INTRODUCTION ................................................................................................................. 5

INTRODUCTION TO PEAKS 4.5 ....................................................................................... 5

HOW TO USE THIS MANUAL .......................................................................................... 5

   Scope ......................................................................................................................... 6
   Terminology and Abbreviations Glossary ................................................................. 6

GETTING STARTED WITH PEAKS 4.5 ................................................................................ 9

SCV WHAT WE WILL NEED ............................................................................................... 9

   Package contents ........................................................................................................ 9
   System requirements .................................................................................................. 9
   Instrumentation ......................................................................................................... 9

SCV INSTALLATION ....................................................................................................... 10

SCV VISTA INSTALLATION ............................................................................................ 11

SC CONNECTING TO A PEAKS ONLINE SERVER .............................................................. 12

SC REGISTERING PEAKS ............................................................................................... 13

SC DATABASE CONFIGURATION ..................................................................................... 14

S inCHORUS (SEARCH ENGINE) CONFIGURATION ......................................................... 18

FEATURES WALKTHROUGH ............................................................................................. 19

SC BEGIN THE WALKTHROUGH .................................................................................. 19

SVC VIEWING RESULTS ................................................................................................. 21

SVC MANIPULATING RESULTS .................................................................................... 23

SC inCHORUS SEARCHING .......................................................................................... 24

SC SPIDER SEARCHING FOR HOMOLOGUES ............................................................... 26

   Merging reports ....................................................................................................... 27

GRAPHICAL USER INTERFACE ..................................................................................... 29

SVC WINDOWS, DIALOGUES, FRAMES AND REPORTS .................................................... 30

SVC PEAKS 4.5 main window ........................................................................................ 30

SC Auto de Novo Parameters Dialogue ........................................................................ 31

SC Protein Identification Parameters Dialogue .......................................................... 32

SCV PEAKS Properties Dialogue .................................................................................. 33

SCV Enzyme Editor Dialogue ....................................................................................... 34

SC PTM Selector Dialogue ............................................................................................ 35

SCV PTM Editing Dialogue ............................................................................................ 36

S Instrument Definition Window .................................................................................. 36

SCV Environment Preferences Window ......................................................................... 38

SCV Protein Identification Result Window .................................................................... 38

   De Novo View ......................................................................................................... 39
   Peptide View .......................................................................................................... 39
   Protein View .......................................................................................................... 39
   Search parameters .................................................................................................. 41
   Filter Pane .............................................................................................................. 41

SCV Main Processing Window ....................................................................................... 43

SCV Ion Editor ................................................................................................................ 45

S marks a section applicable to PEAKS Studio, C for PEAKS client and V for PEAKS Viewer
Data extraction procedure: ................................................................. 70
SCV  TRACKING DATA PROPERTIES ............................................................. 70
SC  REFINING DATA BEFORE ANALYSIS .................................................... 71
SCV  MANUALLY MANIPULATING DATA FILES ........................................ 74
SCV  Editing Precursor information ............................................................ 74
SCV  Manually merging MS/MS scans of the same peptide ..................... 74
SCV  Cutting and Copying Spectrum Data ................................................. 75
SCV  Pasting Spectrum Data ................................................................. 75
SCV  Peptide View ......................................................................................... 104
Search Parameters ......................................................................................... 104
SC  USING PEAKS WITH MODIFICATIONS (PTM) ..................................... 77
SC  AUTO DE NOVO SEQUENCING ........................................................... 78
SCV  VIEWING AUTO DE NOVO RESULTS ............................................... 81
SCV  EDITING SEQUENCING RESULTS (PREPARATION) ....................... 82
SCV  MANUAL DE NOVO SEQUENCING ................................................... 84
SCV  Creating a fresh spectrum for sequencing ..................................... 84
SCV  Manual de novo Operations ............................................................ 84
Selecting a peak ......................................................................................... 84
Measuring distance along the m/z scale .................................................... 84
Measure the m/z difference between two PEAKS .................................. 85
Deselect a peak ......................................................................................... 85
Zoom in on part of the spectrum .............................................................. 85
Add (remove) ions to (from) a peak ......................................................... 85
Using sequence tags ................................................................................ 86
Undoing an edit ......................................................................................... 86
Redoing an edit ......................................................................................... 86
SCV  SUGGESTING A SEQUENCE TO SEE HOW IT FITS THE DATA .......... 87
SC  PROTEIN IDENTIFICATION ............................................................... 89
SC  PEAKS Protein Identification ............................................................. 89
SC  inChorus Protein Identification .......................................................... 92
Options for search engines inChorus ...................................................... 93
Importing results for inChorus ............................................................... 94
SCV  SELECTING UNMATCHED DE NOVO RESULTS ............................. 94
SC  RUNNING PROTEIN IDENTIFICATION ON SELECT SPECTRA ........ 95
SC  SPIDER SEARCH FOR SEQUENCE HOMOLOGY ............................... 95
SC  MERGING SPIDER AND PROTEIN ID REPORTS ................................ 97
SC  CREATING A HIGH-THROUGHPUT WORKFLOW ............................... 98
SCV  USING THE MASS CALCULATOR .................................................... 100
SCV  PREDICTING RETENTION TIME ..................................................... 101
SC  Training the RT prediction algorithm ............................................... 101
SC  Predicting RT after training ............................................................. 102
SCV  VIEWING PROTEIN IDENTIFICATION RESULTS ......................... 104
Search Parameters ..................................................................................... 104
Peptide View ............................................................................................. 104

S marks a section applicable to PEAKS Studio, C for PEAKS client and V for PEAKS Viewer
<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peptide Details and SPIDER Powered Alignment</td>
<td>105</td>
</tr>
<tr>
<td>Protein View</td>
<td>107</td>
</tr>
<tr>
<td>SCV COMPARING PROTEINS – MULTIPLE SEQUENCE ALIGNMENT</td>
<td>109</td>
</tr>
<tr>
<td>SCV THE TABLE OF VALUES</td>
<td>110</td>
</tr>
<tr>
<td>SCV What do the columns mean?</td>
<td>110</td>
</tr>
<tr>
<td>SCV TOGGLING COLUMNS</td>
<td>112</td>
</tr>
<tr>
<td>SCV FILTERING RESULTS</td>
<td>113</td>
</tr>
<tr>
<td>SCV How filters act on de novo sequences</td>
<td>115</td>
</tr>
<tr>
<td>SCV Filter examples</td>
<td>116</td>
</tr>
<tr>
<td>Publication</td>
<td>116</td>
</tr>
<tr>
<td>Digging for a protein by name</td>
<td>116</td>
</tr>
<tr>
<td>Setting a protein mass range</td>
<td>117</td>
</tr>
<tr>
<td>SCV Saving/Loading Filter sets</td>
<td>117</td>
</tr>
<tr>
<td>SCV Using a default filter</td>
<td>117</td>
</tr>
<tr>
<td>SCV EXPORTING RESULTS</td>
<td>117</td>
</tr>
<tr>
<td>SCV WYSIWYG reports</td>
<td>117</td>
</tr>
<tr>
<td>SCV Exporting peak lists</td>
<td>118</td>
</tr>
<tr>
<td>SCV Exporting Sequences by spectrum</td>
<td>118</td>
</tr>
<tr>
<td>SCV Exporting Peptide Sequences</td>
<td>119</td>
</tr>
<tr>
<td>SCV Exporting Exclusion Lists</td>
<td>119</td>
</tr>
<tr>
<td>SCV Exporting high-resolution spectral images</td>
<td>119</td>
</tr>
<tr>
<td>SCV SAVING RESULTS</td>
<td>121</td>
</tr>
<tr>
<td>ABOUT BIOINFORMATICS SOLUTIONS INC.</td>
<td>122</td>
</tr>
<tr>
<td>PEAKS SOFTWARE LICENSE</td>
<td>123</td>
</tr>
<tr>
<td>REFERENCE: PEAKS PAPER</td>
<td>125</td>
</tr>
</tbody>
</table>
Introduction

Introduction to PEAKS 4.5

*PEAKS makes the interpretation of MS/MS data easier and faster.*

PEAKS is an innovative software system designed to derive amino acid sequences and identify proteins from tandem mass spectrometry data. After running MS/MS on a protein sample, PEAKS performs *de novo* sequencing and database search identification of the protein(s) and peptides using raw experimental data.

PEAKS 4.5 provides peptide sequence and protein identification results via an intuitive interface, allowing for rapid visual interpretation. PEAKS provides both auto and manual *de novo* sequencing tools for detailed examination of MS/MS spectra, providing the flexibility to manually modify auto *de novo* results when searching for additional sequence possibilities.

How to use this manual

*There’s no need to read this manual cover to cover.* To get familiar with the software, just read the walkthrough in Chapter 3. Otherwise, it’s best to just keep it as a reference.

This user’s manual is intended to help us get started using PEAKS 4.5, acquaint us with its functionality, show us how to customize PEAKS to our application, allow us to work efficiently with the interface, provide a task based reference, and help us with troubleshooting. As such, this manual is organized into chapters based on these categories. Use the table of contents at the front of this manual to access the relevant section.

---

*S marks a section applicable to PEAKS Studio, C for PEAKS client and V for PEAKS Viewer*
Since PEAKS Studio, PEAKS Viewer and PEAKS Client share the same user’s interface, this manual covers all three software programs. Where a section is applicable to PEAKS Studio, an ‘S’ is noted, ‘C’ denotes PEAKS Client, and ‘V’ denotes PEAKS Viewer. Some sections apply to all three, so ‘SCV’ is noted.

Scope
PEAKS users are assumed to be familiar with computer usage, and the operating system environment. As such, it is beyond the scope of this manual to instruct the user on the use of windows, dialogue boxes, menus, file storage etc. Please refer to the operating system’s manual, or computer help books for such information. Similarly, PEAKS users are expected to be familiar with mass spectrometry, standard operating practices and data.

Terminology and Abbreviations Glossary

**a-ions:** an N-terminal fragment holding at least one charge; similar to b-ions and c-ions. This is a prefix fragment of the peptide. The a-ion’s mass will be the sum of the masses of the N terminal group, plus the intervening neutral amino acid residues, subtract the mass of Carbon Monoxide.

**b-ions:** an N-terminal fragment holding at least one charge; similar to a-ions and c-ions. This is a prefix fragment of the peptide. The b-ion’s mass will be the sum of the masses of the N terminal group, plus the intervening neutral amino acid residues.

**c-ions:** an N-terminal fragment holding at least one charge; similar to a-ions and b-ions. This is a prefix fragment of the peptide. The c-ion’s mass will be the sum of the masses of the N terminal group, plus the intervening neutral amino acid residues, plus the mass of ammonia.

**Deconvolution:** rearrangement of the spectrum to show each monoisotopic peak as if it were singly charged. Thus, to reposition them on the scale, PEAKS multiplies the m/z of ion’s that were doubly charged by two. Note that the deconvoluted scale PEAKS shows is ‘at +1.’

**Fixed modification:** selecting a post-translational modification as a fixed modification tells PEAKS that this modification is applied to all occurrences of the residue(s) that the PTM can act on.

**Enzyme:** The residues PEAKS can find in different positions in the sequence. This is based on information about the enzyme used to digest our protein sample.

**BSI (Bioinformatics Solutions Inc.):** The makers of PEAKS and other fine bioinformatics software.
**ESI (Electrospray Ionization):** A method for ionizing a sample into the mass spectrometer.

**m/z:** mass to charge ratio.

**MALDI (Matrix-Assisted Laser Desorption/Ionization):** A method for ionizing a sample into the mass spectrometer. This has a characteristic effect of only producing singly charged ions.

**Mass accuracy:** this refers to the accuracy of the spectrometer and its resulting data. On a spectrum, this is reflected by how close the PEAKS are to the actual masses of the ions they represent.

**PTM (Post Translational Modification):** A protein, just translated and hence newly formed, may differ from its final form as a result of interaction with the cellular environment (or the experimental environment). As they interact chemically with the environment, residues may gain or lose molecules. This change is referred to as a post-translational modification. Since PTM changes the mass of residues, it must be accounted for when sequencing peptides by mass spectrometry.

- **Built-in PTM:** PEAKS comes equipped with a library of possible post-translational modifications. These can be incorporated into a de novo analysis at the click of a button.
- **Customized PTM:** If the post-translational modification we are looking for is not in the PEAKS PTM set, we may create our own entry, or modify an existing one. This will appear as a customized PTM in the set.
- **PTM library:** A listing of all possible (built-in and custom entered) post-translational modifications that PEAKS can use as a part of its analysis.

**Residue:** as used in this manual, a residue refers to what remains of an amino acid once it has become part of a peptide, or peptide fragment. In this manual, residues are referred to by their original amino acid names.

**Resolution:** refers to the precision of an instrument. On a spectrum, this is reflected by how close together can two PEAKS be and still be told apart.

**Variable modification:** selecting a post-translational modification as a variable modification tells PEAKS that this modification may or may not be applied to any given occurrence of the residue(s) that the PTM can act on.

**x-ions:** a C terminal fragment holding at least one charge; similar to y-ions and z-ions. This is a suffix fragment of the peptide. The x-ion's mass will be the sum of
the masses of the C terminal group, plus the intervening neutral amino acid residues, plus the mass of Carbon Monoxide.

**y-ions:** a C terminal fragment holding at least one charge; similar to x-ions and z-ions. This is a suffix fragment of the peptide. The y-ion’s mass will be the sum of the masses of the C terminal group, plus the intervening neutral amino acid residues, plus the mass of H₂.

**z-ions:** a C terminal fragment holding at least one charge; similar to x-ions and y-ions. This is a suffix fragment of the peptide. The z-ion’s mass will be the sum of the masses of the N terminal group, plus the intervening neutral amino acid residues, subtract the mass of ammonia.
Getting started with PEAKS 4.5

Everything we need to know from the beginning and step by step.

This section of the manual will guide us through the process of installation and configuration of PEAKS 4.5. If problems persist, contact technical support.

SCV

What we will need

Package contents
The PEAKS 4.5 package should contain:

- This manual

- PEAKS 4.5

System requirements
PEAKS 4.5 will run on most platforms with the following requirements:

- Equivalent or superior processing power to a Pentium III at 800 MHz.

- At least 1 GB of memory (RAM). 1.5GB is recommended. For PEAKS Studio / PEAKS Client. PEAKS Viewer requires only 500MB.

- JAVA Virtual Machine 1.6 or better (provided on installation).

Instrumentation
PEAKS 4.5 will work with data from any type of tandem mass spectrometer designed for proteomics work.

PEAKS will accept data in the following formats:
Agilent instrument's data in .d format provided that PEAKS and Agilent's software, including BaseCommon.dll, BaseDataAccess.dll, and MassSpecDataReader.dll. are installed on the same computer.

Applied Biosystems instrument's data in .wiff format provided that PEAKS, the Infochromics converter plug-in and Analyst are installed on the same computer. PEAKS has the ability to read directly from the 4700/4800 Oracle database.

Bruker instrument's data in .yep, .baf and .fid formats.

Thermo Electron instrument's data in RAW format provided that PEAKS and XCalibur are installed on the same computer, or dta format (and concatenated dta formats), with the ability to load an entire folder full of dta’s.

Waters/Micromass instrument's data in .RAW format, provided that PEAKS and MassLynx are installed on the same computer, or pkl files.

All instrument's data as can be converted into mzXML, mzData, .pkl, dta, or .mgf.

SCV Installation

To make sure we only use the latest information available to PEAKS, if we already have PEAKS installed on our system, we must uninstall it before proceeding. For those users running Windows Vista, please refer to the Vista Installation section.

1. Close all programs that are currently running and end all non-system tasks.

2. Insert the PEAKS 4.5 disc into the CD-ROM drive.

3. Auto-run should automatically load the installation software. If it does not, find the CD-ROM drive and open it to access the disc. Click on the exe file.

4. A menu screen will appear. Select the top item “Install”.

5. The installation utility will begin the install. Wait while it does so. Choose ‘English’ as the language for installation instructions. When the “PEAKS 4.5” installation dialogue appears, click the “Next” button.

6. Read the license agreement. If we agree to it, we change the radio button at the bottom to select “I accept the terms of the License Agreement” and click “Next”.
7. Next we choose the folder/directory in which we’d like to install PEAKS 4.5. Press the “Choose…” button to browse our system and make a selection, or type a folder name in the textbox. Click “Next”.

8. Choose where we’d like to place icons for PEAKS 4.5. The default will put these icons in the programs section of our start menu. Click “Next”.

9. Review the choices we have made. We can click “Previous” if we’d like to make any changes or click “Next” if those choices are correct.

10. PEAKS 4.5 will now install on our system. We may cancel at any time by pressing the “Cancel” button in the lower left corner.

11. When installation is complete, click “Done”. The “PEAKS 4.5” menu screen should still be open. One may view movies and materials from here. To access this menu again, we simply insert the disc in our CD-ROM drive.

**SCV Vista Installation**

PEAKS is written in Java, and is therefore platform independent. It runs on Windows Vista, but certain 3rd party programs, including BioWorks, and some of the data conversion utilities, do not work on Windows Vista. X!Tandem, OMSSA, and the Muscle (Multiple Sequence Alignment) have been demonstrated to work within PEAKS on Windows Vista.

However, the installation utility, “InstallAnywhere” by Macrovision, is not capable of completing the PEAKS installation on Windows Vista. Two components fail:

1) creation of shortcuts

2) allocation of memory

To complete the installation:

1) Follow the on-screen instructions. The final screen, shown above, notes the errors.
2) Go to the PEAKS program folder (the place where you installed PEAKS, usually c:\program files\peaks studio\) and find the file called run.bat. This is the file used to launch PEAKS.

3) Edit run.bat in notepad. It contains a command to start peaks, which looks like:

    start jre\bin\javaw -Xmx1024m -jar peaks.jar

the 1024m tells PEAKS to run with 1024MB of memory. This is a default setting, and since installAnywhere was unable to check how much memory is on our system, we should edit this manually. If we don’t have 1GB of memory on our system, we should reduce this number (or preferably, increase the amount of physical RAM available). If we have 1.5GB we can increase it to 1500m. Save the file.

4) Right click on the file and choose ‘Create Shortcut’ from the pop-up menu. A shortcut called ‘Shortcut to run.bat’ will be placed in the same folder. Rename it to ‘PEAKS’ or something familiar, and drag it onto the Desktop, or into the Start menu.

5) To run PEAKS, now we just double click this icon.

SC Connecting to a PEAKS Online Server

PEAKS Client sends all its intensive computation jobs, like PEAKS auto de novo, and PEAKS Protein ID, to a PEAKS Online server, installed somewhere in the lab or computing facility. PEAKS Studio can optionally do this too.

In the Search Engines Details tab of the Environment preferences dialogue, change the “Default PEAKS Invocation” to “PEAKS Online. Then specify the IP address or URL of the PEAKS Online server, the port number to use, and a username and email address. Press Ok. The email is used to notify us that a job has been completed, in the event that we close PEAKS 4.5 before finishing.

It’s important to monitor the task queue when running jobs on the PEAKS Online server. Hold the mouse over a job that’s in the queue to see it’s status.
SC

Registering PEAKS

The first time we run PEAKS, we will be told that the product is not registered. Press the Ok Button and a dialogue will appear. Enter the registration key that came with the product – whether it be a key for the full version or time limited trial version. We must also enter our name, the name of our organization, and the MAC (Media Access Control) address of the machine we are going to use PEAKS on. If we are connected to the internet, registration will be completed automatically. If all is well, a dialogue will show “Registration Successful” and PEAKS will load. If not connected to the internet, onscreen instructions show how to register offline, manually.

