

i2MassChroQ is one of the PAPPSO facility software projects

12MASSCHROQ USER MANUAL

Free and Open Source Protein Identification Software

12MASSCHROQ 1.2.3

12 MassChroQ User Manual: Free and Open Source Protein Identification Soft-

WARE

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Thierry Balliau and Marlène Davanture are warmly thanked for their outstanding technical help while writing this user manual.

i2MassChroQ

http://pappso.inrae.fr/en/bioinfo/i2masschroq/ 🗗

This book is part of the *i2MassChroQ* project.

The *iz MassChroQ* project is the successor of the Java language-based homonymous project. This project is a full rewrite of the former project in the C++ language, with many new features added.

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1.2.3

Revision History

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02 November 2023 Filippo Rusconi

• Added entire section with step-by-step procedure to describe the new *MSstats* interface inside of *i2MassChroQ*.

Revision 0.7.24

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• Start actually documenting the software with the Preface.

Revision 0.7.24

- Filippo Rusconi
- Very first setting up of the user manual using the mineXpert2 project as a template.

28 May 2021

21 April 2021

DEDICATION

To all the admirable people acting in the *"Free Software Movement"* for a better and more ethical computing world

To all the readers who helped with this manual.

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Preface

1 Software feature offerings and intended audience

This manual is about the *i2MassChroQ* protein identification software project.

i2MassChroQ has the following features:

- Load mass spectometry data files in the mzXML or mzML format, thanks to the excellent *libpwiz* library of ProteoWizard¹ fame.
- Configure the way the peptide/mass spectrum matches (PSM) are to be performed;
- Configure the database files to be used (target organism databases and contaminant databases);
- Use the MS/MS data in the file to feed the *X!Tandem* program that produces peptide identification results by matching the measured ion masses with peptide fragments calculated *in silico* on the basis of the databases contents;
- Perform the protein inference step that leads to reliable protein identifications on the basis of the peptide identifications performed by *X!Tandem*
- Display the data obtained at any step in powerful ways in a unified graphical user interface to allow the user to inspect the peptide identifications and also control the way these identifications are used to infer the protein identifications.
- Export the data after the results exploration above in a variety of formats.
- Perform quantitative proteomics on the basis of the results obtained at the previous steps.
- Perform bio-statistical analyses on the quantitative proteomics data obtained at the previous step.

2 FEEDBACK FROM THE USERS

We are always grateful to any constructive feedback from the users.

The PAPPSO software team might be contacted via the following contact page:

HTTP://PAPPSO.INRAE.FR/EN/TRAVAILLER_AVEC_NOUS/CONTACT/ ♂ (search for team members having the "Bioinformatics" specialty mentioned, like Olivier Langella or Filippo Rusconi).

I HTTP://PROTEOWIZARD.SOURCEFORGE.NET/ ₽.

3 Program and Documentation Availability and License

The programs and all the documentation that are shipped along with the *i2MassChroQ* software suite are available at http://pappso.inrae.fr/en/bioinfo/xtandempipeline/ ? . Most of the time, a new version is published as source, and as binary install packages for *MS-Windows* (64-bit systems only).

For *GNU/Linux*, binary packages are created locally (see HTTP://PAPPSO.INRAE.FR/EN/BIOINFO/XTANDEM-PIPELINE/DOWNLOAD/?) but are also built in the *Debian*² autobuilders and are uploaded to the distribution servers. These packages are available using the system's software management infrastructure (like using the *Debian*'s <u>apt</u> command, for example, or the graphical application).

The software and all the documentation are all provided under the Free Software license *GNU General Public License, Version 3, or later, at your option*. For an in-depth study of the *Free Software* philosphy, the reader is kindly urged to visit HTTP://www.gnu.org/philosophy **?**.

² HTTP://WWW.DEBIAN.ORG/ ₽

i Generalities

In this chapter, I wish to introduce some general concepts around the *i2MassChroQ* program, the reference to be used to cite the software in publications, the building and installation procedures.

I.I HISTORY OF THE PROJECT

i2MassChroQ is the successor of the *X!TandemPipeline-Java* project that has seen the following changes along the years:

• Full rewrite of the *X!TandemPipeline-Java* program from Java to C++17. The Java-based software program had been published in Olivier Langella, Benoît Valot, Thierry Balliau, Mélisande Blein-Nicolas, Ludovic Bonhomme, and Michel Zivy (2016) *X!TandemPipeline: A Tool to Manage Sequence Redundancy for Protein Inference and Phosphosite Identification.* in *J. Proteome Res.* 2017, 16, 2, 494–503. https:// doi.org/10.1021/acs.jproteome.6b00632.



