IEKDESAIMK</td>
<td></td>
<td>37.81 703.691 2 25.39 0.8445 - 1.0045 No</td>
<td></td>
</tr>
<tr>
<td>15</td>
<td>TOQQR</td>
<td></td>
<td>44.78 509.2314 2 30.89 0.7234 - 0.0328 No</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>LQVSQNOYGGYSPCAGTR</td>
<td></td>
<td>45.76 608.0412 2 18.47 0.8794 - 1.0828 Yes</td>
<td></td>
</tr>
<tr>
<td>17</td>
<td>TVGQR</td>
<td></td>
<td>28.4 328.7612 2 22.84 0.0897 - 0.0597 No</td>
<td></td>
</tr>
<tr>
<td>18</td>
<td>VQQVCSTDEGQSLRN</td>
<td></td>
<td>36.60 607.8441 2 29.23 0.8445 - 1.0045 No</td>
<td></td>
</tr>
<tr>
<td>19</td>
<td>FQRFQK</td>
<td></td>
<td>34.49 473.7770 2 27.41 0.8035 - 0.0586 No</td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>ANQNGQKNNQK</td>
<td></td>
<td>34.25 446.9015 2 31.21 1.0780 - 1.1359 No</td>
<td></td>
</tr>
<tr>
<td>21</td>
<td>LUQOSHGNAG</td>
<td></td>
<td>27.73 744.4432 2 16.11 1.0519 - 1.0729 No</td>
<td></td>
</tr>
<tr>
<td>22</td>
<td>LEOQDGQAFPHNNAYKVLNG</td>
<td></td>
<td>45.81 916.4226 2 35.21 0.0760 - 1.1352 No</td>
<td></td>
</tr>
<tr>
<td>23</td>
<td>ALQGDRMALQDVD</td>
<td></td>
<td>45.82 672.3532 2 30.89 0.0793 - 1.0151 No</td>
<td></td>
</tr>
<tr>
<td>24</td>
<td>AOSGQGSQGCQGQIHQNHAQVR</td>
<td></td>
<td>43.22 694.9402 2 24.40 0.7650 - 0.1257 No</td>
<td></td>
</tr>
<tr>
<td>25</td>
<td>LSUQGDHRMGRDQDKD</td>
<td></td>
<td>45.82 942.7430 2 28.49 0.8869 - 1.0727 No</td>
<td></td>
</tr>
<tr>
<td>26</td>
<td>YGADQGHIDNQK</td>
<td></td>
<td>45.86 542.6276 2 21.24 0.7322 - 0.0566 No</td>
<td></td>
</tr>
<tr>
<td>27</td>
<td>QGADQQNHQADQG</td>
<td></td>
<td>45.86 304.6409 2 27.17 0.0753 - 0.0329 Yes</td>
<td></td>
</tr>
<tr>
<td>28</td>
<td>TDGQDQAYQDSAYGR</td>
<td></td>
<td>46.95 962.4758 2 20.26 1.0919 - 1.0811 No</td>
<td></td>
</tr>
<tr>
<td>29</td>
<td>VHLDOSQK</td>
<td></td>
<td>45.83 528.2649 2 17.20 0.8059 - 1.0153 No</td>
<td></td>
</tr>
<tr>
<td>30</td>
<td>EVANQGEGHLDEYSHLMQDK</td>
<td></td>
<td>45.52 843.0781 2 24.40 0.0921 - 1.1953 No</td>
<td></td>
</tr>
<tr>
<td>31</td>
<td>LKTVSTQEDGQYDSYKTVSTLIS</td>
<td></td>
<td>47.88 615.3737 2 33.51 0.8495 - 0.0705 No</td>
<td></td>
</tr>
<tr>
<td>32</td>
<td>HKNYC</td>
<td></td>
<td>45.57 539.8885 2 25.96 0.8507 - 1.0111 No</td>
<td></td>
</tr>
<tr>
<td>33</td>
<td>LeGQSTGQVSQGSDRVR</td>
<td></td>
<td>45.85 708.8889 2 18.47 0.8794 - 1.0828 Yes</td>
<td></td>
</tr>
<tr>
<td>34</td>
<td>SDCSTYRQGKSTQSGGSLGKYR</td>
<td></td>
<td>47.69 815.3893 2 18.29 1.0295 - 1.0835 No</td>
<td></td>
</tr>
<tr>
<td>35</td>
<td>ASMQVQQWFK</td>
<td></td>
<td>48.06 720.8808 2 11.06 0.8041 - 1.0291 No</td>
<td></td>
</tr>
<tr>
<td>36</td>
<td>AHQCMNLQGFCGEGFYFK</td>
<td></td>
<td>40.10 549.4312 2 22.19 1.0697 - 1.1358 No</td>
<td></td>
</tr>
<tr>
<td>37</td>
<td>AQPQSPVNYK</td>
<td></td>
<td>45.57 908.8747 2 27.17 1.0869 - 1.7199 No</td>
<td></td>
</tr>
</tbody>
</table>