The software uses your computer’s MAC address as a unique identifier for the computer. As such, if you have more than one MAC address for your computer, you may experience problems in registering PEAKS. The display will be something like this:

If this occurs, try disabling your wireless network card, restarting windows and plug into an Ethernet cable only. Then register with PEAKS. If problems still occur, please contact BSI technical support.

Re-registering PEAKS may be necessary if our license has expired or if we wish to update the license. We will need to obtain a new registration key from BSI. Once we have obtained this new key, select “Register PEAKS” from the Help menu. The “License Upgrade” dialogue box will appear, cautioning us that we are about to update the license. Press the “Ok” button to continue. Follow the on-screen instructions.
Database Configuration

In addition to de novo sequencing of peptides, PEAKS 4.5 also has the ability to search through a database search to identify proteins. But in order to use this function, PEAKS must have access to a protein or EST database in FASTA format or an EST database of DNA sequences. We can point PEAKS to an existing database on our system, or download one. Additionally, we can associate taxonomy with certain databases. This is database configuration.

We can use PEAKS without the database search; PEAKS can perform de novo sequencing.

WARNING: Downloading a database can take a long time (8+ hours), depending on connection speed. Most only take 20 - 30 minutes.

To configure a database:

1. Load PEAKS 4.5. If we have not yet configured a database, the wizard will appear automatically. Otherwise...

2. In the ‘edit’ menu, select “Configuration”, then “Import Database Wizard”

3. The Import Database Wizard will load and ask us to select a database to download – from the dropdown list. If we already have a database we wish to use, we can select “Other database” from the dropdown list and skip to step 6. Click “Next”

4. Having selected a database, the Import Database Wizard will provide us with some information about that database. If this is, in fact, the database we wish to use, click the provided link to begin downloading. A dialogue box will appear with instructions on downloading using file transfer protocol (FTP). It does not matter where we put the download file, but we must remember where it is.
A note on downloading databases

The links in the Wizard may be outdated because the owners of those download locations may change their URL periodically. If this is the case, remove all but the domain name and browse from there.

ftp://ftp.ebi.ac.uk/pub/databases/MassSpecDB/msdb.fasta.z becomes:

ftp://ftp.ebi.ac.uk/

Best practices: configuring databases for use with X!Tandem

At the time of this writing, X!Tandem had trouble searching through large databases, and would crash. It is therefore suggested that X!Tandem only be used with small databases; or if used with a large database, a taxon should be specified. The NCBI nr and Swiss Prot databases are ideal for this purpose.

Best practices: configuring databases for use with OMSSA

At the time of this writing, we could not use OMSSA with databases that were not in NCBI format, or Swiss-Prot format, and have those results available to inChorus.

Also, a bug in OMSSA prevents us from easily using databases with OMSSA when they are stored in a folder that contains a space in its path. This creates problems when PEAKS creates temporary databases on our behalf. To avoid this, best practices suggest we put all our databases in a folder “c:\PEAKS\databases”. The folder “c:\my documents\databases” wouldn’t work because it contains a space between ‘my’ and ‘documents’. Using spaces in the database file name causes the same problem. So after we download and extract our database we should call the database file “ncbinr.fas”, or “ncbi_nr.fas” rather than “ncbi_nr.fas”.

5. The database we downloaded may be in a compressed file, perhaps a .zip or a .gz file. We must find the file and use a decompression utility, such as WinZip,
or WinRar to extract its contents. The file inside the compressed file will be a FASTA format text file (a .fas or a .fasta file).

The taxonomy options are only available if the NCBI nr or Swiss-Prot database is selected and the Apply button has been pressed, or either database was selected on a previous screen.

6. Return to the Import Database Wizard and click the “Next” button. This screen will allow us to configure the database. Click on the hyperlink next to each field for more information.

- PEAKS will ask us to enter the database nickname. This is a nickname that we chose to represent the database we are configuring. It doesn’t matter what name we enter, but we must enter at least one character.
• The Path textbox shows where the database is located. It will be blank, so we must tell PEAKS where the database is. Type the location of the file into the textbox, or we can browse to find the file on our system. We must be sure to select the FASTA database, not the compressed file of the same name (see step 5).

• If we chose one of the public standard databases (in step 3) its format style will be displayed in the advanced options box. The selected database format is shown in the dropdown list. Accession number information and the way PEAKS parses the database headers – i.e. the parsing rules – are shown in the textboxes below.

• If our database is an EST database containing DNA sequences, check the ‘EST database’ checkbox.

• If we chose an “other database” (in step 3) we must enter parsing parameters ourselves by typing in the textboxes. Alternatively, if our database format is the same as one of the public databases, we can choose to apply that database’s format when PEAKS reads our database. Select the database that is similar to ours from the dropdown list and press the apply button to fill the textboxes with the appropriate parsing rules.

  The delimiter is the character used to separate multiple headers.

• If we are configuring the NCBI nr database or the Swiss-Prot database, we may choose to point PEAKS 4.5 to the location of the taxonomy files associated with that database. Under “Taxonomy Options” we must type the location of the taxonomy files, or click browse to find the file on our system. If we do not specify these taxonomy files, we will not be able to limit our database search to a specific taxon. We can use the compressed (.zip or .gz) files; no decompression is required for the taxon files.

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<thead>
<tr>
<th>A note on choosing the taxonomy files for NCBI nr</th>
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<td>At the time of printing, the gi_taxid file was called: gi_taxid_prot.dmp.gz and the taxdmp file was called: taxdmp.zip. Select these files.</td>
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<table>
<thead>
<tr>
<th>A note on choosing the taxonomy files for Swiss Prot</th>
</tr>
</thead>
<tbody>
<tr>
<td>At the time of printing, the gi_taxid file was called: gi_taxid_prot.dmp.gz and the taxdmp file was called: taxdmp.zip. Select these files.</td>
</tr>
</tbody>
</table>
At the time of printing, the gi_taxid file was called: `spedist.txt` and the taxdmp file was called: `taxdmp.zip` – the same one as used by NCBI.

7. Press the “Finish” button to complete the database configuration.

We can repeat this process to configure a number of other databases. Once configured, a database need not be configured again unless we update the database itself.

### Trouble shooting

Some problems with a database may not appear until we run a search. While PEAKS is quite tolerant of format errors in databases, other search engines called from the inChorus tool may not be. If there is an error in the search, it will be reported in a summary screen after the work has finished. If there is a problem, check the best practices outlined in this section. If the problem persists, it is possible that the database download was corrupted; try downloading again. Please contact technical support for help.

---

**inChorus (Search Engine) Configuration**

To configure a search engine for inChorus, select from the main window toolbar, Edit -> Configuration -> Environment Preferences and go to the Search Engine Details tab. Once there, you just have to type in the location of your server or search engine. The only difficult part is knowing where this is. We recommend asking your IT person, or the person who set up the server/software. Note that you usually don't have to worry about preceding or trailing slashes. You can use your regular user account for the username and password. It doesn't necessarily have to be an administrator account. *However, some of the built in user accounts like 'daemon' have some special properties that cause problems here. Don't use the daemon account, but you can use an account in the daemon group*. The email is not used by PEAKS but can be requested by the Search Engine server.

(For further explanation, refer to Chapter 5, PEAKS Configuration > Search Engine Details).
Features Walkthrough

Let’s familiarize ourselves with PEAKS

This section of the manual will walk us through most of the basic functionality of PEAKS 4.5. After completing this section we will have seen how easy it is to load and view a data file, perform de novo sequencing, and database search protein identification.

SC

Begin the walkthrough

Run PEAKS 4.5, then download and configure the SwissProt database. The procedures for doing so are outlined in the previous section.

The demo sample data should load automatically on startup under the heading OrbiSample.anz. If it is not loaded, open the data file by clicking the icon on the toolbar in the upper left corner of the PEAKS window, or selecting “Open” from the “File” menu. Sample data is located in the PEAKS program folder – for example “C:\Program Files\PEAKS Studio\data\”. Load the file “OrbiSample.anz” by clicking on it, then clicking “Open.”.

The data file will appear in the left-hand frame. Make sure “OrbiSample.anz” (i.e. the data file) is selected.

In the “Tools” menu, select “PEAKS Protein ID”. The protein identification options dialogue will appear.
Enter the settings as shown. Settings can be changed by clicking on the drop down list and selecting one of the options.

After entering the settings, as shown, click the “Save As” button to save these parameters for future use (PEAKS Studio only). When prompted, type OrbiSample and press enter. Click “OK” to commence analysis. Analysis will be initialized (most of this time is spent creating a partial database, which only has to be done once) this takes one or two minutes.

After this, de novo sequencing will commence. For this sample it takes just over a minute, after which PEAKS database search will proceed. In total, the process takes
less than a minute for this sample (depending on the system’s processing speed and memory). The PEAKS auto de novo algorithm derives sequence candidates for each of the 14 spectra in our example data file. These sequence candidates are then used for the database search component of PEAKS 4.5. PEAKS uses a unique sequence tag plus peptide fragment fingerprinting approach to protein identification.

### SVC Viewing Results

After the search is finished, the protein identification results will appear on screen. The ‘Peptide View’ is displayed by default. The display shows each spectrum for which PEAKS found a matching peptide. The spectra are grouped/sorted by index number. Since a spectrum may match to more than one peptide, there may be more than one entry per spectrum. The list is sort-able; click the heading on each column to experiment with sorting by score, by mass, etc.

Click the ‘Protein View’ tab. PEAKS 4.5 presents a list of proteins that it believes to be the best match for the sample. The top section is an index, listing them by accession number, ranked in descending order from highest score on downward.

The correct proteins, Serum Albumin and Serotransferrin, are shown at the top of the list and with high score. Since one cannot distinguish between the different forms of Albumin, PEAKS 4.5 groups them all together, thus avoiding cluttering the report. Click the plus sign next to the top entry for a listing of other possible Albumin. The peptides matching these homologues will be the same set or a subset. Collapse this list of homologues by clicking the minus sign.

The listing, as shown above, is simply an index. We will find this useful, in the future, when dealing with complex mixtures. Clicking on any protein will display the peptides matched to that protein in the bottom pane.
Above, 7 of the original 14 spectra indicated a peptide sequence matching with Bovine Serum Albumin. Most peptide matches show a high confidence – strong evidence for having found the correct protein. We can also see exactly where the peptide fits into the protein sequence; with the matching sequences highlighted in red at the bottom.

As mentioned above, the peptide sequence results are based on a database search guided by an initial de novo sequencing analysis. The RSD number listed here measures how well the de novo sequence agreed with the peptide returned from the database. In most cases it is zero! so we can have that much more confidence in the answers. The last peptide didn’t quite agree. Let’s see how where the difference lies. Click on the last entry (number 13); this will bring up the main processing window for that spectrum 871.41 2. Look in the top left frame to see the de novo and database results.

The RSD measure was proposed by Sergey Pevtsov, Irina Fedulova, Hamid Mirzaei, Charles Buck, and Xiang Zhang. J. Proteome Res., 5 (11), 3018 -3028, 2006

Color coding shows positional confidence scores. We can see that the PEAKS auto de novo analysis returned, with >99% confidence, the partial peptide sequence (OxM)P(CarboxymethylC)TEDYLSLILNR, but was not as sure of the middle residues. The PEAKS DB Search was able to confirm this result, returning the peptide: (OxM)P(CarboxymethylC)TEDYLSSLILNR. Clicking on either sequence will highlight supporting ions on the original profile data.
Selecting another spectrum from the Peptide Data frame (left), e.g. 537.25 2, will allow us to view the results from that spectrum without having to return to the protein identification result. Click on the time and date stamp beneath the filename to return to the report. Now try clicking on the Peptide View tab to see the same data, but simply listed by peptide.

**SVC Manipulating Results**

We already know that the sample was a simple digest of Albumin and Serotransferrin; but let’s pretend we’re only interested in Serotransferrin. Click on the Filter Pane tab. Here we can add a filter that shows only proteins with mass greater than 70000 or any other value we choose. Scroll down the Possible Filters list and select the Protein Filter called Mass Filter. In the [Edit Filter] frame, remember to select "Greater than" and type 70000, then press the Add Filter button. The filter now appears in the list of Selected Filters. The filter can be edited while in either listing and can be added multiple times (in case, you only wanted proteins with a mass greater than 700000 but less than 90000). Hit the Apply Filters button and let’s examine the results. Click the ‘Protein View’ tab.

![Protein View Tab](image)

Ok, so we removed the other proteins, and we’re left with Serotransferrin and a few variants (click the + sign to see them).

![Variant View](image)

Remember, all proteins in this group contain the same set or a subset of the same peptides. We’ll see that there are still some hits down there that aren’t very relevant. Yes, they contain a good peptide, but only one. Let’s see if we can remove these one hit wonders. Go back to the Filter Pane tab. The Selected Filters list still shows our Protein Mass Filter > 70000. Let’s add one to it. Scroll down the list of Possible filters and select the protein filter called Query filter. In [Edit Filter], select the ‘Greater than’ option and type ‘1’ in the box, then press the Add Filter button.
Now both filters appear in the list. Press the Apply Filters button to use them. Going back to the Protein View tab, and expanding the plus sign reveals that we are only left with variants of Serotransferrin.

**SC inChorus Searching**

Let’s try another kind of search. This time we’ll use inChorus database searching – this technology, unique to PEAKS, allows us to launch other search engines that will help improve the results. The best confirmation of results comes from using two or more methods to confirm the peptide matches.

Select ‘OrbiSample.anz’ from the Peptide Data frame (left) and choose ‘inChorus protein ID’ from the Tools menu. The inChorus Database search dialogue appears:
Make sure that “PEAKS Protein ID” and “X!Tandem Search” are selected. Notice that there are six ‘Options’ icons on the right. They correspond to each search engine.

Click the PEAKS Protein ID Options button. The options pane is similar to the one we’ve seen already. The settings that we used before should still be there. If not, select “OrbiSample” from the drop down list in the top right corner. Before pressing the “OK” button, we can make one change. Since we already have de novo sequencing results, we don’t need to do de novo sequencing again. Click the option ‘I have already run de novo, don’t do it again’, then press the OK button.

Click the X!Tandem search options button (top). This window allows us to set options for the X!Tandem search tool. This window is set up to behave almost exactly the same as the X!Tandem interface, so it may look familiar. In section 2, set the fragment error to 0.02. In section 3, choose ‘Carboxymethyl (C)” as a complete mod, and “Oxidation (M)” as a potential mod. Under “7. Predefined methods” choose FTICR. To learn more about X!Tandem settings, double-click any of the question marks. Press the OK button.

Now that we’ve set everything up for the inChorus search, press the “Launch inChorus” button on the bottom of the search dialogue pane. inChorus will call on each search engine, wait until they are finished, then compile their results together, ensuring the integrity of the data-results relationship. It will do a final pass through a common database to reconstruct the protein identities.

Watch the task queue (bottom-left of PEAKS). After everything is finished, new search results will appear in the Peptide Data frame (left), stamped with the date and time. The task queue will be empty, and the results will display. There’s also a nice little report to tell us if there were any errors.
PEAKS displays the same ‘Peptide View’ as before. Now, XTandem search results are considered as well. Where XTandem agrees with a PEAKS assignment, there’s a checkmark in the PEAKS column and a checkmark in the XTandem column. The search engine's individual score is presented, and a meta-score is calculated by examining the two – in this way we increase confidence in the assignment. Since the two tools take different approaches, we may discover that PEAKS finds some peptides that XTandem misses, and vice-versa. Where this is the case, only one checkmark will be displayed and the score is penalized slightly in some cases. Sometimes we can find a good hit that the other search engine would have missed. In this way we increase coverage. On this remarkably clean data, it’s not surprising that both search engines found everything.

**SC**

**SPIDER searching for homologues**

But wait! We missed a peptide – spectrum number 8. In a larger sample, this would be hard to spot, but fortunately we can filter for it. Go to the Filter Pane Tab, and remove all the filters we had selected. Then choose the option shown on the right. Basically, this makes the De Novo View tab show spectra that could not be explained.

Click the Apply Filters Button, then hop over to the De Novo tab. There’s our peptide! The de novo sequence seems to be a good one, and matches the spectrum quite well, but we couldn’t match it to the database because it is a mutated peptide from Serotransferrin.

To identify it, we’ll use SPIDER. But first we have to create a little database for SPIDER to search through. Go back to the Protein View tab for a second. Mark all the proteins, and right click on either of them. In the popup window that appears, choose Export DB.
Save this new database as temp.fasta and press the Ok button. It is a miniature database containing only the protein sequences we’ve marked – only Albumin and Serotransferrin. Now we can use SPIDER to help consolidate the mutant with the known sequence of the protein we’ve already identified.

Select the spectrum of interest, 680.3 2, by clicking in the Peptide Data Tree (left), or by selecting it’s entry on the De Novo View tab. Now we’ll run SPIDER; click the icon on the toolbar, or choose Tools -> SPIDER Search. The SPIDER Parameters window will appear:

Enter the settings as shown. For the “Database to search:” we’ll have to add our database by clicking the New Database Button.

Press the Ok button. After a few seconds, SPIDER returns a match, to Serotransferrin. Click the Peptide View Tab to see how the mutated peptide aligns with the known peptide in the database.

**Merging reports**

Adding this SPIDER hit back together with our PEAKS Protein ID and X!Tandem search results will give us a complete picture of the protein. To do so, we just run the inChorus feature again. The difference is that now, we’ll be using existing results. Select the filename again (OrbiSample.anz), and click the inChorus icon on the toolbar.
Check PEAKS and SPIDER. For PEAKS, press the Import… Button, and choose the existing PEAKS result that contains PEAKS and X!Tandem. For SPIDER choose the SPIDER results we just finished.

Press Launch inChorus on the inChorus window. PEAKS will display the results in a few seconds. Now, importantly, the new SPIDER peptide is shown in red on the Serotransferrin sequence:

Thus concludes our walkthrough of PEAKS 4.5’s basic features.
This chapter deals with interface elements. It isn’t meant to be read from start to finish; it is to be used as a reference so we can look up certain interface elements when we get stuck. For instructions on how to use PEAKS to perform certain tasks, the Chapter entitled “Analyzing data with PEAKS 4.5” will be more instructive.

The first part of this chapter describes windows, dialogues, frames and reports. This tells us what certain dialogue boxes, windows and frames do and how to read them.

The second part of this chapter deals with toolbars. Toolbars are a very useful way to quickly get at the functions we use most.

Graphical User Interface

A reference section to help us find our way around.
SVC Windows, Dialogues, Frames and Reports

PEAKS 4.5 main window

Comprises:

- **Peptide data frame (left):** This displays a listing of parent ions by m/z and charge. Clicking on one will bring up its MS/MS spectrum. The colored dot by each spectrum appears dark green for unprocessed, or light green for sequenced (or partially sequenced). An asterisk (*) next to a spectrum shows that it contains unsaved information.

  Spectra are grouped by data files, or by nodes (which act like data files). Select a data file or node by clicking on its name (i.e. click on CytC-ESI.anz in the above example), or a spectrum within a data file by clicking on it. Use the ‘ + ’ and ‘ – ’ boxes to expand and collapse the view.