Tip

Before the integrations described below, the product of the rewrite has been called transitorily *X!TandemPipeline++* (or *xtpcpp*). That name might appear in some places while the code/documentation is being revised to change its name to *i2MassChroQ*.

- Integration into the new software of the *MassChroQ* software project that was developed as a standalone C++ software piece. *MassChroQ* is a software project that was developed to perform quantitative proteomics in a variety of modes (label-free or with labelling).
- Unfinalized integration of the *MCQ~R* project that was developed as a standalone project. *MCQ~R* is a GNU R project aimed at performing bio-statistical analyses on the quantification analysis performed by *MassChroQ*.

The *i2MassChroQ* project encompasses three main quantitative proteomics fields of endeavour:

- Database search, peptide identification and protein inference. The database search is actually performed by *X!Tandem* and is started seamlessly by *i2MassChroQ*. Protein grouping is performed by original code in *i2MassChroQ*.
- Quantitative proteomics, mainly based on area-under-the-curve processes (requires the full mass data set to extract ion current chromatograms, XIC). This part was historically performed by the *MassChroQ* software program.
- Bio-statistical analysis of the quantification data. This part was historically performed by the *MCQ~R* GNU R-based package (unpublished software as of yet).

I.2 WHAT DOES *i2MassChroQ* Stand for?

The *i2MassChroQ* software project aims at providing users with an integrated software solution for quantitative proteomics. As described in detail in another chapter of this book, quantitative proteomics involve a number of steps that can be enumerated in sequence below:

- Search databases to connect MS/MS spectra to peptide sequences. This step is called *identification*;
- Apply logic to reliably identify proteins based on the peptides identified at the previous step. This step is called *inference*;
- Optionally perform quantification of the *i*dentified peptides and *i*nferred proteins. *i2MassChroQ* has area-under-the-curve quantitative proteomics capabilites that are based on precursor peptide ion current extraction from the mass spectrometric data. The extracted ion currents are then plot like chromatograms: intensity as a function of retention time. This analytical process thus somehow involves "*Mass Chromatograms*" for the *Q*uantification.

From the sequence above, the *izMassChroQ* name becomes self-explanatory!



Tip

It is however possible (and encouraged) to mentally read *i2MassChroQ* as "*I too MassChroQ*!"

1.3 TRANSITIONING FROM X!TandemPipeline то i2MassChroQ

The previous X!TandemPipeline version of this software did store configuration data in the local configuration directory and in the <u>PAPPSO/xtpcpp.conf</u> file. In order to preserve these configuration data after having transitioned from X!TandemPipeline to i2MassChroQ, please, rename that configuration file to <u>PAP-</u> PSO/i2masschroq.conf.

I.4 GENERAL CONCEPTS AND TERMINOLOGIES

This section describes the general concepts at the basis of the analysis of proteomics data that one needs to grok in order to properly assimilate the workings of the *i2MassChroQ* software.

1.4.1 BOTTOM-UP PROTEOMICS OR TOP-DOWN PROTEOMICS?

Proteomics is a mass spectrometry-based field of endeavour that is aimed at characterizing the "protein complement" of a given genome. The protein complement of a genome is the set of proteins that are expressed at a given instant in the life of a cell, a tissue or an organ, for example. Characterizing that protein complement actually means identifying the proteins expressed by a given living cell or tissue or organ. Optionally, if feasible, the characterization of post-translational modifications might be desirable.

There are two main variants of protemics: "bottom-up" proteomics and "top-down" proteomics:

- The first variant—bottom-up proteomics—identifies proteins on the basis of the identification of all the peptides obtained by first digesting all the proteins of the sample using an enzyme of known specificity. In this variant, the sample that is injected in the mass spectrometer is the resulting peptide mixture (first resolved by high performance liquid chromatography). The identification of the proteins contained in the initial sample is performed in a number of steps that are actually the focus of *i2MassChroQ*. Indeed the *i2MassChroQ* software is a bottom-up-oriented software program.
- The second variant—top-down proteomics—identifies proteins on the The second variant identifies proteins on the basis of intact proteins directly injected in the mass spectrometer. Of course, it might be necessary to fragment the proteins in the mass spectrometer and to use the fragments to actually identify the protein. However, the fact that the protein is first detected and analyzed as one entity (and not as set of peptides), allows for some very useful discoveries, like the identity and number of post-translational modifications, for example.