The following columns are in the Protein compare table:

- **Peptide**: The peptide sequence. Used as the key for comparison.
- **PTM**: The modifications in the peptide. Mouse-over the PTM to see more details in a tooltip.
• **Cover Map**: Indicates which results contain this peptide. The cover map order is the same as the other of the results in the table.
• -10lgP: Top PSM -10lgP score.
• m/z: The m/z of this peptide in this result.
• z: The charge of this peptide in this result.
• RT: The retention of this peptide in this result.
• 1/k0: For timsTOF data only. Shows the 1/k0 value of the result.
• CV: For FAIMS data only. Shows the CV (compensation voltage) of the result.
• Multi-Hit: Indicates if there are multiple PSM hits.

### 12.3.2.1 Peptide Compare Table - Pagination

By default, 1000 entries are shown in the table at any one time. This is to ensure performance so that only 1000 entries will be displayed and refreshed at one time. Click on the dropdown or the left/right arrows to switch between the 1000 entry segments.

### 12.3.2.2 Peptide Compare Table - Peptide Result Filter

Click on the button to open the Peptide Result Filter.

Peptide contains: Only keeps the peptides that contain the sequence. Multiple sequences can be listed here if separated by semicolon. E.g. search for “LSS;GV”.

Score Threshold: Sets a peptide -10lgP threshold for each of the up to four results in the comparison.

Show peptides: Only displays proteins that meet the criteria of the cover map below:

- **All**: All peptides are listed.
- **Common**: Only peptides found in every result are listed.
- **Unique in X**: Only proteins found in the selected result but not other results are listed.
Click on the **Reset** button to remove all filters.

Click on **OK** to apply the Protein View Filter to the Protein table.

Click on **Cancel** to discard changes and close the dialog.

### 12.3.2.3 Peptide Compare Table - Search Function

An easy-to-use search function is available for the peptide compare table. Use this to search for certain sequences of interest.

![Search Function](image)

**seq contains** LAL

1/19

### 12.3.2.4 Peptide Compare Table - Exporting

Click on **Export** to open the Export pop-up and export the contents of the Peptide table.

![Export Peptide Table](image)

The export content is the same as what is displayed in the table, which consider the result filters.

### 12.3.2.5. Peptide Compare Table - Venn Diagram

A Peptide Number Venn Diagram is generated which shows the overlap of Peptides between the selected results. The type of venn diagram depends on the number of results nodes selected. As an example, the venn diagram below is comparing 2 result nodes. Different venn diagrams are also available for four sample and three sample comparisons.

The venn diagram considers the result filter and will update accordingly. Right-click on the venn diagram to get the option to export the image.
12.3.2.6 Peptide Compare Table - Score Distribution

The Peptide Score Distribution plot shows the number of peptides at a certain score. The higher the rank, the higher the peptide score. This gives a quick overview of how many proteins are at certain score thresholds.

The score distribution chart considers the result filter and will update accordingly. Right-click on the score distribution chart to get the option to export the image.

12.3.3 Compare Result - Glycan Compare Table

The set of glucan peptides are compared between the different results. The key used for comparison is the Peptide sequence.
The following columns are in the Protein compare table:

- **Peptide**: The peptide sequence. Used as the key for comparison.
- **Glycan**: Lists the Glycan moiety for this peptide.
- **Glycan Type**: Indicates if it is an N-linked or O-linked glycopeptide.
- **PTM**: The modifications in the peptide. Mouse-over the PTM to see more details in a tooltip.
- **Cover Map**: Indicates which results contain this peptide. The cover map order is the same as the other of the results in the table.
- **-10lgP**: Top PSM -10lgP score.
- **Glycan Score**: The glycan score of each result.
- **Peptide Score**: The peptide score of each result.
- **S-score**: The S-score of each result.
- **m/z**: The m/z of this peptide in this result.
- **RT**: The retention of this peptide in this result.
- **1/k0**: For timsTOF data only. Shows the 1/k0 value of the result.
- **CV**: For FAIMS data only. Shows the CV (compensation voltage) of the result.
- **Multi-Hit**: Indicates if there are multiple PSM hits.