- **Task Queue frame (bottom left):** Shows running tasks, sorted by priority.

S marks a section applicable to PEAKS Studio, C for PEAKS client and V for PEAKS Viewer
- **Working area (right):** This is where the Protein Identification Result Window and the Main Processing windows appear.

- **Menu bar:** access file, edit, view, tools, windows, and help commands.

- **Main window toolbar:** quick access to many commands. See “Toolbars” section below

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**Auto de Novo Parameters Dialogue**

- **Instrument:** choose the type of spectrometer that produced our data. Choose from a dropdown list. To edit or add a new instrument to the list, press the Edit Button adjacent to the dropdown list.

- **Parent mass error tolerance:** determines how much random and systematic experimental error on the parent/precursor ion PEAKS will account for in its analysis. Type a number in the textbox. Select the units (PPM or Daltons) from the dropdown list.

- **Fragment mass error tolerance:** determines how much random and systematic experimental error on the fragment/daughter ion PEAKS will account for in its analysis. Type a number in the textbox. Daltons are the units here.
- **Enzyme**: choose from a dropdown list of enzymes that we used to digest our protein sample. Click the “Edit Enzymes” button to edit the enzymes defined in this list, or to add to it.

- **Report up to**: set how many de novo sequence candidates PEAKS will report. Choose from a dropdown list.

- **PTM selected for search**: this box displays the modifications currently selected for analysis, these will be considered during auto de novo sequencing. To change this, click the Add/Remove PTM button.

- **Max variable PTM per peptide**: this allows us to restrict the number of variable PTM that will appear on any given peptide.

**Instrument**: choose the type of spectrometer that produced our data. Choose from a dropdown list. To edit or add a new instrument to the list, press the Edit Button adjacent to the dropdown list.

**Parent mass error tolerance**: determines how much random and systematic experimental error on the parent/precursor ion PEAKS will account for in its analysis. Type a number in the textbox. Select the units (PPM or Daltons) from the dropdown list.
- **Fragment mass error tolerance**: determines how much random and systematic experimental error on the fragment/daughter ion PEAKS will account for in its analysis. Type a number in the textbox. Daltons are the units here.

- **Enzyme**: choose from a dropdown list of enzymes that we used to digest our protein sample. Click the “Edit Enzymes” button to edit the enzymes defined in this list, or to add to it.

- **PTM selected for search**: this box displays the modifications currently selected for analysis, these will be considered during database searching. To change this, click the Add/Remove PTM button.

- **Database to search**: This dropdown allows us to choose which FASTA format database to search

- **Taxonomy selection**: if the ‘Database to search’ has been configured for taxon based searching, this list will allow us to select limit which taxa are searched.

- **Paste fasta sequences**: Paste a few sequences in this box and PEAKS 4.5 will search through those sequences as opposed to the database selected.

- **Preprocess this data ‘on the fly’**: PEAKS has its own built-in preprocessor for removing noise, centroiding, and peak charge recognition from MS/MS data. Check this box to turn preprocessing on.

- **Advanced options (de novo sequencing)**: The PEAKS approach to protein identification uses de novo sequences to help out in the search. This section allows you to decide how to obtain the de novo sequences required for the search.
- **Enzyme list tab**: Displays a list of built-in and user-defined enzymes. We may edit and create enzymes from here.

- **PTM library tab**: Displays a list of built-in and user-defined PTMs. We may edit and create PTM from here.

- **Database tab**: Displays a list of databases available to PEAKS. We may make new databases available to PEAKS from here.

- **Instrument list tab**: Displays a list of instrument types and acquisition modes available to PEAKS. New instruments/modes can be defined from here.

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**SCV Enzyme Editor Dialogue**

- **Digestion Rules**: This is how we specify where our enzyme will cleave the protein between two amino acids to create peptides. Use set brackets {} around a residue to denote “any amino acid except the ones enclosed in these brackets”. Use X to denote “any residue”. Listing several amino acids in one box means “any one of these residues”.

- **Specificity Parameters**: Peptides can break down such that only one end is a cleavage site. Check the boxes to tell PEAKS to search for only for peptides
that have proper cleavage sites on both ends, or to require that only one end be a proper cleavage site.

- **Shorthand notation:** Advanced users may specify their enzyme cleavage in shorthand notation, but it is not required.

- **Saving/Loading Enzymes:** After setting up an enzyme, we can save it for future use. Click the "Save Parameters" button, and choose a name for future reference if prompted. Don't worry, we can't accidently overwrite the defaults. Any enzyme we save will be available in the drop-down list at the top of the window. To see what's inside, just select one, and the enzymes digest rules boxes will be populated.

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### SC

This dialogue allows us to create or edit a PTM

Here we can form a list of available post-translational modifications. We can choose any PTM as Fixed PTM or Varied PTM (to tell PEAKS that it may or may not occur). To make this selection, click on a PTM in the list at left, and then click the “Select As Fixed=>” or the “Select As Varied=>” Button. If a PTM is already selected as a fixed PTM, it cannot be selected as varied PTM, and vice versa.

If we change our mind about a PTM after having selected it, it is still possible to unselect it. Click the erroneous PTM from the list of ‘Selected Fixed PTM’ or ‘Selected Varied PTM’, and then click the “<=Unselect” Button to remove it from either list of Selected PTM.
PEAKS software ships with some pre-defined PTMs. These are listed as <Built-In>. If we want to create a new PTM, we can click ‘New PTM’ to create a new one. The **Editing a PTM** and **Creating a New PTM** sections below describes how this is done.

### SCV

**This dialogue allows us to create or edit a PTM**

#### SCV PTM Editing Dialogue

- **Name:** This will appear in the PTM list
- **Abbreviation:** This will appear in the auto *de novo* results, if it is found.
- **Mass (monoisotopic):** The mass that the residue gains or loses as a result of the PTM. Enter this numerically here, or enter the chemical formula below.
- **Neutral Loss Mass:** The mass that the modified residue loses as a result of fragmentation. E.g. 28 would signify a loss of 28 Daltons.
- **Formula:** The chemical formula of the PTM. This will automatically enter the mass.
- **Residues that can be modified:** Enter residues that can be modified anywhere, residues that can only be modified if they are at the N-terminus and residues that can only be modified at the C-terminus, and residues that can only be modified if they are not on either terminus.
- **Rule:** user entered, a comment for our reference.

### Instrument Definition Window

This window allows us to define new instrument configurations for use by PEAKS algorithms. Users are free to set any ionization mode, fragmentation method and mass analyzer. Also, we can set suggested default error tolerances, and make notes for all our instruments.
- **Dropdown-List of Saved Instruments**: Select from this list of built in and user defined instruments to display it's properties.

- **Ion Source**: define an instrument with a MALDI/SELDI source to tell PEAKS Data Refine tools that this data will always have parent charge of +1.

- **MS - Precursor Scan**: This selection will tell PEAKS Data Refine tools if the survey scan is of sufficient resolution to determine charge and monoisotopic peak from examination of the survey scan.

- **Fragmentation Type**: This selection will tell PEAKS what type of ion-series to expect for PEAKS auto de novo sequencing and PEAKS protein ID database search. Select CID/ECD if alternating fragmentation is used, allowing the algorithm to determine fragmentation from each scan header.

- **MSⁿ or MS/MS - Fragment Scan**: This selection will help PEAKS decide which internal parameters (for weighting fragments and amount of noise) to use during PEAKS auto de novo sequencing and PEAKS protein ID database search. Select LIT/FT if alternating hi-res/low-res mode is used, allowing the algorithm to determine mass analyzer from the scan header.

- **Nickname**: A name, for our reference only. We’ll select the instrument, using this nickname, for the various PEAKS tools.

- **Precursor Mass Search Type**: For ion-trap instruments, it is sometimes beneficial to allow the PEAKS Protein ID database search to use an average mass.
- **Default Error Tolerances:** The values set here will be suggested on the *PEAKS Protein ID* and *PEAKS auto de novo* options screens when the instrument is selected.

- **Notes:** For our reference only.

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**SCV Environment Preferences Window**

This window controls the basic, user configurable environment. Colours, displays, file handling and default settings can all be changed from here. The window contains 8 tabs:

- **Parameters tab:** Our saved parameters for de novo sequencing and protein ID are shown and can be deleted from here.

- **Basic Ion Table Editor tab:** allows configuration of the basic ion table

- **Advanced Ion Table Editor tab:** allows configuration of the advanced ion table

- **Environment tab:** Allows us to set default file locations, default file handling functions, and miscellaneous behaviour defaults.

- **Colour tab:** from here we can set the colours to use for various spectrum annotations.

- **Display tab:** from here we can set the type-face to use across the PEAKS 4.5 user interface. The width of spectrum PEAKS can also be set from here.

- **Manual De Novo tab:** default options for manual de novo sequencing can be set from here.

- **Search engines details:** allows us to define where the search engines are located on our system, and/or where the servers are located.

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**SCV Protein Identification Result Window**

The protein identification result window contains the results from one protein identification run on one data set. It is organized into three tabs: *peptide view, protein view* and *search parameters.*
De Novo View
The De Novo View summarizes the de novo sequencing results for each spectrum in the data file. The exact contents of the list are determined by filters set on the Filter Pane. By default the peptides are grouped by spectrum, but the list is sort-able by any of the columns in the table. Highlighting one or more (click and drag, or use shift+click) items in this list selects those spectra in the peptide data tree (found on the left hand side).

Peptide View
The Peptide View summarizes the results for each MS/MS spectrum. All peptides that match to each spectrum are displayed. The exact contents of the list are determined by filters set on the Filter Pane. By default the peptides are grouped by spectrum, but the list is sort-able by any of the columns in the table. Highlighting one or more (click and drag, or use shift+click) items in this list selects those spectra in the peptide data tree (found on the left hand side). The bottom panel of this view shows more details about the peptide identification that is highlighted in the top section. In this view, we can see the protein that the peptide came from, and a simple alignment between the original de novo sequence for this spectrum (if available) and the peptide found in the database.

Protein View
The protein view is most useful as a summary of what proteins were present in a sample, and the peptides matched to them. It has two sections:
- **Index (top section):** PEAKS presents a list of proteins that it believes to be the best match for the sample. This index lists them by accession number, ranked in descending order by score. Very similar proteins (i.e., those that contain most of or all of the sequences identified by PEAKS) are grouped together—only the first entry in this group is shown here. Show the whole group by clicking the + sign. In the example below, lactoglobulin beta is the top ranked protein candidate.

- **Peptide Match Reports (bottom section):** PEAKS presents each protein candidate with a peptide match list beneath it. Each peptide that matched the protein sequence is shown, in order, by spectrum. The confidence that the correct peptide sequence was found is displayed next to each peptide sequence. At the bottom of this list, the complete protein sequence is shown with matching peptides highlighted in red.

- **MSA Pane (bottom section):** This panel displays a multiple sequence alignment between any protein that is 'marked' in the Index (top section). The MSA is only displayed when the button is pressed.
**Search parameters**
This tab displays the protein identification parameters that were used to guide the search that generated these results. For inChorus searching, all parameters used for each search engine are exposed here.

**Filter Pane**
The Filter Pane allows us to control what to display on the other tabs of the multi-part protein ID report by setting up customizable filters. The various parts of this screen are described as follows:

- **Possible Filters list**: a list of the filters, currently available, that can be used to build a particular filter set.

- **Active Filters list**: a list of the filters, currently selected, that will be used to filter your results.

- **Add Filter Button**: clone the currently selected filter in the Possible Filters listing to the Active Filters.

- **Apply Filter Button**: Apply your filters; any/all selected filters will not be used until this button is actually pressed.

- **Edit Filter Frame**: Edit the settings for the currently selected filter. Filters in the Possible Filters list and the Active Filters list can be edited.
**Options frame:** The options listed here fundamentally change what is listed in the De Novo View, regardless of the specific filters selected.

- **Option 1:** De Novo View shows peptides that could not be explained by peptides from the Peptide View – In this mode, the De Novo View only shows peptides for spectra that could not be satisfactorily explained by searching the database. As such, removing a peptide/spectrum from the database search results – the Peptide View list – will cause the de novo sequence for that spectrum to **appear on the De Novo View.** The de novo peptides still have to pass the filters defined for the De Novo View.

- **Option 2:** De Novo View shows all peptides that are not filtered – In this mode, the De Novo View shows de novo sequences for all spectra that have not been removed by filtering out database peptides or proteins. As such, removing a peptide/spectrum from the database search results – the Peptide View list – will cause the de novo sequence for that spectrum to be **removed from the De Novo View.** The de novo peptides still have to pass the filters defined for the De Novo View.

- **Checkbox:** remove de novo peptides with no database hits – This option removes all peptides that have no corresponding database peptide hits. Selecting Option 2 (above) and this checkbox shows essentially the "reverse" of Option 1.

**Save As Button**– Save and re-use your filters between different sessions of PEAKS. All active filters and selections from the Options Frame are saved.

**Delete Set Button** – Remove the selected filter from the list of saved parameters.

**Set Saved as Default Button**– Select a saved filter to be automatically applied to every report we load.

See the section “Post analysis of results – preparing for publication” for help on using filters.
Main Processing Window

- **Main Processing Window Toolbar**: quick access to many processing functions. See “Toolbars” in the next section.

- **Peptide Candidates Frame (top left)**: PEAKS shows peptide sequence candidates, ranked by score for the selected spectrum. Peptide sequences are grouped by the headings: “Auto de novo”, “Manual de novo”, “user defined result type” and “database search results” depending on how they were derived. For de novo results, positional confidence is color coded on each residue. More specific positional confidence appears when the mouse is held over a sequence – this shows the confidence in each of its parts.

- Under the “Manual de novo” heading, the masses of the residues are displayed. That is to say, the mass of the whole peptide minus water.
The Ion Table Frame (top right): The Ion Table shows the proposed ions with their corresponding masses -- i.e. the mass of the b1 ion is shown in the top right corner. The default Ion Table will display b, a, immonium, yH2O, yNH3 and y ions in basic mode; it will display b, b-H2O, a, c, immonium, y, y-H2O, z, z’ and y(2+) ions in advanced mode. To switch from basic mode to advanced mode, choose ‘Show ion table’ from the ‘View’ menu.

The Ion Table Frame also contains an error plot (it may be necessary to scroll down to see the error plot). The error plot shows the confidence each ion is assigned. The most confident results lie on the centerline. Clicking a cell or column in the Ion Table highlights the corresponding points on the error plot and corresponding PEAKS on the spectrum.

Spectrum View Frame (middle): Shows a graphical representation of the spectrum. Peak masses are labeled, as are the PEAKS associated with identified ions. We can zoom in on the spectrum by clicking and dragging over an area.

Spectrum Alignment Frame (bottom): Shows a graphical representation of the spectrum. This view always shows the whole spectrum and is used as a tool to help navigate the spectrum view frame. A blue bar along the horizontal axis of the alignment view indicates the range of the spectrum view in the Spectrum View Frame.

The Spectrum Alignment Frame can also show the positions of major ions that delimit the proposed sequence. By default, the Spectrum Alignment Frame displays b-ion and y-ion PEAKS and the derived peptide sequence between them. The Spectrum Alignment Frame can also show the position of c-ion and z-ion PEAKS.
The Ion Editor is used when performing manual de novo sequencing.

- **Selected peak information**: displays information about the currently selected peak.

- Under “Please choose ion type”, the radio buttons set whether the ions in the “ion choice list” are C terminal ions or N terminal ions.

- **Ion choice list: (left)**: lists the ions we can apply to the selected peak.

- **Selected ion list (right)**: lists the ions we have selected add or remove them using the “Add” and “Remove” buttons.

- **Apply button**: applies the ions in the “selected ion list” to the selected peak.
SCV

Export Image Dialogue

- **File Format**: select an image file format from the drop down list. Bitmap, JPEG, PNG, GIF and SVG are supported.

- **Width and Height**: together these determine the size of the output image (measured in pixels).

- **Filename**: type in the textbox, or browse to a file, to enter the file name of the image that will be created.

- **Resolution**: in most cases the image resolution can be upscaled for printing large pictures. More resolution (i.e. more pixels) means a higher quality picture when blown up or printed.

- **Image types**: Select one of these options to choose the image that will be output. The ‘annotated spectrum with alignment’ will suit most purposes.
**SCV Print Image Dialogue**

- **Orientation**: paper orientation is shown in the picture at the top. Change this by clicking the “Portrait” or “Landscape” radio buttons.

- **Paper**: Set the paper size and source by selecting from the appropriate dropdown list.

- **“Printer” button**: pressing this will bring up another dialogue where we can select from a list of printers installed on our machine.

- **“OK” button**: this will commence printing.

The default print output is the full spectrum as shown in the spectrum alignment frame. If we wish to print something else, we must use the export image functions and then print the image from another application.

**SVC Toolbars**

**SCV Main window toolbar**

- **"Open data file"** button: This allows us to open a raw data file built by our mass spectrometer, or a PEAKS data file (in ANZ format) that also contains peptide analysis data. The file should be in PKL, DTA, MGF or ANZ format.

- **"Close data file"** button: Close the selected data file. Press this after selecting a data file in the Peptide Data Frame.

- **"Save data file"** button: Save any changes made to the file (a “*” will appear next to any file that has been changed and not yet saved). The file will be saved in the ANZ format. Press this after selecting a data file in the Peptide Data Frame.
"Save all files" button: Save all files. Any changes to files will be saved in the ANZ format.

“Copy” button: Copy selected spectrum data.

“Cut” button: Cut selected spectrum data.

“Paste” button: Paste spectrum data into the selected data file.

"Manual merge spectra" button: After selecting more than one spectrum in the peptide data tree, this button becomes enabled. Right click it to merge these spectra into a single MS/MS spectrum, and remove the old ones.

"Data Refinement" button: Merge scans of the same peptide, remove noise spectra, preprocess within each MS/MS spectrum and recover peptide charge state. The data refinement options dialogue will allow us to choose and to set parameters for each of these refinement tools.

"Auto De Novo" button: perform auto de novo for a selected data file, spectrum or list of data files. Press this after selecting one or more data files (or spectra) in the Peptide Data Frame. An auto de novo options dialogue will allow us to set parameters before we begin.

"PEAKS Protein ID" button: perform protein identification a selected data file. Press this after selecting one or more data files (or spectra) in the Peptide Data Frame. A protein identification options dialogue will allow us to set parameters before we begin.

"inChorus Protein Identification" button: perform protein identification on a selected data file using multiple search engines. Press this after selecting a data file in the Peptide Data Frame. A protein identification options dialogue will allow us to set parameters before we begin.

"Environment Preference Configuration" button: configure the environment, spectrum color-coding, and manual de novo parameters.

“PEAKS Properties Configuration” button: define PTM, Enzymes, and add FASTA protein or EST databases.

“Import Database Wizard” button: help user download and configure database.
Main Processing Window Toolbar

“y ion Alignment” button: toggle (show/hide) the location of PEAKS corresponding to y-ions and the corresponding proposed peptides between them.

“b ion Alignment” button: toggle (show/hide) the location of PEAKS corresponding to b-ions and the corresponding proposed peptides between them.

“Deconvolute” button: toggle (on/off) deconvolution of the mass spectrum scale.

“1:1 zoom” button: return spectrum to original 1:1 zoom.

“Undo Zoom” button: return to previous zoom ratio.

“Edit Ion” button: set or edit the type of ion associated with a peak in manual de novo. Press this button after having selected a peak in the spectrum view frame.

“Next Peptide” button: redo changes to the peptide in manual de novo.

“Previous Peptide” button: undo changes to the peptide in manual de novo.

“Export Results” button: export the spectrum view, ion table, or to a picture (bmp, gif, or jpg format) with ions, masses, PEAKS and peptides marked.

“Print Results” button: print the spectrum view with ions, masses, PEAKS and peptides marked.

“View Results” button: show, in HTML format: the spectrum view with ions, masses, PEAKS and peptides marked; peptides and confidence scores; the ion table; and the error plot.
Chapter 5

PEAKS Configuration

How to set up PEAKS just the way we like it.

This chapter deals with configuration. PEAKS 4.5 is a versatile and flexible tool. But in order to use the software to its full extent, we must learn how to configure it to make it do what we want it to. Additionally, PEAKS 4.5 allows us to set up many defaults and presets to help us be quick and precise.

We can use PEAKS 4.5 without the need to configure; default settings will be used. However, to increase efficiency we should set environmental preferences, and PEAKS properties. This will enable us to customize the tool to our requirements. It is recommended that we configure PEAKS 4.5 before processing data files.
PEAKS Properties Configuration

One of PEAKS 4.5 preferences, PEAKS Properties configuration sets the parameters that the algorithm will use while processing our data files. PEAKS properties include: enzyme, PTM and database. When using PEAKS Client, all PEAKS properties are controlled by the server, please contact your administrator. PEAKS Studio provides tools to edit PEAKS properties for convenient use in de novo sequencing and protein identification in PEAKS Studio. To edit PEAKS Properties:

- Click the icon in the main window toolbar;
- Or from the “Edit” menu, select “Configuration” then “PEAKS Properties”

The PEAKS Properties dialogue will then appear. This dialogue box has three tabs: “Enzyme list”, “PTM Library” and “Database”. Clicking a tab will allow us to edit the PEAKS Properties corresponding to that tab. We can also import or export our preferences to/from a file.

Creating and Defining Enzymes

PEAKS allows us to use almost any enzyme, or combination of enzymes in our analysis. Identified/sequenced peptides will be limited to those conforming to the enzyme’s cleavage rules. Users can select from any of the built in enzymes, or define their own. PEAKS Client uses the enzyme list defined on the server, editing here will have no effect on the list available during analysis.

All enzymes are listed here including <built-in> and user defined. From here, we can create a new enzyme, edit an existing enzyme or remove an enzyme from the list. See the sections below for help with these operations. Built-in PTM cannot be removed from the list, but can be edited.

Creating/Editing an Enzyme

To define an enzyme, we open the PEAKS Properties dialogue, ensure that the ‘enzyme’ tab is selected and click the New button, or select one from the list and press the Edit button. To define an enzyme ‘on the fly’ while setting up PEAKS auto de novo or PEAKS Protein ID, click the Edit Enzymes button. The Enzyme Definition dialogue will appear.
Now simply type/edit the digestion rules for our enzyme or set of enzymes by specifying what residues can appear at the end of a peptide and what can appear at the beginning. The example here shows a combination of Trypsin and Asp-N. The letter X denotes ‘any amino acid in this position’, while {set brackets} indicate ‘any amino acid except the one in the brackets’. Furthermore, on this window, we can set specificity rules. For example, a peptide may degrade after digestion, and so would no longer conform to the rules. The specificity allows us to set ‘semi-specific’ cleavage, denoting that the peptide should conform to the rules at one end and/or the other, not necessarily both. Click the checkboxes to specify.

After defining the digestion and specificity rules, press the Save Enzyme Button to save it. We will be prompted for a name. This name will appear in the enzyme list, and will be available so we can use it later.

**Creating and defining PTM**

If we know that our sample protein may have been modified since translation, we need to apply this information to our analysis. To edit the list of PTM available to PEAKS we open the PEAKS Properties dialogue and select the PTM Library tab. **PEAKS Client uses the PTM library defined on the server, editing here will have no effect on the list available during analysis.**

All PTM are listed here including <built-in> PTM and user defined PTM. From here, we can create a new PTM, edit an existing PTM or remove a PTM from the list. See the sections below for help with these operations. **Built-in PTM cannot be removed from the list, but can be edited.**
Editing a Built in PTM

It is possible to modify a built in PTM. PEAKS will save the modification and treat this PTM as a customized PTM. It will temporarily overwrite the built-in PTM - we will not be able to see the original, built-in, PTM until we remove the customized one. We can remove this customized PTM at any time and the built-in PTM will reappear.

Creating a New PTM

To create a new PTM, we open the PEAKS Properties dialogue, ensure that the ‘PTM’ tab is selected and click the ‘new’ button. To create a new PTM ‘on the fly’ while setting up PEAKS auto de novo or PEAKS Protein ID, click the “new PTM” button. The PTM Editing dialogue will appear:

Now we type information pertaining to our PTM in the appropriate boxes (see above section on Interface for a more in depth explanation of these fields). At a minimum, we must enter a name, a mass and one residue that may be modified.

Enter the mass of the modification either by typing in its monoisotopic mass difference directly, or by entering its empirical formula. It is unnecessary to do both; each will override the other.

Click the “OK” button to save changes and create our new PTM, or click the “Cancel” button to exit, discarding changes.

After clicking the “OK” button we return to the PEAKS Properties dialogue to find that our new PTM is listed at the top of the PTM list.

Editing a PTM

To edit a PTM, we open the PEAKS Properties dialogue, ensure that the ‘PTM’ tab is selected, and select a PTM from the list by clicking on it and click the ‘Edit’ button. To
edit a PTM ‘on-the-fly’ while setting up PEAKS auto de novo or PEAKS Protein ID, click the “add/remove PTM” button, to bring up the Modification window for that search, then click the “new PTM” button. The PTM Editing dialogue will appear. Now we follow the same procedure (see above) as we would if creating a new PTM.

Removing a PTM
To remove a PTM, we open the PEAKS Properties dialogue, ensure that the ‘PTM’ tab is selected, select a PTM from the list by clicking on it, and click the ‘Remove’ button. Any built-in PTM cannot be removed.

Database Manager
PEAKS 4.5 needs a protein or EST database (in FASTA format) to identify protein candidates. Since databases are being constantly updated, PEAKS does not ship with a protein or EST database. Thus, we need to download it from the Internet and tell PEAKS where the database is located. PEAKS provides the Database Manager as a tool to help us do this. To see a list of databases available to PEAKS 4.5, load the PEAKS Properties dialogue and click the ‘Database’ tab. From here we can edit a database’s properties, load a new database, or remove an existing database.

PEAKS Client uses the databases configured on the PEAKS Online server.

Load/Configure a new database
For an in depth look at configuring a database, see the Database Configuration section in Chapter 2.

To configure a new database, we open the PEAKS Properties dialogue, ensure that the ‘Database’ tab is selected and press the “New” button. Now we open up our web browser to find a database to download. Find one, download it, and unpack it. If taxonomy is available for this database, download those files too. Return to PEAKS 4.5 and find the file on our system where we unpacked it. Name the database and select the header format to use (or we can define our own). If taxonomy is available for the database, find those files too. Click “OK”. The new database will now appear, listed by our chosen name, in the list of databases.
**Remove a Database**
To remove a database, we open the PEAKS Properties dialogue, ensure that the ‘Database’ tab is selected, select a database from the list of databases, and press the “Remove” button. This will not permanently remove it from our system, it may be reloaded (follow the procedure for configuring a new database) at any time.

**Edit a Database**
This tool allows us to change the name that PEAKS 4.5 associates with a database, taxonomy files and the header parsing rules for that database. To edit a database, we open the PEAKS Properties dialogue, ensure that the ‘Database’ tab is selected, select a database from the list of databases, and press the “Edit” button.

**Moving/Updating a Database**
If we choose to move a database to another directory, or delete it entirely, we should tell PEAKS. We must remove the database from the list and re-load it. Until we do so, the database name will appear in red in the list of databases and any protein identification using that database will fail.

If we choose to update the database (perhaps by downloading the latest database file and overwriting the old database file), PEAKS will show the database information in light gray. A light grey colour could also mean that the database does not have header information.
Instrument List

PEAKS 4.5 is optimized for many different types of instruments. Weighting the analysis according to the specific fragmentation patterns, resolution, accuracy and charges generated by different types of instruments allows us better precision in analysis. To avoid having to set all these options on every search parameters dialogue, we wrap up commonly re-used parameters that are instrument specific.

For example, when using the data refine tool, and correcting charge, it’s important to notice that an instrument has a MALDI source, as all peptides will have charge 1. Or, when de novo sequencing, it’s essential to note if a spectrum was produced using ETD fragmentation, so that c and z ions are most important.

The instrument list shows the <built-in> and user defined instruments available to all PEAKS tools. From here we can create new instruments or edit existing ones.

Creating/Editing Instrument Settings

To define an instrument, we open the PEAKS Properties dialogue, ensure that the ‘Instrument list’ tab is selected and click the New button, or select one from the list and press the Edit button. To define an instrument ‘on the fly’ while setting up PEAKS auto de novo or PEAKS Protein ID, click the Edit button next to the Instrument dropdown list on the options screen for any PEAKS tool. The Enzyme Definition dialogue will appear.

From here we can define the ion source, fragmentation, and mass analyzers used to generate the data.
Importing and Exporting PEAKS Properties

We may wish to use PEAKS 4.5 on another system. However, if we have a large number of user defined PTM, enzymes and saved parameters it could take a great deal of our time to re-input those. This is where importing and exporting of PEAKS properties is useful.

The export function will save PEAKS Properties information in a XML file. The import function can read a PEAKS properties XML file and overwrite local PEAKS Properties with the information from XML file. If we wish to use our PEAKS properties on a colleague’s system, we must remember to export our colleague’s properties to a separate file so that it will not be lost and can be imported later.

<table>
<thead>
<tr>
<th>A note on sharing sequences with PTM</th>
</tr>
</thead>
</table>

Sequence data and protein identification results for a given spectrum are stored in an .ANZ file. Any modifications that were found in the sequence are also included. As such, user-defined modifications will still show up if the file is viewed on another machine. It is not necessary to import all PEAKS properties to view these modifications. Also, user-defined modifications can be extracted from an .ANZ file and added to the local PEAKS properties.

**To export PEAKS properties to a file**, open the PEAKS Properties dialogue and press the “Export” button. Type in a file name and press the “Save” button.

**To import PEAKS properties from a file**, open the PEAKS Properties dialogue and press the “Import” button. Select a file or type in a file name and press the “Open” button. This must be a PEAKS configuration file in XML format.

**To import a user defined PTM from another user's .ANZ file**, we open the .ANZ file and find a sequence containing the user defined modification. Right click on that sequence to bring up the popup menu. Click the “View Modifications” menu item. This brings up a dialogue box named “Modifications”. Select the PTM of interest from the dropdown list (in this example: ‘Lab 2 custom PTM’) and click the import modification button.
PEAKS Environment Preference Configuration

One of PEAKS 4.5 preferences, PEAKS Environmental Preference allows us to customize PEAKS 4.5 to our needs. PEAKS Environmental Preferences include: Environment, Color, Manual de novo, and Parameters. To edit PEAKS 4.5 Environment Preferences:

- Click the icon in the toolbar;
- Or, from the “Edit” menu, select “Configuration” then “Environment Preference”

The Environment Preferences dialogue will then appear. This dialogue box has seven tabs: ‘Parameters’, ‘Basic Ion Table Editor’, ‘Advanced Ion Table Editor’ ‘Environment’, ‘Color’, ‘Display’, ‘Manual De Novo’ and ‘Parameters’. Clicking a tab will allow us to edit the Environmental Preferences corresponding to that tab.

Removing Saved Parameters

After months of use, our list of parameters saved for use with PEAKS Protein ID or PEAKS auto de novo may become cluttered with infrequently used parameter sets. It makes sense to clear them out from time to time. To do so, open the Environment...
Preferences dialogue and ensure that the ‘Parameters’ tab is selected. A list is shown for each tool that has savable parameter sets. Select one or more (using shift+click and ctrl+click) and click the adjacent “Remove” button to remove it from the list.

**SCV**

**Configuring the ion table**

The ion table, displayed in the top right of the main processing window, displays all the ions that were found as evidence for the selected sequence. There are two presets: the Basic Table and the Advanced Table. Select which one to display by choosing “Show Ion Table ►” from the “View” menu.

The Basic Table will display a maximum of 6 ions. The Advanced Table can be configured to display as many as are available. To configure the ion table, we choose “Edit menu” ➔ “Configuration” ➔ “Environment Preferences” and then choose either the ‘Basic Ion Table Editor’ tab, or the ‘Advanced Ion Table Editor’ tab. Both tabs look and work the same way:

The ions types that will be displayed in the ion table are shown on the right. The complete list of ion types available is shown on the left. To add an ion type to the ion table (i.e. add a column to the ion table):
1. Select one or more ions from the list on the left. Use shift+click and ctrl+click to select multiple list items.

2. Select a charge (from 1-4) from the drop down list in the middle.

3. Click the ‘Add with charge’ button

For example, configure the ion table to display $y^{2+}$ ions by selecting ‘y’ from the list on the left, and ‘2’ from the dropdown charge list.

Remove ion types from the ion table (i.e. remove columns from the table) by selecting one or more items in the list on the right, and clicking the ‘Remove’ button.

**SCV**

Environment - Changing default file handling settings

To change the working environment, we open the Environment Preferences dialogue and ensure that the “Environment” tab is selected.

We can change the environment settings so that when we are browsing our system to find or save data files, PEAKS always starts looking in the folder we specify. The current working folders for data input and data output are shown. PEAKS 4.5 can use the last folder we loaded from (saved to) as the current working folder or by toggling the appropriate radio button to “User directory” we can set it ourselves so that it will be the same each time. The directory where PEAKS stores its preference information cannot be changed.

The option to load a new spectrum view window for each spectrum or open one at a time is available. We can choose to show the sample data at startup. Further, adjustments can be made so that the GNU (General Public) license displays whenever GNU governed software/libraries are called. Click the appropriate checkbox at the bottom of the window.

Once we’ve chosen from these options, pressing the “OK” button will exit, saving changes. The “Cancel” button will exit discarding changes.

**SC**

Environment - PEAKS Client to PEAKS Online

In the Search Engines Details tab of the Environment preferences dialogue, change the “Default PEAKS Invocation” to “PEAKS Online”. Then specify the IP address or URL of the PEAKS Online server, the port number to use, and a username and email address. Press ok. Now we can use PEAKS 4.5 in the same way we normally would, except that processing will occur on the PEAKS Online server. Email is used to notify us that a job has been completed, in the event that we close PEAKS 4.5 before finishing.
SCV **Changing Colours**

For ease of viewing, we can choose which colours we would like to represent which items on the spectrum view. To change the colour of an object on the spectrum, we open the Environment Preferences dialogue and ensure that the ‘Colour’ tab is selected.

Choose the object whose color we’d like to change from the list at the left. Then we select a swatch to choose the colour. After we’ve chosen colors we may click the “OK” button to exit and save changes.

SCV **Display - Changing typefaces and peak width**

We can customize the various typefaces to use on certain parts of the PEAKS 4.5 interface. To do so, open the Display tab on the Environment Preferences window. Select the typeface from the dropdown list and/or change the font size, then press ok. The width of PEAKS on the spectrum display can be adjusted from here as well.
SCV

Changing Manual de novo defaults

We may wish to sequence a peptide manually, using spectrum data. PEAKS 4.5 provides us with a set of tools to help us do so. We may need to tweak these tools to adjust for error tolerance and to customize the working environment. This can be done on an individual basis, by right clicking on the sequence we’re working on, but we can set defaults.

To adjust Manual de novo options, open the Environment Preferences dialogue and ensure that the ‘Manual de novo’ tab is selected.
When sequencing a peptide using the manual de novo tools, we can get PEAKS to help us by searching to the left or right of a selected peak and returning a set of possible sequence tags (see the Manual de novo section in the next chapter). We can choose how many search results we’d like to see and choose how the maximum (number of amino acid residues) length. To choose how long tags will be, we click on the “Maximum tag length” dropdown list box, and making a selection. To choose the number of search results displayed, click on the “Maximum return” dropdown list and make a selection.

Changing the default machine error sets the amount of error PEAKS will tolerate when tagging a residue. For example: we have a mass difference of 113.19 between two y-ions that we have labeled. We are fairly confident that this should be tagged L (Leucine, with actual mass of 113.08), but PEAKS is not labeling it for us. This may be because 113.19 is too far out of PEAKS error tolerance for the mass of L. We can tweak the settings until we get the desired result. To do so, type a value for error (larger numbers indicate greater tolerance) into the “Default machine error” textbox.

After having made all desired changes, click the “OK” button to save changes and exit the dialogue box. Click the “Cancel” button to exit, discarding changes.

Search Engine Details (inChorus)
PEAKS Studio not only performs de novo sequencing and identifies proteins, but compares its results with those of such MS/MS databases as Mascot, OMSSA, SEQUEST, X!Tandem and sequence tag search tool SPIDER with no extra work to you! Simply configure any/all of these search engines, which you may already use for PEAKS.

For each of these options, navigate to the Search Engine Details tab (Edit -> Configuration -> Environment Preferences).

Mascot Server Configuration
In “Default Mascot Information”, type in the location of your Mascot server (We recommend asking your IT person, or the person who set up the server).

Host Name or IP Address – name of the machine on which Mascot search engine is installed;

Port – port number for the Mascot search engine, by default: 80;

Virtual Directory – directory of Mascot MS/MS Ions Search, by default: /mascot.

You can use your regular user account for the username and password. It doesn't necessarily have to be an administrator account. However, some of the built in user accounts like 'daemon' have some special properties that cause problems here. Don’t use the daemon account, but...
you can use an account in the daemon group. The email is not used by PEAKS but is requested by the Mascot server.

Then press OK!

**OMSSA Server Configuration**
OMSSA is configured automatically in PEAKS Studio, therefore, users already have access to this search engine.

**Sequest Server Configuration**
For Sequest, we simply enter in the location of various important parts of BioWorks. Click the Browse button next to each box to find them. They will automatically look for the default location. It's most important to find the sequest.exe, but it doesn't matter quite as much which .params file is chosen as the default as changes will likely be made to it. Finally select your desired output directory for the results.

Then press OK!

**X!Tandem Server Configuration**
In “Default XTandem Information”, type in the location of your XTandem server (We recommend asking your IT person, or the person who set up the server).

Host Name or IP Address – name of the machine on which XTandem search engine is installed;

Port – port number for the XTandem search engine.

Then press OK!
Loading and preparing data

A task based guide to getting our data into PEAKS 4.5.

The following chapters deal with usage. They are broken up into tasks that a typical user might perform. It assumes we can identify parts of the Graphical User Interface and that we are familiar with how PEAKS 4.5 can be configured. The preceding two chapters provide in depth help on these subjects and should be used as a reference. Such detail has occasionally been omitted from this chapter in the interest of succinctness.
SCV

Loading data into PEAKS 4.5

PEAKS 4.5 can be used to process data from any MS/MS instrument, provided the data is accessible, or can be converted to an accessible format. PEAKS handles data files in the following formats:

- PKL: The file format associated with MassLynx software.
- DTA: The file format associated with SEQUEST software
- MGF: The file format associated with Mascot software.
- ANZ – the zip compressed XML based file format associated with PEAKS
- XML format files using the mzXML schema
- XML format files using the mzData schema
- RAW files from Thermo Electron instruments
- WIFF files from ABI QSTAR and QTRAP instruments
- RAW files from Waters QTOF instruments
- .BAF, .YEP and folders of .FID files from Bruker instruments
- .D files from Agilent QTOF instruments
- DAT files created by BSI’s ABI converter software

In order to do any data processing we must first load our spectrum data into PEAKS 4.5. To open a data file, click the icon on the toolbar in the upper left corner of the PEAKS window. Alternatively, to help us find a file more quickly, we can choose from one of the several options in the File Menu.