### 12.3.3.1 Glycan Compare Table - Pagination

By default, 1000 entries are shown in the table at any one time. This is to ensure performance so that only 1000 entries will be displayed and refreshed at one time. Click on the dropdown or the left/right arrows to switch between the 1000 entry segments.
12.3.3.2 Glycan Compare Table - Peptide Result Filter

Click on the button to open the Peptide Result Filter.

Peptide contains: Only keeps the peptides that contain the sequence. Multiple sequences can be listed here if separated by semicolon. E.g. search for “LSS;GV”.

Score Threshold: Sets a peptide -10lgP threshold for each of the up to four results in the comparison. There is a separate filter for Glycan score and Peptide score. There is also a separate filter for N-linked glycopeptides and O-linked glycopeptides.

Show glycan: Only displays proteins that meet the criteria of the cover map below:

- All: All peptides are listed.
- Common: Only peptides found in every result are listed.
- Unique in X: Only proteins found in the selected result, but not other results are listed.

Click on the button to remove all filters.

Click on to apply the Protein View Filter to the Protein table.

Click on to discard changes and close the dialog.
12.3.3.3 Glycan Compare Table - Search Function
An easy-to-use search function is available for the glycan compare table. Use this to search for certain sequences of interest.

seq contains LAI

12.3.3.4 Glycan Compare Table - Exporting
Click on to open the Export pop-up and export the contents of the Peptide table.

The export content is the same as what is displayed in the table, which consider the result filters.

12.3.3.5. Glycan Compare Table - Venn Diagram
A Peptide Number Venn Diagram is generated which shows the overlap of Glycopeptides between the selected results. The type of venn diagram depends on the number of results nodes selected. As an example, the venn diagram below is comparing 2 result nodes. Different venn diagrams are also available for four sample and three sample comparisons.

The venn diagram considers the result filter and will update accordingly. Right-click on the venn diagram to get the option to export the image.

12.3.3.6. Glycan Compare Table - Glycan Score Distribution
The Glycan Score Distribution plot shows the number of glycan peptides at a certain glycan score. The higher the rank, the higher the glycan score. This gives a quick overview of how many proteins are at certain score thresholds.

The score distribution chart considers the result filter and will update accordingly. Right-click on the score distribution chart to get the option to export the image.
12.3.3.7. Glycan Compare Table - Peptide Score Distribution

The Peptide Score Distribution plot shows the number of glycan peptides at a certain peptide score. The higher the rank, the higher the peptide score. This gives a quick overview of how many proteins are at certain score thresholds.

The score distribution chart considers the result filter and will update accordingly. Right-click on the score distribution chart to get the option to export the image.

12.4 Performance Configuration

In the Configurations tab, click on the icon to open the Performance Configuration dialog.

The Computer Resources shows the detected memory and number of processors of the computer.
By default, performance configuration tries to set a reasonable total thread and allocated memory count to make use of the computer resources while leaving some for other programs.

You can manually change these default values and click ‘Apply’, but only do so if you understand the system requirements for other programs.

The Detach Service JVM is recommended to be checked. 2 threads and 8 GB memory will be reserved for processing user interface updates. If this is unchecked, the user interface processing and project/analysis processing tasks will be shared and PEAKS Studio may freeze more often if no resources are available for refreshing the user interface.

GPU resource checkbox will only be enabled if the computer has an NVIDIA CUDA compatible GPU. The GPU should have an updated driver and at least 8 GB memory to be considered usable. The GPU is only used for DIA Database step, and can be disregarded if not running DIA Database Search.

Click on ‘Cancel’ to discard any changes.

12.5 Worker Monitor

Click on  in the Tools tab to open the Worker Monitor.

The Worker Monitor shows which tasks are currently running in PEAKS Studio. In addition, it indicates how many threads and how much memory is being used for each task. This is useful in checking the utilization of PEAKS Studio, and also for debugging purposes.
Click on the ‘Refresh’ button to update the worker monitor. Check ‘Auto Refresh’ to continuously update the worker monitor information every 2 seconds.

<table>
<thead>
<tr>
<th>Task ID</th>
<th>Task step type</th>
<th>CPU usage</th>
<th>Total memory</th>
<th>Free memory</th>
<th>Used memory</th>
<th>Threads</th>
<th>GPU</th>
</tr>
</thead>
<tbody>
<tr>
<td>0x46c1f-5c45-4c23-b406-7461994593af</td>
<td>denovoService</td>
<td>16.3%</td>
<td>5.0 GB</td>
<td>2.6 GB</td>
<td>2.1 GB</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>5c60ee65-7ew4-47ba-9e6a-770de677407</td>
<td>denovoService</td>
<td>16.5%</td>
<td>2.6 GB</td>
<td>1.9 GB</td>
<td>704.9 MB</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>