It is often best to import RAW data directly, so that PEAKS has access to all the rich data, including the MS survey scan and retention time information, and that our analysis does not suffer from poor preprocessing.
Select a file. Click the Open button.

The data file we just opened appears in the Peptide Data Frame on the left. It is represented by its file name. Each spectrum contained in the data file is represented by its precursor ion information (m/z value followed by the charge of the precursor ion that generated the spectrum).

**SCV Loading Thermo RAW data**

PEAKS 4.5 can load RAW data from our Thermo Electron mass spectrometer; provided that the XCalibur software is installed on the same computer as is PEAKS 4.5. To load Thermo RAW data, simply choose File → Import RAW → Thermo (.raw), and browse to the file.

**SCV Loading Agilent .D data**

PEAKS 4.5 can load native data from our Thermo Agilent QTOF, provided that the MassHunter software is installed on the same computer as is PEAKS 4.5. To load
Agilent .d data, simply choose File → Import RAW → Agilent (.d folder), and browse to the file.

SCV Loading Bruker data
PEAKS 4.5 can load data from our Bruker mass spectrometer; provided that the CompassXport software is installed on the same computer as is PEAKS 4.5. To load Bruker data, simply choose File → Import RAW → Bruker (.yep/.baf/.fid files), and browse to the file. If loading .fid files, which are stored in a network of folders, we can select the top level folder to load them all at once.

SCV Loading Shimadzu data
PEAKS 4.5 can load data from our Shimadzu mass spectrometer; provided that the Shimadzu software is installed on the same computer as is PEAKS 4.5. To load Shimadzu data, simply choose File → Import RAW → Shimadzu AXIMA (.run), and browse to the file.

SCV Importing Applied Biosystems WIFF data
PEAKS 4.5 can load WIFF data from our Applied Biosystems QSTAR (or QTRAP) mass spectrometer; provided that the Analyst QS (Analyst 1.4.1 for QTRAP) software and the MSX plug-in are installed on the same computer as is PEAKS 4.5. The MSX tool is produced and sold by Infochromics Ltd., and is available (at cost) from Bioinformatics Solutions Inc. Please contact a BSI sales representative to obtain a license. To load QSTAR WIFF data, simply choose File → Import wiff raw data, and browse to the file.

SCV Importing MassLynx RAW data
PEAKS 4.5 can import RAW data from our Waters/MicroMass QTOF instrument. To do so, we choose “Import RAW data” from the File menu. As above, the file browser appears. Choose the .RAW data, and click the Open button. PEAKS uses a utility called wolf.exe (originally created as part of the Sashimi Project) to access MassLynx libraries and convert the data. PEAKS ships with a version of wolf.exe designed to work with MassLynx 4.1. If we need a different wolf, we can check:

www.bioinfor.com/products/PEAKS/support/watersmicromass.php

Additionally, we must make sure that the following MassLynx libraries are installed on the same computer as PEAKS and wolf.exe:

- DACServer.dll
- Genutil.dll
- MetaGD32.dll
- raw.dll
- securityAccess.dll
● securitySettings.dll
● securitySignature.dll

They should be stored in the folder C:\MassLynx\ as part of the MassLynx software. If they are not stored here, or MassLynx is installed on another computer, the automatic loading will not work. If the automatic loading is not working for either reason try this:

1. We should be able to find the listed files on our computer, or another computer in our lab. If you can copy them, do so.

2. We can then create a folder called “C:\MassLynx\” on our computer, and place the files we copied here. But we’re not finished, we must also register these files with Windows.

3. We can find a file on our system called regsvr32.exe using the ‘Find’ or ‘Search’ tool in our windows start menu. It is probably in “C:\WINDOWS\System32\”. If it’s not there, substitute the correct location in step 4.

4. Open a command prompt or the Run tool from the start menu and type the following:

   C:\WINDOWS\System32\regsvr32 C:\MassLynx\DACServer.dll

   All on one line, with one space in the middle as shown. Press the enter key. If successful, windows will pop up a success message.

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### Please check the license

The libraries mentioned in this section are part of the MassLynx software, distributed by Waters Corp. Please check the MassLynx license agreement, or contact a Waters representative to make sure it is okay to copy and use the libraries in this way.

---

### SCV

**Importing Data from the ABI 4700 or ABI 4800**

BSI has created a converter to extract the data from an ABI-Oracle database. If we require this separate, free tool, we must ask a BSI representative. Once installed, we can start up the ABI 4700 Data Extractor from the Start menu.

**System Requirements**

This extractor can be installed on the same machine as ABI 4700 Explorer and the Oracle database (we will call this machine the 4700 SERVER in the following instructions) or another machine that has direct network access (no firewall, no proxy
required) to the 4700 SERVER. Windows 2000 or Windows XP is recommended for use with this tool.

**Configuration:**
Before using the ABI 4700 Data Extractor, we should configure it. To do so, we can choose “Settings” from the “File” menu. Configuration needs the following:

- **4700 SERVER Name or IP Address:** input “localhost” if the Extractor is running on the 4700 SERVER (this is the default value), otherwise enter the IP address of the 4700 SERVER.

- **The socket used by the 4700 SERVER:** this is the port that the Oracle database listens to (the default is 1521).

- **Username to access the Oracle database:** most likely we do not need to change this (the default is “tsquared”).

- **Password to access the Oracle database:** mostly likely we do not need to change this one either.

**Data extraction procedure:**
1. Load Spot Set List from the database: (Do it via menu File | Load Spot Set List) The extractor will export the peak list of a spot set into a PKL file.

2. Open a Spot Set: (menu File | Open Spot Set) Spot Set Chooser will help the user to choose a spot set. After selecting a spot set, click 'OK' to open it. The job run information of a spot set will be shown.

3. Select a job run: There is a radio button before each job run, only the MS/MS job run can be selected for export, because we need the precursor information. Select a job run and click 'Convert' to do the extraction.

4. Choose a filename to save: After clicking the 'Convert' button, the user needs to input a file name and the peak lists of the selected job run will be exported.

**SCV Tracking data properties**
PEAKS reads and tracks information about the experiment and data for use in the analysis and for our future reference. This information is displayed automatically after a file loads. If any information cannot be found in the file, PEAKS will prompt us with a warning; since some information is used to suggest parameters in PEAKS analysis, it
makes sense to complete the information before proceeding. Additionally, we can enter notes about the data or experiment that will be saved along with the results.

**SC Refining data before analysis**

Since mass spectrometry data often contains noise and redundant data, it makes sense to purify the data before analysis. This will increase the quality of the results, while saving time spent on database searching and/or de novo sequencing. MS/MS spectra that are purely noise can be removed from the data:

- peptide charge information can be verified and recovered
- multiple low quality scans of the same peptide can be merged into one scan with intense signal PEAKS
- the MS/MS scans themselves can be centroided, filtered for noise, and deconvoluted.

When PEAKS is connected to a PEAKS Online server, we’ll also save time by uploading smaller, preprocessed data. Data refinement is done locally, before uploading to the server.

To begin the refinement of data from a whole MS/MS run:

1. In the Peptide Data Frame, select the data file(s) containing the data that we wish to refine.
2. Click the Data Refine toolbar icon

Or

Select “Data Refine” from the “Tools” menu.

Or

Right click on the selected data file and select “Data Refine” from the popup menu

The Data refinement options dialogue appears:

3. Choose the data refinement tools we wish to use by clicking the “yes” radio button next to each one.

- **Instrument Type:** choose the type of spectrometer that produced the data to be analyzed. This setting is important as MS resolution and ion source are important for peptide charge calculation.

- **Merge Scans of the same peptide:** in DDA mode, a mass spectrometer will often produce several tandem ms (i.e. ms/ms) scans of the same peptide. To increase the intensity of real signal PEAKS within these scans and to reduce
the size of the whole data set, it makes sense to merge ms/ms scans of the same peptide together. To avoid improper merging (of ms/ms scans of different peptides) we make sure that the measured parent ion masses of these peptides are very close and that they have similar retention times in the LC column. The units here are m/z values in Daltons. For retention time, we use whatever units are recorded in the data file (usually minutes or seconds).

- **Correct precursor charges:** Since a mass spectrometer measures mass-to-charge ratios, we must know the charge on a peptide before we can determine its mass. The standard method of finding the charge is to look at the spacing of the isotope ladder in the survey scan. However many Ion-Trap instruments don't have enough resolution for this. So PEAKS will look at the MS/MS data to determine if it's charge 1+, 2+ or 3+. For data where the survey scan is available, PEAKS will examine the precursor ion’s isotope distribution to confirm or correct the charge assignment.

- **Remove low quality MS/MS scans:** Scans of contaminants and electrical noise should not be included in analysis. Removing them from the data set will save time, and reduce the risk of random matches to the database. PEAKS presents an effective tool for removing these low quality ms/ms scans. Type in the boxes to set ranges of retention time and m/z – PEAKS between these values will be considered for analysis. Additionally, PEAKS examines the MS/MS spectrum to determine its quality (using various characteristics including total intensity, n-term + c-term pairing, sequence tag calling, etc.), so you can choose a threshold of quality score (a value from 0 to 1) for accepting a scan. Set to 0.01 to disable this filtering.

- **Preprocess MS/MS scans:** Deconvolution (de-isotoping), centroiding and noise filtering within the MS/MS data. Data is always saved in the ANZ file along with the PEAKS results. Preprocessing can save hard disk space or upload time. But make sure to have the original data available in case we need to refer to it later.

To see how your data is changed, refer to the data properties window (above section).

A note on preserving data-results integrity

| Protein ID and de novo sequencing results obtained for a given dataset prior to use of this tool may become invalid, since some spectra are removed/merged/corrected and the data-results relationship may be broken. PEAKS 4.5 will warn us when this may occur and prompt us to save a separate copy. |

Parameters used to refine the data before analysis are saved in the .ANZ file. These and the reduction in the number of MS/MS scans are summarized on the data properties window.
Manually Manipulating Data Files

Editing Precursor information

It is possible that the precursor information, as listed in the Peptide Data Frame, is incorrect. If the charge listed is wrong or if the m/z listed is even slightly incorrect (more than the selected error tolerance) it could really affect the quality of the results. In this case it is imperative that we change the precursor information. The change will only affect the .ANZ file we are working on.

To edit precursor information, select a spectrum by clicking on its name, then right click the mouse while holding it in position over the name. A small menu will appear; click on “Edit Precursor”.

In the dialogue that follows, type the new precursor information into the appropriate textboxes. Click the “Apply” button when finished to apply the changes. Click the “Cancel” button to exit, discarding changes.

The precursor information will be updated, reflected by a change in the name of the spectrum in the Peptide Data Frame. A * will also appear in front of that name, indicating that there is unsaved information pertaining to that spectrum.

Manually merging MS/MS scans of the same peptide

If we’ve done several MS/MS scans of the same peptide, we may want to reduce the amount of data to process and at the same time, improve the data quality by merging all of a peptide’s MS/MS scans together. Often, we choose to automatically merge appropriate spectra from the whole data file using the Data Refine tool (see above). But this can also be done manually.

To manually merge spectra after opening a data file:

1. Select those spectra we wish to merge together from the Peptide Data Tree (left) using shift+click and ctrl+click.
2. Next, right click in the Peptide Data panel, and choose Merge Spectra from the popup window that appears, OR click the ‘Manual Merge Spectra’ toolbar button.

3. A dialogue will appear, asking what should be the correct value for the precursor mass and charge. After reviewing and/or correcting the value, press “OK”. The spectra will be merged.

SCV

Cutting and Copying Spectrum Data
If we wish to move spectrum data from one data file to another we may do so by copying and pasting it (see below for pasting instructions). Also, we may wish to make a copy of the spectrum in the same data file in order to re-sequence an individual spectrum using different preferences. Cutting spectrum data will remove it completely until pasted. Copying spectrum data will duplicate the spectra when pasted. To cut/copy spectrum data:

Changes made to the original spectrum after duplication will not affect the duplicated spectrum.

Select a spectrum by clicking on its name (select multiple spectra by holding down the ‘control’ key and clicking on any number of spectrum) in the Peptide Data Frame.

Right click on one of the selected spectra. A small pop-menu will appear. Select “Cut” or “Copy”

Or click the “Copy” button or “Cut” button in the main toolbar.

Copied/Cut items will remain on the clipboard until replaced by another copied/cut item. Warning: Unless pasted, a cut item will be lost, as subsequent cut/copied items will displace it from the clipboard.

SCV

Pasting Spectrum Data
After having copied or cut spectrum data, we would like to paste it into another data file, or the same data file. To paste spectrum data:

1. Select the data file into which we wish to paste the spectrum (or spectra) by clicking on its name in the Peptide Data Frame. We may only choose to paste into one data file at a time.

2. Right click on one of the selected spectra. A small popup menu will appear. Select “Paste” from the popup menu.

Or click the “Paste” button in the main toolbar.

The pasted spectra will appear in the Peptide Data, under the data file into which we pasted.
Chapter 7

Analyzing data with PEAKS 4.5

A task based guide to processing our data with PEAKS 4.5.

This chapter will step us through the analytical techniques available to PEAKS 4.5 users. It is broken up into tasks that a typical user might perform. It assumes we can identify parts of the Graphical User Interface and that we are familiar with how PEAKS 4.5 can be configured. Such detail has occasionally been omitted from this chapter in the interest of succinctness.
Using PEAKS with modifications (PTM)

PEAKS 4.5 provides the most flexible handling of post-translational modifications of any software built for de novo sequencing and protein ID. Users are free to create their own modifications (see the Creating a New PTM section in Chapter 5), and search for any combination and any number of modifications. Modifications can be considered as part of auto de novo sequencing or protein identification. The search is set up the same way for both tools. The options screen for each tool has an area titled “PTM selected for search”. Any modifications to be considered during the search will be shown here (and labeled as ‘Fixed’ or ‘Var’). When we first run PEAKS, the box will be blank, meaning no PTM are selected.

To add modifications to this list, click the Add/Remove PTM button. The Modification dialogue appears. The entire PTM library — i.e. the list of all <built in> and user defined modifications that are available to PEAKS — is displayed in the list on the left. We'll choose the modifications we need from this list. To add a new modification to the PTM library, click the ‘New PTM’ button. To edit a modification in the PTM library, select it from the list on the left and press the ‘Edit PTM’ button. To remove a modification from the PTM library, select it from the list on the left and press the ‘Remove PTM’ library.

If you remove/edit a PTM, it will be removed or changed from/in any saved parameter set that refers to it.

The lists on the right show what PTM will be enabled for the search. Use the ‘Select as Fixed=>’, ‘Select as Variable=>’ and ‘<=Unselect’ buttons to move them in and out of these lists. Press the ‘OK’ button when finished and the changes we made will be reflected on the protein ID options dialogue.
Remember, when doing auto de novo sequencing or PEAKS Protein ID on a complex mixture, we will get best results if we choose the correct fixed PTM and a few variable PTM. When using PEAKS Protein ID to characterize a protein, it is best to search against a small database that contains only a few proteins and turn on all modifications. Furthermore, to limit spurious hits, we can assume that it is less likely that a tryptic length peptide will not be modified more than a few times and as such, limit the number of variable modifications that can occur on each peptide.

**Auto de novo Sequencing**

To begin auto de novo sequence derivation:

1. In the Peptide Data Frame, select the data file(s) containing the spectra that we wish to sequence by Auto de novo. We can also select an individual spectrum, or a few spectra within a data file; auto de novo will proceed on only the spectra selected.

2. Click the Automatic de novo toolbar icon

   Or

   Select “Auto de novo” from the “Tools” menu.

   Or
Right click on the selected spectra or data files and select “Auto de novo” from the popup menu.

The Auto de novo Parameters dialogue window will appear:

Selecting an instrument fills in suggested error tolerances. Try some slightly higher or lower ones to find the best result.

3. If we wish to change any of these parameters, we do so now.

- **Instrument Type:** Choose the type of spectrometer that produced the data to be analyzed. To define a new instrument, press the Edit Button.

- **Parent mass error tolerance:** Determine how much random and systematic experimental error on the parent/precursor ion PEAKS will allow for in its analysis. Type a tolerance in the textbox and choose units from the dropdown list. Using PPM allows for larger errors at larger m/z values. PEAKS will be very stringent concerning this value, so new PEAKS users should try setting this a little higher than past experience may suggest, if sensitivity is a concern.

- **Fragment mass error tolerance:** Determine how much random and systematic experimental error on the fragment/daughter ion PEAKS will allow for in its analysis. Type a tolerance in the textbox. Again, new PEAKS users should try setting this a little higher than past experience may suggest.
- **Enzyme:** Tell PEAKS what kind of enzyme was used to digest the sample. Choose from a dropdown list of enzymes, or if our enzyme (or combination of enzymes) is not in the list, click the "Edit Enzymes" button.

- **Report top:** Set how many peptide sequences PEAKS will report from its de novo sequencing analysis.

- **PTM selected for search:** This list tells PEAKS what kind of post-translational modifications to include in its analysis. Each is marked Fixed or Variable. To edit this list, click the "Add/Remove PTM" button.

- **Max variable PTM per peptide:** To reduce uncertainty, we can limit PEAKS' de novo sequencing 'vocabulary', by restricting the number of variable PTM we can find on a peptide. Specify a number by typing it into the box. To lift such restrictions, type a very large number (longer than the length of the peptide).

- **Saving/Loading Parameters:** After setting up parameters, we can save them for future use. Click the "Save Parameters" button, and choose a name for future reference when prompted. Don't worry, we can't accidentally overwrite the defaults. Any parameters we save will be available in the drop-down list at the top of the window. To see what's inside, select one and the parameters boxes will be populated.

- **Preprocess before auto de novo:** PEAKS has its own built-in preprocessor for removing noise, centroiding and peak charge recognition from MS/MS data. Check this box to turn preprocessing on.

### Notes on pre-processing

BSI highly recommends using PEAKS to preprocess all data, as opposed to using instrument vendor software, if the data is to be used by PEAKS. PEAKS preprocessor should not be used on data that has already been pre-processed as this will have adverse effects on the results (unless it is ion-trap data).

4. Press the “OK” button to commence Auto de novo sequencing.

Once a job is submitted to PEAKS 4.5, it is added to the Task Queue for processing. After processing, the job is removed from the task queue list and the dark green icons beside the spectra (in the Peptide Data Frame) change to light green and/or an asterisk (*) appears.
Viewing Auto de novo Results

After performing auto de novo on a spectrum, we may wish to see what the algorithm determined the peptide sequence to be and review the results for ourselves. To do so, we click on the spectrum of interest in the Peptide Data Frame. This brings up the Main Processing Window for that spectrum.

The most likely peptide sequence candidate, as determined by PEAKS auto de novo, will be automatically selected. This is found in the Peptide Candidates Frame as the top listed candidate under ‘PEAKS Auto de novo’. In this example, the highlighted sequence is VDVEK. Any modifications that have been found will be shown abbreviated and in sequence before the amino acid residue they are associated with. If the PTM was defined/created by another PEAKS user, the PTM would still be shown and it can be imported into the local PEAKS configuration as desired.

Right next to the proposed sequence, the auto de novo confidence score is shown. Positional confidences (that is, confidence that the correct residue in each position has been identified) are readily available by color coding. 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 a low confidence (less than 60%). For more detailed positional confidence, we can place our mouse over the sequence of interest. A Position Confidence Table will appear, showing the confidence that each tag/subsequence is correct.

In the Ion Table frame, select a cell from the Ion Table – each cell represents an ion. This will highlight its position on an error plot (scroll the Ion Table frame down if the error plot is not visible). A point close to the centerline indicates a more confident
result. We can also notice that the peak corresponding to the Ion we selected is highlighted on the Spectrum View. Select a whole column to highlight all the points for that type of ion. The types of ions displayed in the ion table can be configured, choose Configuration → Edit Ion Table from the Edit menu. Users might find this particularly useful when sequencing data acquired using ECD or ETD.

By looking at the Spectrum View Frame, we can see the strength of the MS/MS PEAKS that PEAKS 4.5 has set as ions. The view also displays the mass of the ions at that peak and the type of ion. Click on a peak to mark it and display its information at the top left corner of the Spectrum View Frame.

Zoom in by clicking and dragging horizontally on an area of the Spectrum View. The area over which we dragged will now take up the whole spectrum view. To un-zoom, press the undo zoom icon or press the 1:1 icon to return to the full spectrum view.

We may also zoom in on the spectrum using the Spectrum Alignment Frame. Again, click and drag horizontally on an area of the Spectrum View. The area over which we dragged will now take up the whole spectrum view. The blue bar beneath the Spectrum Alignment view shows where we are zoomed in. The white portion of the bar represents the area that we are zoomed in on.

We can toggle whether or not we’d like to see the positions of the y-ions and b-ions (and the proposed residues in sequence between them) on the alignment view by pressing the y-ion alignment (yellow and red) and b-ion alignment (yellow and purple) icons in the main processing window toolbar.

To view another peptide candidate, as determined by auto de novo, click on another peptide in the Peptide Candidates Frame and under ‘PEAKS Auto de novo’. The information in the Ion Table will change, as will the tags on the spectrum, to reflect the selected peptide candidate’s sequence.

**SCV Editing sequencing results (preparation)**

We cannot change the results provided by PEAKS auto do novo or PEAKS database search. However, we can make a copy of any sequence and edit it using manual de novo techniques. To copy a sequence for editing:

1. Select a peptide sequence candidate from within the Peptide Candidates Frame. We can only select one peptide sequence candidate at a time.

2. Right click the mouse button while holding the mouse over that sequence. A pop-up menu will appear.
3. We can select the pop-up menu item “Copy for manual de novo”. In this case, the sequence will be automatically placed under the ‘Manual de novo’ heading. A ‘Manual de novo’ heading will be created if there wasn’t one there already.

4. Now we select our newly copied sequence, under the ‘Manual de novo’ heading, to display this sequence in the Ion Table Frame, Spectrum View Frame, and Spectrum Alignment Frame.

Now we are ready to edit the sequence, using manual de novo techniques.
**Manual De Novo Sequencing**

We can use manual *de novo* sequencing to fine tune the results of an auto *de novo* analysis or to perform our own sequencing analysis from scratch. PEAKS 4.5 provides a set of tools to help us sequence a peptide, using graphic cues from the spectrum.

**Creating a fresh spectrum for sequencing**

We cannot change the results provided by PEAKS auto *de novo* or PEAKS database search. Thus, to begin manual *de novo* sequencing, we must either copy a sequenced peptide (see above section: Preparing to edit sequence results) or create a new peptide candidate for sequencing.

To create a new peptide candidate for sequencing:

1. Right click on the ‘Peptide Candidates’ heading, the ‘Manual *de novo*’, or any ‘user defined type’ heading. This will bring up a pop-up menu.

2. Select “New candidate for manual *de novo*” from the pop-up menu

A new candidate will be created under the ‘Manual *de novo*’ heading, or under the ‘user defined type’ heading if we selected a user defined type. The new candidate will not have been sequenced, so it will be represented by the mass of the peptide less the mass of water (e.g. [945.15]).

**Manual *de novo* Operations**

*All operations occur in the Spectrum View Frame of the Main Processing Window.*

When the mouse is placed in the Spectrum View Frame, a blue (by default) bar follows the movement of the mouse. This is the *Position Bar* and it is used as a cursor for all manual *de novo* operations. The cursor’s position on the m/z scale is enumerated on the top of the *Position Bar*.

**Selecting a peak**

To select a peak, click on it. An orange (by default) bar, called *Freeze Bar*, indicates the selected peak.

Alternatively an ion peak can be selected by clicking on its corresponding cell in the Ion Table.

**Measuring distance along the m/z scale**

Once a peak is selected with the *Freeze Bar*, moving the mouse left or right will display the *Position Bar* along with a value that represents the m/z difference (as an absolute value) between the selected peak (orange) and the *Position Bar* (blue). In the example below, the distance between the selected peak and the position bar is 51.02 Daltons.

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S marks a section applicable to PEAKS Studio, C for PEAKS client and V for PEAKS Viewer
Measure the m/z difference between two PEAKS
Select a peak (orange line by default) with the Freeze Bar and move the mouse to the left or right. Hold the Position Bar above another peak. The number above the Position Bar is the difference between the two PEAKS.

Deselect a peak
Double click anywhere in the Spectrum View Frame

Zoom in on part of the spectrum
In the Spectrum View Frame or the Spectrum Alignment Frame, click and drag the mouse horizontally. The selected area will be shown in the Spectrum View Frame.

Add (remove) ions to (from) a peak
Select a peak, then right click the mouse anywhere in the Spectrum View Frame. Select “Set Y Ion” from the pop-up menu to designate the peak as a y ion, “Set B Ion” from the pop-up menu to designate the peak as a b ion, Select “Ion Edit” from the pop-up menu to view the Ion Editor dialog box and designate the peak as another ion.
The Ion Editor dialogue allows us to add or remove ion designations to/from a peak. Select an ion from the ion choice list and press the “Add” button to add it to the selected ion list. Remove an ion from the selected ion list by selecting it and pressing the “Remove” button. We can type any comments we wish to make about the ion/peak, then press the “Apply” button to apply the changes to the selected peak.

Two short-cut keys may also be used to label a peak. Select a peak, then hit the 'y' key to add a y-ion and or the 'b' to add a b-ion to the peak on your keyboard.

After setting an ion, both the alignment view and the peptide sequence candidate name (as displayed in the peptide candidate frame) will change to reflect the mass remaining to be sequenced on either side of the ion. After setting two ions, PEAKS 4.5 will estimate the residue found between them (if a residue corresponds closely to the mass difference). The peptide sequence candidate name (as displayed in the peptide candidate frame) will change to show the residue and the mass remaining to be sequenced on either side of the residue.

Using sequence tags

Searching the left or right side of the spectrum for the first/last y or b ion: right click anywhere in the Spectrum View Frame to trigger the popup menu. From the menu, select the search of interest. PEAKS will select the appropriate terminal tags and show them in the Ion Table Frame. We may test the suitability of a tag by clicking on its radio button; the tag will be shown, in position, on the Spectrum View. We may insert one or more tags by clicking on their checkboxes, then clicking the “Apply” button. Press the “Cancel” button at any time to exit the search discarding changes.

Search a sequence tag: select a peak with a defined ion (i.e. an ion that has been labeled with a peptide). Right click to trigger the popup menu, then select “Search Right” or “Search Left” to search peptide tags either to the right or left of the selected peak. PEAKS will select the appropriate tags and show them in the Ion Table Frame. We may test the suitability of a tag by clicking on its radio button; the tag will be shown in position on the Spectrum View. We may insert one or more tags by clicking on their checkboxes, then clicking the “Apply” button. Press the “Cancel” button at any time to exit the search discarding changes.

Undoing an edit

If we have made an error in our sequencing it is possible to undo the change. With the Peptide candidate still selected in the Peptide Candidates Frame, click the previous peptide button to return to the previous peptide sequence. We can click this button multiple times to return to successively earlier stages in our edit.

Redoing an edit

If we have undone one too many changes, we can redo that change by clicking the next peptide button. We can click this button multiple times to proceed to successively later stages in our edit.
Suggesting a sequence to see how it fits the data

If the data is ambiguous, PEAKS 4.5 may not have displayed a particular candidate that we wish to evaluate, after auto de novo, or protein ID. We may enter this sequence and have PEAKS 4.5 find if there is any evidence for it in the data.

For instance, PEAKS may give the sequence RMYNVHGC(phosphorylationS)K for a particular spectrum, and we may wish to see if there’s any evidence for the phosphorylation being on the Tyrosine. As such, we may type in our own version of the sequence and have PEAKS find ions that might support our hypothesis.

To do so, open the spectra in the main processing window, and right click on “Peptide Candidates” (in the Peptide Candidates Frame). Then, from the pop-up menu that appears, choose “New Candidate for Manual De Novo”.

A new node will appear with the heading “Manual De Novo” and beneath it will be the mass of the residues yet to be sequenced (in square brackets). Right-click on this heading. In the pop-up menu that appears, choose “New Candidate with user input sequence”, and the Sequence Input dialogue box will appear.
We can now enter our proposed sequence. The total mass of the residues, modifications, and un-sequenced masses should equal the total mass of the peptide (minus water). We might find the mass calculator tool (Tools menu), useful in this regard.

Short forms for the modifications may also be used.

Enter sequences in the format:

- MPELAYLK
- [228.09]ELAYLK
- DE[226.168]AYLK
- EDLLA(phosphorylationY)LK
- DE[226.168]A(phosphorylationY)LK

Then press the OK button. The sequence we just entered will appear under the “Manual De Novo” heading and when selected, the ions that PEAKS has found to match the proposed sequence will appear on the spectrum, spectrum alignment view, and ion table.
Protein Identification

The PEAKS protein identification method is unique – an improvement on and the ideal compliment to existing tools. The unique approach is a combination of sequence tag searching and fragment ion mass matching.

PEAKS also introduces an amalgamative approach to protein identification called inChorus. With inChorus protein identification technology, we can use PEAKS together with several other protein identification software programs. This will deliver more protein coverage and more confidence in results than any one method on its own.

The following two sections deal with usage of PEAKS protein identification on its own and usage of inChorus protein identification.

PEAKS Protein Identification

The PEAKS Protein ID search engine takes a hybrid approach that uses sequence tag information to filter the protein or EST database before fragment ion fingerprinting. Thus to get useful protein identification results, we must first perform de novo sequencing on the spectrum data.

If we already have sequence information for this data, we may use this existing sequence information (manual or auto de novo sequences) to filter the database. If we do not have existing sequence information, or if we wish to refine our database search by providing brand new sequence information, we can ask PEAKS to perform auto de novo before searching the database. Brand new results will not overwrite any existing sequence data that we have.

1. In the Peptide Data Frame, we select the data file(s) that we wish PEAKS to use to identify our protein(s). This can be done by clicking on a data file’s name in, the peptide data frame.

2. Click the "Protein identification" toolbar icon.

Or

Select “PEAKS Protein ID” from the tools menu.

The Protein Identification Parameters dialogue window will appear

3. If we wish to change any of the protein identification search parameters, we do so now.
- **Instrument Type**: choose the type of spectrometer that produced the data to be analyzed. To define a new instrument, press the Edit Button.

- **Parent mass error tolerance**: Determine how much random and systematic experimental error on the parent/precursor ion PEAKS will allow for in its analysis. Type a tolerance in the textbox and choose units from the dropdown list. Using PPM allows for larger errors at larger m/z values. PEAKS will be very stringent concerning this value, so new PEAKS users should try setting this a little higher than past experience may suggest, if sensitivity is a concern.

- **Fragment mass error tolerance**: Determine how much random and systematic experimental error on the fragment/daughter ion PEAKS will allow for in its analysis. Type a tolerance in the textbox. Again, new PEAKS users should try setting this a little higher than past experience may suggest.

- **Enzyme**: Tell PEAKS what kind of enzyme was used to digest the sample. Choose from a dropdown list of enzymes, or if our enzyme (or combination of enzymes) is not in the list, click the "Edit Enzymes" button.

- **Max missed cleavages**: determine the most missed cleavages to allow, internal to the peptide, in a de novo sequence. For instance, if we set this to 2, and Trypsin is the enzyme, then PEAKS will return de novo sequences with up to 2 R's or K's internally.

- **PTM selected for search**: this list tells PEAKS what kind of post-translational modifications to include in it's analysis. Each is marked Fixed or Variable. To edit this list, click the "Add/Remove PTM" button.

- **Max variable PTM per peptide**: To reduce uncertainty, we can limit PEAKS' de novo sequencing 'vocabulary', by restricting the number of variable PTM we can find on a peptide. Specify a number by typing it into the box. To lift such restrictions, type a very large number (longer than the length of the peptide).

### Best practices for setting modifications (PTM)

The developers have discovered that database searching often returns better results if the auto de novo analysis is run with no variable PTM (perhaps one or two if necessary), but with the correct enzyme and fixed PTM. Modifications should be then turned on for the database search function. m/z tolerance can also be adjusted separately for each phase to allow us to tweak the results.
- **Database to search**: Select from this dropdown list, one of the FASTA databases that we’ve set up in PEAKS. If the database we'd like to search is not in this list, click the "Load new database" button.

- **Taxonomy selection**: This list displays the taxa we've chosen for our search. If the database selected has taxon information available, we can click the aptly labeled "Add/Remove Taxa" button. Otherwise, the whole database will be searched. The selections correspond to established hierarchy -- i.e. selecting 'Mamalia' will search all of 'horse, cow, rat, mouse, human, etc.'

- **Paste FASTA sequences**: If we already know the sequence of the protein(s) we are studying, we can paste it here (must be FASTA format) and run a search against it. Alternatively, if we search the same sequence regularly, it is recommended to simply create a small text file and configure it as a database for PEAKS.

- **Preprocess before auto de novo**: PEAKS has its own built-in preprocessor for removing noise, centroiding and peak charge recognition from MS/MS data. Check this box to turn preprocessing on.

**Notes on pre-processing**

BSI highly recommends using PEAKS to preprocess all data, as opposed to using instrument vendor software, if the data is to be used by PEAKS. The PEAKS preprocessor should not be used on data that has already been deconvoluted by PEAKS, or if a peak list was loaded into PEAKS software as this will have adverse effects on the results (unless it is ion-trap data).

If we choose to do pre-processing ‘on the fly’ during protein ID (as opposed to using the Data Refine tool beforehand) PEAKS preserves the original data, and does not save the results of its preprocessing.

- **Advanced Options - de novo**: We must have some de novo sequences before database searching since PEAKS sequence tags to help in database searching. As such the option of doing de novo prior to protein ID is presented here. In most cases, the same values for instrument, error, enzyme and PTM can be used in de novo and in protein ID, but we have the option of using one of our saved de novo parameter sets for the de novo portion. Select one from the drop down list.

- **Saving/Loading Parameters**: After setting up parameters, we can save them for future use. Click the "Save Parameters" button, and choose a name for future reference when prompted. Don't worry, we can't accidently overwrite
the defaults. Any parameters we save will be available in the drop-down list at the top of the window. To see what's inside, just select one, and the parameters boxes will be populated. Note: the Advanced Options selections will not be saved.

4. Press the “OK” button to commence Auto de novo (if we have so chosen) and subsequent protein identification.

If we have chosen to perform auto de novo prior to our database search, the auto de novo process will appear first in the task queue. Once this is finished the database search will begin. If PEAKS finds protein candidates after searching the database, a Protein Identification Results window will appear:

**inChorus Protein Identification**

inChorus Protein Identification will call upon several search engines for protein identification. Once we load our data into PEAKS, we can invoke (start searches running on) several search engines. When all the results are returned, PEAKS 4.5 will compare the answers and summarize everything in one simple report.

1. In the Peptide Data Frame, we select the data file(s) that we wish PEAKS to use to identify our protein(s). This can be done by clicking on a data file's name in the peptide data frame.

2. Click the "Protein Identification" toolbar icon.

   Or

   Select “inChorus Protein ID” from the tools menu

The inChorus database search launch window will appear:
3. First select each of the protein identification tools we would like to use by putting a checkmark in their respective checkboxes. Search parameters for each program can be set by clicking the corresponding Options icon.

4. To tell inChorus to launch a search engine, click the Options Button for that search engine. To import existing results, click the Import Button for that search engine.

5. Select a database to consolidate all the results, in the Database Options frame at the bottom. Since we'll have more peptide information than any one search engine could have had, we'll get a more complete protein identification by going back to the database to rebuild the peptide hits. Old search results can also be improved with new databases in this way. PEAKS will suggest a database and taxon to use (based on our selection for one of the search engines). If we wish to use a different database, select one of the databases configured for use with PEAKS from the dropdown.

6. Press the Launch inChorus Button

Options for search engines inChorus
Option screens for each of the programs available to inChorus are designed to work in the same way as options screens for the original programs. For help in setting search parameters for each program, please refer to that program’s user manual. For help with PEAKS Protein ID, please refer to the above section.
For the Sequest search engine it is preferable to index the database prior to searching. The parameters (e.g. peptide mass, enzyme and static PTM settings) for database indexing must be the same as those used for searching. As such PEAKS will index the selected database, then search said database with the parameters as selected on the Sequest Options Screen.

**Importing results for inChorus**

PEAKS inChorus reads X!Tandem and OMSSA results in their respective XML formats, Mascot results in .dat format and Sequest results in .srf format. inChorus may not be able to import older results files, if the format varies significantly.

It is up to the user to make sure the imported results were generated from the correct data. PEAKS inChorus will attempt to match the results back to the dataset we have open – but the data results relationship may not be preserved in all cases. PEAKS and SPIDER results are selected from the file we currently have open, so this is not an issue.

**SCV Selecting unmatched de novo results**

When working with unknown organisms we are interested in finding novel peptides, but abundant, uninteresting proteins (like keratin) can get in the way. Even when working with human samples, we may have some interesting mutated peptides that couldn’t be matched to the database. Having identified proteins using PEAKS Protein ID or inChorus, we can focus on the new, interesting peptides.

Open a Protein ID report and click on the Filter tab to make sure that Option 1 (de novo view shows peptides that could not be explained …) is selected. Then, if desired, add some filtering criteria, then press the Apply Filters button.

Click on the De Novo View tab. This now displays de novo sequences for spectra that could not be explained by the identified proteins (as listed in Protein View). Select some sequences using click and drag, shift+click or ctrl+click…

---

**Table:**

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>HLYFTTPK</td>
<td>99.98</td>
</tr>
<tr>
<td>DPAVER</td>
<td>99.92</td>
</tr>
<tr>
<td>MCFDTrace</td>
<td>99.9</td>
</tr>
<tr>
<td>TESCLMLBNSK</td>
<td>99.88</td>
</tr>
<tr>
<td>TESCLMLBNSK</td>
<td>99.85</td>
</tr>
<tr>
<td>TESCLMLBNSK</td>
<td>99.41</td>
</tr>
<tr>
<td>DAI REIK</td>
<td>99.41</td>
</tr>
<tr>
<td>BCRF</td>
<td>99.43</td>
</tr>
<tr>
<td>MALHCMQVAVGSLLDSQSPPLR</td>
<td>9.85</td>
</tr>
<tr>
<td>LDGQGAK</td>
<td>9.26</td>
</tr>
<tr>
<td>LDGQGAK</td>
<td>9.26</td>
</tr>
<tr>
<td>LDGQGAK</td>
<td>79.79</td>
</tr>
<tr>
<td>KTAAVFII</td>
<td>7.82</td>
</tr>
</tbody>
</table>
... they appear highlighted in the peptide data tree on the left. A good next step is to run SPIDER on these data to see if we can find homologous peptides.

SC

Running protein identification on select spectra

When searching, PEAKS may not have found a hit for certain spectra. If these are good data, we may wish to try searching them against a more general database. Before we do so, we must create a new data set with these “good spectra that did not match”. This is essential so that we can organize our data well, and because PEAKS will only run Protein ID on all the spectra in a data node. To create a new data node:

1. Make a new data node by right clicking on the peptide data node. The new node appears as “Data1”.
2. Select the relevant spectra using shift+click and ctrl+click. Then press the cut button.
3. Click the new node and press the paste button. Pressing the + next to Data1 will expand it and reveal the pasted spectra.

After pasting, we must save Data1 to continue

Now we’ve essentially removed the already matched peptides from our dataset. We can now run protein identification on Data1 or on the remaining spectra in our original dataset. We can save that dataset in a new file or any of the other functions that apply to regular nodes. Make sure the new node is saved before running protein ID (or any other function) on it.

SC

SPIDER search for sequence homology

After having obtained de novo sequences for peptides that are not in the database, it’s a good idea to look for a homologous peptide in the database. This will help us learn more about the proteins in our sample. To search with SPIDER we must first have some good de novo sequences.
1. Select some spectra (either directly from the Peptide Data Tree (left) or from a Protein ID report). SPIDER will use the best de novo sequence candidate from each selected spectrum.

2. Click the SPIDER icon on the toolbar – OR – choose SPIDER Search from the Tools menu. The SPIDER Options Window appears.

3. Choose a Query Type. They are, in order of increasingly rigorous analysis:

   a. **Segment Match:** this is not a true mutation search, it will insist that the mass of the peptide returned is the same as that of the de novo sequence.

   b. **Non-gapped Homology Match:** this search will allow for transpositions, and single point mutations but not insertions or deletions.

   c. **Gapped Homology Match:** this search is the most rigorous, will find all types of mutations, but it is the slowest of the three search modes.

4. Choose if Leucine should match to Isoleucine without penalty, and/or if Lysine should be allowed to match Glutamine without penalty.

5. Choose how many of the best homologous peptides should be displayed after searching. For “Report top:”, 10 is the default, 200 is the maximum.
6. Choose PTM for consideration in analysis. Due to the ambiguity involved in mass sensitive homology searching, only fixed PTM are allowed for this type of analysis.

7. Choose a database to search. For this type of analysis it makes sense to rigorously search a small database that is known to contain homologous proteins.

8. Press the Ok Button.

SPIDER will search the database for homologous peptides, and attempt to consolidate these into protein hits as well. The result report will look much like the results for PEAKS Protein ID or inChorus searching.

After finding a homologous peptide in the database, SPIDER will decide what is likely a mutation and what is more likely a simple de novo sequencing error [resulting from certain combinations of amino acids having exactly the same mass – L/I, N/GG, AG/G, etc. As such it reconstructs the ‘real’ sequence from a de novo sequence and it’s homologue. This is highlighted on the Peptide Details frame of Peptide View.

The ‘real’ sequence takes some time to compute, so to save time, it is not constructed as soon as we load the report, only when we click on each peptide. Instead the peptide returned from the database is displayed in the Peptide View report. To display the constructed ‘real’ peptide, right click anywhere in the report and choose “Toggle Reconstruction” from the pop-up menu.

**SC**

**Merging SPIDER and Protein ID reports**

Mapping homologous peptides returned by SPIDER back to previously identified proteins can help us gain coverage. For this reason it’s advantageous to display the results from these two different types of analysis into one report. It’s easy to do using the inChorus search tool. Just click the inChorus button on the toolbar, or choose Tools -> inChorus Protein ID. Then check both SPIDER and PEAKS Protein ID checkboxes. Click the Import Button for each of these and choose the SPIDER and PEAKS Protein ID (or previous inChorus Protein ID) results that we want to combine. In the resulting report, SPIDER peptides will be in red, and all other peptides will be in blue.
Mutated peptides found only by SPIDER are shown in red.

SC

Creating a high-throughput workflow

In some situations, we may have many data sets that we wish to process all at once, and in the same way. PEAKS 4.5 allows us to do this kind of work, and with minimal effort on our part. By setting up a workflow, we can start a batch process of several data files and not worry about it until it is finished.
It is important to note that all the files we load will be processed in exactly the same way, using exactly the same parameters. If we want to do some differently than others, we must set up another workflow.

- File Options – Click the browse button (Folder Icon) to open a file chooser. From the chooser, select several files using shift+click or ctrl+click, and pressing the “OK” button. The list on the left will then show the selected files, the list on the right will show the names of the files that will be saved after analysis. Load more files by pressing browse again or remove them from the
list by right clicking on them. If these files comprise a large analysis, we should choose to ‘Close file after saving’ to save system resources.

- Data Refine – Choose how to filter and correct data for maximum utility.
- Auto de novo – Choose whether or not to do auto de novo sequencing. Note that PEAKS database search requires some de novo sequencing results.
- inChorus Database Search – Choose which protein identification programs to run the data.

**SCV**

**Using the Mass Calculator**

The Mass Calculator is a simple tool to help us determine the molecular weight of a peptide. To access the Mass Calculator, open the “Tools” menu and click “Mass Calculator”. The Mass Calculator will appear.

We can also load the mass calculator outside of PEAKS. To access the mass calculator without having to load PEAKS, click on the Mass Calculator’s icon in the start menu. It will appear in the same program group as PEAKS.

When using the mass calculator, remember to start with water.

We can click any of these buttons multiple times to repeatedly add that mass.

- Amino acids are represented by their single letter symbols. Clicking on an amino acid’s button will add it to the ‘sequence’ above, and add its mass to the mass of the peptide. Note that the peptide’s monoisotopic and average masses are both computed.
- Add a proton by clicking the “Proton” button. It will be represented by an \(<H^+>\) in the ‘peptide’ above.

- To compute the mass of the peptide as if it had been modified, select a PTM from the list, and press the “PTM” button to apply them to the peptide.

- If the PTM we wish to add does not appear in the list, we may wish to enter it's mass manually. To add a mass numerically, click the “Num” button and enter a numeric value in the dialogue box that appears. Press the “OK” button on the dialogue and the mass will be added to the sequence.

- To remove a mass that we’ve just added to the peptide, press “Undo”.

### SCV Predicting Retention Time
We can use PEAKS to predict the retention time of a peptide. Comparing the predicted retention time against the measured retention time can help us to evaluate correctness. The chemical properties of an amino acid sequence are used to predict its retention time. The prediction is adjusted using the actual retention time, as listed in the training data.

### Training the RT prediction algorithm
Since retention time varies between columns, and depends on experimental procedures (like flow rate), it is recommended that we keep a different training set for each column/procedure. The more data we have, the better, so keep adding to the training data, but only use peptides that are trustworthy. To train parameters for the prediction algorithm:

1. Select some trustworthy peptides from the Peptide View Tab to help with the training.

2. Right click, and from the pop-up menu that appears, select "Predict Retention Time" -or- Choose "Predict Retention Time" from the tools menu.

3. The Retention Time Prediction window appears.

4. Select a training file from somewhere on our system or define a new one. If we're adding to this training set, we should make sure that the data we're adding is for the
same type of column, and same procedure. Put a checkmark in the 'Add selected sequences to the training data file' checkbox. Otherwise, we can overwrite the training data file. If we check neither box, the training will proceed using the existing data in the training file.

5. Choose the type of column we used (both for the training data and the current analysis). 100 or 300 Angstrom column.

6. Choose a file to save the training results. These will be used as parameters for predicting RT on the peptides in our current analysis. Type in the Save Param textbox, or press the Browse Button and select or define a file.

7. Press the Start Training Button to begin the training. If we've chosen to add-to or overwrite the training data file, this will be done just before training begins. Training can take some time depending on the size of the training data. Press the Stop Training Button at any time to abort.

The training results in a parameters file being saved for future use. To judge how good our retention time predictions will be, based on this training, a Correlation value is given. This measures the quality of the trained prediction against the actual retention time. A correlation of 1 means a perfect match, above .9 is an excellent correlation, and above .85 is good. If the training is good, we can use it to predict retention time for our current analysis.

SC

Predicting RT after training

If the training is good enough, we can use the retention time prediction algorithm on our current analysis. It's likely that we'll use the RT prediction algorithm immediately after training. If this is the case, skip to step 3.

1. Open an ANZ file containing peptide sequences, and ensure the filename is selected.

2. Choose “Predict Retention Time” from the tools menu

3. Choose a Saved Parameters file by typing in the Save Param textbox, or clicking the Browse Button, and choosing an existing .par file.

4. Press the Ok Button. Retention time prediction will commence. It takes only a few seconds, and predicts retention time for all peptides in all reports saved in the file.

5. To view the predicted retention time, we'll have to toggle that column on the De Novo View tab and/or Peptide View tab in any report. If there's already a report open, we'll have to close and open it again, or untoggle and retoggle the column. Right click on any row in the report and mouse down to 'Toggle Column'. Here is presented a list of available columns. Click Predicted RT and DeltaRT.
Working with Results

Viewing, Post-Analysis, Exporting and Saving of results.

Results are what count. So the PEAKS 4.5 developers have made sure to present them simply, effectively and allow you to get them out of PEAKS easily. This chapter covers how to interpret the results, how to get the most out of them with minimal effort, how to present them to others, and how to save data and results together for future reference.

This chapter assumes that the user is familiar with all preceding chapters.
Viewing Protein identification results

To view Protein Identification results for a data file, we must have performed PEAKS protein identification or inChorus protein identification on that data file. The result from each protein identification search is represented by the time stamp and database searched (found just under a data file’s “Protein ID Result”). Click on one to display the results report. We can view results by peptide or by protein and check on the search parameters we used to generate these results. We can manipulate the contents of this multi-part report by filtering out what we don’t want to keep. Finally we can export the report and sub reports in a number of ways.

Search Parameters

Search parameters that we used when generating this report are preserved for future reference and are available by clicking on the search parameters tab.

Peptide View

Peptide View is available by clicking on the Peptide View tab. In tabular format, it displays relevant information about each peptide found. Since two peptides may match to one MS/MS spectrum, they are visually grouped together (using colour) by the MS/MS scan.

When we first load the report, it is sorted by MS/MS index number. This is analogous to the scan number or DTA file name (unless spectra have been merged). In the example above, there was one match returned for MS/MS spectrum #1. For spectrum #2 there were two; LDAINENK and IDALNENK. For spectrum #3, there were also two, and so on. Each peptide is given a score and the protein it matches is displayed for reference under “Accession No.” If the report was generated after inChorus protein ID, each peptide that had a search program agree (identify the peptide) is given a checkmark under its column.

The columns themselves are customizable. Right click anywhere in the report and choose Toggle Column from the pop-up menu. The sub-menu that appears shows a checkmark in each of the columns that are currently showing. Click any one of them to show or hide a column. These settings will apply to all our reports.

The peptide view list is sort-able. Click on a column header to sort the list using that column’s values. For example, sort by score. Clicking again on the same column header will toggle ascending/descending sort. Use ctrl+click to set a second sort – one can sort by Accession, then by score in this way. When sorting the grouping by MS/MS scan number will be retained except when sorting by Accession No.
Scores associated with each peptide represent the quality of the match and the number of programs in agreement on the peptide.

Clicking on this report will highlight the spectrum in the Peptide Data tree on the left. Select multiple spectra by clicking and dragging, using shift+click or ctrl+click. In this way, the highest scoring peptides may be selected and isolated for further analysis.

**Peptide Details and SPIDER Powered Alignment**

Whenever we click on a peptide in Peptide View, it’s details are shown in the panel below – the Peptide Details. The peptide details shows an alignment between the original de novo sequence for the selected spectrum, and the proposed peptide identified from the database. We can visually confirm a good match, strong evidence for having identified the peptide correctly. SPIDER is very good at mass-aware alignment of peptide sequences. The alignment is displayed in the following way:
If SPIDER searching is used to find homologous peptides, the alignment will account for differences between the de novo sequence and the homologue in the database, deciding what is a real mutation, and what is more likely a de novo sequencing error – thus constructing the 'Real' sequence.

Letters on a green background, and with vertical bars, indicate agreement. Letters on a red background indicate sequencing error. Colour codes on the de novo sequence letters still indicate positional confidence.

Letters on a blue background indicate uncertainty or mutation. + signs represent more likely mutations.

[brackets] indicate an equal mass substitution, common non-critical de novo errors. <these brackets> indicate an equal mass substitution and a mutation.

When simply identifying exact peptides from the database, using PEAKS Protein ID, or inChorus, there’s no need to reconstruct the ‘real’ sequence.

Letters on a green background indicate agreement. Letters on a red background indicate sequencing error – an equal mass substitution. Letters on a blue background indicate that it's uncertain why the sequences don’t match.

SPIDER powered alignment also generates an **RSD – a measure of the similarity between the two sequences**. A lower value indicates better agreement between the two; an RSD of 1 indicates no agreement.

\[
\text{RSD} \ (h=0, 1, 2, 3):
\begin{align*}
0.2 & \quad 0.1333 & \quad 0 & \quad 0
\end{align*}
\]

The h value is the tolerance for equal mass errors of length 0, 1, 2, or 3. Thus, in the example above, when we allow equal mass errors of length 2, the sequences agree exactly.
The ‘real’ sequence takes some time to compute, so to save time, it is not constructed as soon as we load the report, only when we click on each peptide. Instead the peptide returned from the database is displayed in the Peptide View report. To display the constructed ‘real’ peptide, right click anywhere in the report and choose “Toggle Reconstruction” from the pop-up menu.

**Protein View**

**Protein View** is accessed by clicking on the Protein View tab. It collects all the peptide identifications together, summarizes which proteins were present in the sample, and groups homologous proteins together. The same information is displayed in the Peptide View as in this Protein View; however, the results are organized to best enable us to evaluate at the protein level.

The top section of this view (shown above) behaves like an index, listing each protein found in the sample. **Very similar proteins, containing the same set or a subset of the matched peptides, are clustered together.** To see the full list of proteins within each cluster click the ‘+’ sign. In the example above, the Bovine Serum Albumin cluster has been expanded to reveal several similar proteins. “gi|162648 albumin [Bos Taurus]” is listed at the top, having the most high scoring unique peptides – reflected in a high UP (Uniqueness of Protein) score. Clicking in the same place (now a ‘—’ sign) will collapse the cluster. Not all similar proteins can be clustered, not necessarily having the exact same set of peptides – maybe they have an extra peptide hit associated. So we have to decide which cluster is best; fortunately, the ‘extra peptide’ is usually low scoring, so we can reject it. This will be reflected in a low UC (Uniqueness of Cluster) score.

This view is helpful when building a summary that can be sent to a customer/collaborator. Simply right click to export to a MS Excel file. We can export
interesting parts of the report or a whole summary. Mark proteins of interest by clicking their checkboxes and export protein and peptide information for those.

- Or highlight a homologue group and export proteins and peptides in that group.
- Or just export the whole report.

The columns themselves are customizable. Right click anywhere in the report and choose Toggle Column from the pop-up menu. The sub-menu that appears shows a checkmark in each of the columns that are currently showing. Click any one of them to show or hide a column. These settings will apply to all our reports.

Whenever we click on a protein, the Matched Peptides panel (bottom) changes to display the spectra (and peptides) that were found to be supporting evidence for that protein. The columns in this list are the same as the ones we choose for ‘Peptide View’. It may be necessary to scroll this panel down to see the complete list. Clicking a peptide in this list brings up the Main Processing Window for the corresponding spectrum, and displays the ions that were found in support of this peptide (shown below).

Scrolling down the Matched Peptides panel reveals the complete sequence of the highlighted protein with the matched peptides highlighted in red. The strength of the red colour is a visual indication of the confidence in that portion of the protein. Where
an EST database was used, the translated sequence is shown (with all six reading frames concatenated).

**De Novo View** usually shows de novo sequences for spectra that could not be explained by matching to the identified proteins (as listed in Protein View), but the exact behaviour is customizable. See below for a discussion of filters and how they relate to the De Novo View. The list of peptides in the De Novo View behaves quite like the list of peptides in Peptide View. Differences are as follows:

- Note that since PEAKS 4.5 uses de novo sequences to help out in the database search, it is sometimes useful to see which de novo sequences were use to help in the search. Each sequence that is used in this way has a checkmark in it’s PID column.

- The parameters used to generate a de novo sequence may be slightly different from the database search parameters. As such, PEAKS 4.5 records how each de novo sequence was generated – this information is in the [Source program] column.

- Positional confidence information is available in de novo sequencing results. Colour codes show the strength of confidence in an individual amino acid. The colour codes are the same as those used in the main processing window (red is the best, then purple, then blue, then black). Positional confidence can also be seen by holding the mouse over a sequence.

**Result filters** are accessed by clicking on the ‘Filter Pane’ tab. These allow us to manipulate the contents of the whole multi-part report. For instance, one may choose to discard any protein hit that was not supported by at least two high scoring peptides. The filters are very flexible, allowing you to create filters on any of the fields (score, mass, search engine, etc) and any number of fields. The filters are also cascading such that removing a protein from the results removes all peptides associated with only that protein and vise verse. But all this can be a bit complicated, so please read on to the section “Post analysis of results – preparing for publication”.

**SCV**

**Comparing proteins - multiple sequence alignment**

After having identified some proteins, it may be useful to see how their sequences compare. This is particularly useful when trying to identify specific isoforms of proteins or antibodies. To display a multiple sequence alignment between two proteins, click on the time and date stamp underneath a filename in the peptide data tree (left). This brings up a multi-part protein ID report. Click on the **Protein View tab** to see a list of identified proteins. **Mark a few proteins by clicking on their checkboxes** in the ‘Marked’ column. Then, in the bottom panel of this view, click the **MSA tab**. It will be blank to start with, but click one of the two buttons on this panel to:
generate the MSA and display it in the default web browser

generate the MSA and display it on the MSA panel.


SCV

The table of values

PEAKS 4.5 tracks all kinds of useful information about our experiments, the results associated with them, and calculated measures of fit between them.

SCV

What do the columns mean?

<table>
<thead>
<tr>
<th>Protein View</th>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accession</td>
<td>The GI, Accession or other unique identifier for this protein as recorded in the database that was searched.</td>
<td></td>
</tr>
<tr>
<td>Mass</td>
<td>The calculated mass of this protein</td>
<td></td>
</tr>
<tr>
<td>Display</td>
<td>A graphical coverage map. Blue areas represent parts of the sequence that have been explained by the identified peptides.</td>
<td></td>
</tr>
<tr>
<td>Score(%)</td>
<td>A value from 1 to 99 representing the confidence we have in this protein identification – calculated from the confidence on the ten best peptide hits for this protein, and normalized against the other identified proteins.</td>
<td></td>
</tr>
<tr>
<td>Coverage(%)</td>
<td>The number of amino acids in the protein sequence that have been explained by the identified peptides. Expressed as a percentage of the total protein’s sequence length.</td>
<td></td>
</tr>
<tr>
<td>Query Matched</td>
<td>The number of spectra explained by matching to a peptide from this protein.</td>
<td></td>
</tr>
</tbody>
</table>
Marked
A multi-function checkbox. By default unchecked, but we can use this to select proteins for export or multiple sequence alignment.

Description
The part of the protein's header information as parsed from the database, usually it contains the name of the protein.

UniqueIDs
The number of peptides that uniquely match to this protein.

UC Score
A measure of uniqueness, calculated from the confidence on the ten best peptide hits that are unique to this cluster of proteins. Remember, PEAKS groups similar proteins together. So UC measures how unique the cluster is.

UP Score
A measure of uniqueness, calculated from the confidence on the ten best peptide hits that are unique to this protein. Usually we use this only when deciding which is the best protein within a cluster.

### Peptide View

<table>
<thead>
<tr>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpecID</td>
<td>A unique identifier for the MS/MS spectrum. This differs from a scan number since we may have merged several scans together.</td>
</tr>
<tr>
<td>Scan</td>
<td>The scan number.</td>
</tr>
<tr>
<td>RT</td>
<td>Retention time (elution time) for the peptide as recorded in the scan header.</td>
</tr>
<tr>
<td>File</td>
<td>If using DTA files, the file name can be displayed here.</td>
</tr>
<tr>
<td>m/z</td>
<td>The measured mass/charge value, in Daltons, for the peptide.</td>
</tr>
<tr>
<td>z</td>
<td>The calculated charge value for the peptide.</td>
</tr>
<tr>
<td>Mass</td>
<td>Calculated using the measured m/z and calculated z, we use this as the experimental mass of the peptide.</td>
</tr>
<tr>
<td>Peptide</td>
<td>The sequence of the peptide (including modifications if present) as identified from the database.</td>
</tr>
<tr>
<td>Mr(Calc)</td>
<td>The sum of the theoretical mass of the residues that form the identified peptide sequence from the database.</td>
</tr>
<tr>
<td>*RSD</td>
<td>A measure of the similarity of the identified peptide's sequence and the original de novo sequence obtained for the same spectrum. This is a value from 0 to 1, with zero being perfect agreement. Useful for finding false negatives and false positives.</td>
</tr>
<tr>
<td>Con. Peptide</td>
<td>For peptides returned by SPIDER, we must decide what is a true mutation and what is just a de novo sequencing error. The Constructed peptide does just this. Because of the complexity of the calculation, this can take a while to compute.</td>
</tr>
<tr>
<td>Delta (Mass)</td>
<td>The difference between Mr(Calc) and Mass, in Daltons.</td>
</tr>
<tr>
<td>Spectrum Quality Score(%)</td>
<td>A value from 0 to 1 estimated from the spectrum to refer to spectrum quality. Attributes like signal to noise, total intensity, and spectrum tagging are used. The confidence we have in the identified peptide, represented as a percentage value (1 to 99). 50% and greater represents a good match. If only one search engine is used, that search engine’s score will be displayed here. If inChorus searching is used, a meta-score will be displayed here.</td>
</tr>
<tr>
<td>Start</td>
<td>The start position of this peptide in the protein sequence.</td>
</tr>
<tr>
<td>End</td>
<td>The end position of this peptide in the protein sequence.</td>
</tr>
<tr>
<td>Accession No.</td>
<td>The accession, or unique identifier of the best protein this peptide matches to.</td>
</tr>
<tr>
<td>HitID</td>
<td>A unique identifier for the peptide match. This differs from SpecID since the same spectrum may have several likely matches in the database.</td>
</tr>
<tr>
<td>Predicted RT</td>
<td>The retention time as predicted from the peptide sequence.</td>
</tr>
<tr>
<td>Delta RT</td>
<td>The difference between the measured RT and the Predicted RT. If our peptide identification is good, and the prediction is accurate, the Delta RT should be low.</td>
</tr>
<tr>
<td>Search engines</td>
<td>For inChorus searching, we list a checkmark in this column to indicate which...</td>
</tr>
</tbody>
</table>
search engines reported this peptide. Where two search engines agree, there will be two checkmarks – good evidence for having found the right match!

Search engine scores

For inChorus searching, we expose the original score from each search engine.

*The RSD measure was proposed by Sergey Pevtsov, Irina Fedulova, Hamid Mirzaei, Charles Buck, and Xiang Zhang, J. Proteome Res., 5 (11), 3018-3028, 2006

<table>
<thead>
<tr>
<th>De Novo View</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Value</td>
<td>De Novo View Description</td>
</tr>
<tr>
<td>Scan</td>
<td>The scan number.</td>
</tr>
<tr>
<td>RT</td>
<td>Retention time (elution time) for the peptide as recorded in the scan header.</td>
</tr>
<tr>
<td>File</td>
<td>If using DTA files, the file name can be displayed here.</td>
</tr>
<tr>
<td>m/z</td>
<td>The measured mass/charge value, in Daltons, for the peptide.</td>
</tr>
<tr>
<td>z</td>
<td>The calculated charge value for the peptide.</td>
</tr>
<tr>
<td>Mass</td>
<td>Calculated using the measured m/z and calculated z, we use this as the experimental mass of the peptide.</td>
</tr>
<tr>
<td>Peptide</td>
<td>The sequence of the peptide (including modifications if present) as determined by de novo sequencing.</td>
</tr>
<tr>
<td>Score(%)</td>
<td>The confidence we have in the peptide sequence, represented as a percentage value (1 to 99). Calculated by statistical analysis of all possible peptide matches to a spectrum.</td>
</tr>
<tr>
<td>Source Program</td>
<td>Since we may have done de novo sequencing several times on any given spectrum, this column records which parameters were used to get a given de novo sequence.</td>
</tr>
<tr>
<td>Predicted RT</td>
<td>The retention time as predicted from the peptide sequence.</td>
</tr>
<tr>
<td>Delta RT</td>
<td>The difference between the measured RT and the Predicted RT. If our peptide identification is good, and the prediction is accurate, the Delta RT should be low.</td>
</tr>
<tr>
<td>Rank</td>
<td>Starting from 0, this simply counts the best de novo sequences for a given spectrum and parameter set, in order of score.</td>
</tr>
<tr>
<td>Data Quality</td>
<td>A value from 0 to 1 estimated from the spectrum to refer to spectrum quality. Attributes like signal to noise, total intensity, and spectrum tagging are used.</td>
</tr>
</tbody>
</table>

SCV Toggling Columns

Still, the information in the report might be too intensive. Some users might not care to see the start and end position of the peptide in the protein, while some users may wish to see even more information. PEAKS 4.5 allows us to customize which columns are shown. Right-click anywhere in De Novo View or the top section of Peptide View or Protein View. In the pop-up menu that appears, hold the mouse over Toggle Column. A sub-menu will appear showing a checkmark in each of the columns that are currently showing. Click any one of them to show/hide a column. PEAKS will remember which columns we select, and present us with these columns each time we load a report.
SCV

Filtering results

PEAKS 4.5 provides us with an exhaustive list of all proteins and peptides that can be found in the sample. But since everyone has their own criteria of what information is required in the report and what is an acceptable result, a good hit, we must have a way of filtering out the less critical information to leave us with the essentials.

In order to manipulate results, we must, of course, have done some protein identification using PEAKS Protein ID or inChorus searching. Click on the time and date stamp associated with this result under its file name in the peptide data panel on the left. Once the report loads, click on the “Filter Pane” tab to bring up the filtering options.

Choose a filter from the ‘Possible Filters’ list on the left by clicking on it. Options for this filter will appear below, in the [Edit filter] frame. Once we’ve set these options, we press ‘Add filter’ to bring it to the list of ‘Selected Filters’. If we’d like to add another filter, we can repeat the process, continuing to add as many filters as necessary. In this way it is also possible to have two filters on the same property; we can set a range of
protein mass, for instance, by applying one filter on the upper bound of the mass and adding another filter to be the lower bound of the mass. We can also have more complex filters that involve multiple properties. The example above shows filters that allow only proteins with more than one high scoring (greater than 60% score) peptide, a standard requirement for publication.

Filters are grouped into three basic types to reflect what they act on: De Novo filters act to remove proposed de novo sequences, Peptide Filters act to remove peptides found in the database from the report and Protein filters act to remove proteins from the report. **All columns, as described above, can be filtered upon.**

The filters cascade through each view in the multi-part report. For instance, filtering out a protein (by mass perhaps) will remove it from the Protein View and remove all peptides associated with that protein from the Peptide View list. The manner in which the De Novo View is linked can be specified by the user (illustrated on the next page).

Filter sets can be saved and re-used between sessions, but remember to hit Apply Filters to use them.

Once all the filters we need are set up and showing in the Selected Filters list, we must press the Apply Filters button to use them on our results. For larger reports this can take a few seconds, during which time the tabs of the multi-part report will be grayed out. After completion, we can click on any of the tabs to see how the results have been affected.
How filters act on de novo sequences

PEAKS 4.5 automatically relates de novo sequencing results to protein ID results. This makes it easy to see which de novo sequences correspond to known contaminants and which are truly new peptides. The relationship is always drawn back to the data, not necessarily the sequence, which helps highlight important sequence variations. PEAKS 4.5 users can set what is displayed on the list of de novo sequences, according to the following diagrams:

Option Selected
(on Filter Pane tab)

What De Novo View Displays
(represented in grey)

How Filters are linked

Filtering out (removing) a de novo result does not affect what is displayed in the Protein View tab or Peptide View tab.

Filtering out (removing) a de novo result also removes that spectrum from consideration in the Peptide View tab and Protein View tab.

Again, filtering out (removing) a de novo result also removes that spectrum from consideration in the Peptide View tab and Protein View tab.
Filter examples

Any column of information in any part of the Protein ID report can be filtered. Filters cascade throughout the report – i.e removing a protein removes all its associated unique peptides, and removing all peptides from a protein removes that protein too. Each filter can be applied several times over. So it can get a little complex. To illustrate, here are a few examples:

Publication

Show proteins that have two high-scoring hits:

- Add the Protein Filter called Query and in the [Edit Filter] section choose ‘greater than’ and type ‘1’ in the box (without the quotes).
- Then add the Peptide Filter called ‘Score’ and in the [Edit Filter] section choose ‘greater than’ and type ‘50’ in the box (without the quotes).
- Press the ‘Apply Filters’ button.

Digging for a protein by name

To illustrate this process, let us find a protein that contains the word ‘human’ or ‘rat’ in the database entry’s description, but not Keratin or Trypsin.

- Add the Protein Filters called ‘Desc’
  - In the [Edit Filter] section, we’re required to type in a regular expression (regex). This allows us to use wildcards.

<table>
<thead>
<tr>
<th>Wildcard</th>
<th>Meaning</th>
<th>Example</th>
</tr>
</thead>
</table>
| .*       | "Anything of any length" | .*human.* Will find anything that contains the word 'human', with anything before and anything after.
| | | .*(human|rat|RAT).* Will find anything that contains the word ‘human’, or the word ‘rat’ or the word ‘RAT’, with anything before and anything after.
| ?!       | "Not" (use brackets) | (?!.*(Keratin|Trypsin).)*.*(human|rat).* will find anything containing human or rat but not Keratin or Trypsin
| [ ]      | "Any of these characters" | .*((Hh)uman|[Rr]at).* will find anything containing the words Human, human, Rat or rat.

- So type in the regex: (?!.*(Keratin|Trypsin).)*.*(human|rat).* and press the Enter key. If PEAKS confirms that this is a valid regular expression, it will put a check in the ‘Valid Java Regex’ box. You may then press the Add Filter button.

The Regex must match the whole description, you can’t just type “human”.


Press the ‘Apply Filters’ button.

**Setting a protein mass range**
If we know the approximate mass of the proteins we’re interested in, we should eliminate all proteins that are not close in mass.

- Add two filters: “Protein Filters: Mass >12000” and “Protein Filters: Mass < 32000”
- Press the ‘Apply Filters’ button.

**SCV Saving/Loading Filter sets**
We’re sure to use some common filters repeatedly. To save a Filter Set, press the ‘Save As’ button on the filter pane. This saves all filters that we’ve added to the ‘Selected Filters’ list. To recall a saved Filter Set, select it from the drop down list in the top right corner of the filter pane. This will simply populate the ‘Selected Filters’ list with our saved filters. Don’t forget to press the ‘Apply Filters’ button to apply them.

**SCV Using a default filter**
We may prepare our results the same way each time; in which case it makes sense to set up a filter that will be automatically applied each time we load a report. Select a filter from the list of saved filters in the top right corner of the Filter Pane. Press the Set Saved as Default Button at the bottom of the Filter Pane. This filter will be displayed in the Default Filters box at the top of the Filter Pane, and will be applied automatically just after a report is loaded. Be careful, if your default filter is very stringent, it can sometimes remove everything! To remove a default filter, press the Clear Default Button at the bottom of the Filter Pane. At any time you can choose or create another filter which will override the default.

**SCV Exporting Results**

**WYSIWYG reports**
PEAKS 4.5 allows us to create interactive reports to share with collaborators, colleagues and clients. The reports are available in HTML or Microsoft Excell (.xls) formats and follow a ‘What you see is what you get’ (WYSIWYG) philosophy. All the information you see on screen in PEAKS 4.5 will appear in the exported report. For this reason, it is important that we complete results filtering and toggling columns before exporting a report.
In order to export a report, first choose the view to export by clicking on the De Novo View, the Peptide View tab or the Protein View tab. Next, right-click anywhere in the top section of the selected view and choose ‘Export Report’ from the pop-up window that appears.

The report export dialogue will appear. One must choose either HTML or XLS format and specify a file name. PEAKS 4.5 will load the report after it has been created, but it makes sense to always place the report in a place we can find it later on. By default the filename of the report will include the time and date the report was created. To change the location/filename one can either type it in to the Filename box or click the folder icon to browse to a new location.

The bottom part of the Report Export dialogue asks us what we would like to include in the exported report. When exporting from Peptide View or De Novo View we have the choice of exporting the whole list or the selected ones. When exporting from Protein View we have some more options, allowing us to include peptide details, the complete protein sequence, a listing of proteins containing the same set or a subset of the same peptides, or to include only marked proteins. Press the OK button to complete the export.

**SCV**

**Exporting peak lists**

To export data to a PKL file, we select the data file (not an individual spectrum) to export. Then, from the “File” menu, select “Export”, then “Export PKL File”. The spectrum data will be saved in PKL format, but all sequencing and protein data will not be preserved.

**SCV**

**Exporting Sequences by spectrum**

For most users, the WYSIWYG reports exporting will be enough, but PEAKS 4.5 also provides other ways to export. To export peptide sequencing results to a FASTA format file, select the data file (not an individual spectrum) to export. Then, from the “File” menu, select “Export”, then “Export Peptide Sequence”. The peptide sequencing data will be saved in FASTA format, one entry for each spectrum, but will not retain any spectrum data.
**SCV Exporting Peptide Sequences**

To export peptide sequencing results to an HTML table, select the data file (not an individual spectrum) to export. Then, from the “File” menu, select “Export”, then “Export HTML File”. PEAKS will then ask us which results we would like to export. We can choose from any *de novo* sequencing or Protein ID run we have done. Each will be listed with the parameter set we used. The difference here is only that the results are organized by spectrum rather than by peptide as in the WYSIWYG exporting.

**SCV Exporting Exclusion Lists**

If we are repeating an experiment, it can be useful to exclude the peptides we have already scanned and analyzed. In this way, we can focus on some potentially interesting but lower intensity PEAKS. To tell the instrument to do this we must create an exclusion list. PEAKS 4.5 has the ability to create an exclusion list that will be compatible at least with Waters/Micromass instruments. To create the exclusion list, simply select a few spectra from the peptide data tree on the left, then right click and choose “Export to Exclusion list” from the pop-up menu that appears. The selection of spectra can also be done using by clicking in the Peptide View or De Novo View of a Protein ID report.

**SCV Exporting high-resolution spectral images**

It is often useful to provide visual aids in reports, posters and papers. As such, PEAKS 4.5 provides the facility to export high quality images of a spectrum or part of a spectrum zoomed in. In any case, PEAKS allows us to export exactly what’s shown on the screen. So to export an image:

1. Bring up the main processing window for an MS/MS spectrum by double-clicking on its precursor ion mz, or by double clicking a peptide in any of the reports. The main processing window shows us the spectrum of interest, along with any peptides matched to it.

2. Set up the colour, typeface and size of labels and PEAKS on the spectrum. To do so, choose Edit menu → Configuration → Environment Preferences. Within the Environment preferences dialogue, choose the ‘Colour’ or ‘Display’ tab to get at the relevant settings.

3. *(optional)* If we want to export a picture of the Ion Table, we should configure what columns are shown in the ion table before the export. To do so, select one of the Ion Table Editor tabs from the Environment Preferences dialogue.

4. *(optional)* If we want to export a zoomed in region of the spectrum, we must do so now by clicking and dragging horizontally over the area of interest.
5. Click the export results button on the main processing window toolbar, or right-click on any frame within the main processing window and choose “Export Image” from the pop-up menu.

6. The Image Export Options Dialogue appears. On this window you can export:

   a. Choose what you want to export.
   b. Select a file type. Four popular formats are supported, but either GIF or PNG formats are recommended.

   **SVG is a vector graphics format**

   Vector graphics means the information is stored as math representing lines and shapes, not pixels. Essentially, this means that your spectrum will be easier to edit, and won’t be affected by resizing. Several SVG editors are available (Adobe Illustrator, or freeware like Inkscape), or you can convert the SVG into an EMF file for MS Word.

   c. Choose a file name. We can browse by clicking the folder icon or just type a file name in the box.
   d. *(optional)* Specify width and height dimensions in pixels. In some cases the size is predetermined by the size of the frame on our screen. Otherwise we can increase these numbers to make the picture bigger.
e. To export high resolution images, increase the resolution by a few hundred percent.

f. Press ok and you’re done.

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**SCV Saving Results**

Saving results will preserve our work for later use. Saving files in PEAKS’s .ANZ format will preserve spectrum data, manual *de novo* sequence information, automatic *de novo* sequence information, protein identification results, and information about any PTM that were found in sequence.

To save the results of our analysis, we first select the data file we wish to save in the Peptide Data Frame.

To save, click the save icon in the main window toolbar, select “Save” from the “File” menu or right click on the data file and select “save” from the popup menu. This will save the processed spectra in .ANZ format and of the same name as the data file we opened.

To change the name of the ANZ file, choose “Save as” from the “File” menu, or right click on the data file and select “Save as” from the pop-up menu. We may then change the file name.

To save all currently opened data files, select “Save all” from the “File” menu.
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Reference: PEAKS Paper

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Bin Ma, Kaizhong Zhang, Christopher Hendrie, Chengzhi Liang, Ming Li, Amanda Doherty-Kirby, and Gilles Lajoie. PEAKS: Powerful Software for Peptide De Novo Sequencing by Tandem Mass Spectrometry. Rapid Communication in Mass Spectrometry 17(20): 2337-2342. 2003