Note

At the moment, *i2MassChroQ* does not handle top-down proteomics data: it is a bottom-up proteomics software project.

1.4.2 Typical cycle of a mass spectrometer data acquisition

Once the initial sample, containing all the proteins to identify, has been digested using a protease of known cleavage specificity (trypsin, typically), the peptidic mixture (that might be highly complex) needs to be resolved as much as possible using chromatography. In the vast majority of the proteomics experimental settings, the chromatography setup is connected to the mass spectrometer so that when the gradient is developed, all the peptides are immediately injected "on line" to the mass spectrum ion source.

The mass spectrometer runs an analysis cycle that can be summarized like the following:

- Acquire a full scan mass spectrum of the whole set of ions at a given chromatography retention time. This kind of mass spectrum is called a MS spectrum;
- Enter a loop during which ions having the most intense signal are subjected in turn to collision-induced dissociation (CID), that is, are fragmented by accelerating them against gas molecules in a fragmentation cell. The mass spectra that are collected at each one of these fragmentation acquisitions are called MS/MS spectra because they are obtained after two mass analysis events: the first event is the measurement of the intact peptide ion's m/z value (full scan mass spectrum) and the second event is the measurement of all the obtained fragments' m/z values (MS/MS scan).

Each instrument records all the MS and MS/MS spectra in a raw data format file that is specific of the vendor. Free Software developers cannot know the internal structure of the files. To use the mass spectrometric data, they need to rely on a specific software that performs the conversion from the raw data format to an open data format (mzML). That program is called *msconvert*, from the *ProteoWizard* project.



Note

Mass spectrometrists used to call ions that were analyzed in full scan mass spectra "parent ions". They also used to call fragment ions arising upon fragmentation of a parent ion "daughter ions". This terminology has been deprecated and has been replaced with "precursor ion" and "product ion", respectively. In our document, we thus use the new terminology.

1.4.3 Outline of an i2MassChroQ working session

i2MassChroQ loads mzXML- and mzML-formatted files and needs for its operations to have accesss to all the MS and MS/MS spectra. Once data files have been loaded, *i2MassChroQ* allows the user to perform the following tasks, that will be detailed in later chapters:

- Configure the *X!Tandem* database searching software (that is, the software, external to *i2MassChroQ* that actually performs the peptide-mass spectrum matches);
- Run the *X!Tandem* software and load its results;
- Display the results to the user in a way that they can be scrutinized and checked. The peptide identification results serve as the basis for another processing step that is integrally performed by *i2MassChroQ*: the "protein inference". That step aims at using the peptide identifications to actually craft a list of proteins identities. The user is provided with various means to control that step in various ways.
- Optionally start the *MassChroQ* module to perform the quantitative proteomics on the identification data checked at the previous step.
- Optionally start the *MCQ~R* module to perform the bio-statistical analysis of the quantitative proteomics data obtained at the previous step.

1.5 CITING THE *i2MassChroQ* SOFTWARE.

The *i2MassChroQ* software is published : Olivier Langella, Thomas Renne, Thierry Balliau, Marlène Davanture, Sven Brehmer, Michel Zivy, and Filippo Rusconi (2024). *Full Native timsTOF PASEF-Enabled Quantitative Proteomics with the i2MassChroQ Software Package* in *J. Proteome Res.* 23, 3353–3366. doi: HTTPS:// DOI.ORG/10.1021/ACS.JPROTEOME.3C00732

I.6 INSTALLATION OF THE SOFTWARE

The installation material is available at $http://pappso.inrae.fr/en/bioinfo/xtandempipeline/down-load/<math>\mathbf{Z}$.

1.6.1 INSTALLATION ON MS WINDOWS AND MACOS SYSTEMS

The installation of the software is extremely easy on the MS-Windows and macOS platforms. In both cases, the installation programs are standard and require no explanation.

1.6.2 INSTALLATION ON DEBIAN- AND UBUNTU-BASED SYSTEMS

The installation on Debian- and Ubuntu-based GNU/Linux platforms is also extremely easy (even more than in the above situations). ; is indeed packaged and released in the official distribution repositories of these distributions and the only command to run to install it is:

\$ ^I sudo apt install <package_name> RETURN

In the command above, the typical *package_name* is in the form <u>i2masschroq</u> for the program package and <u>i2masschroq-doc</u> for the user manual package.

Once the package has been installed the program shows up in the *Science* menu. It can also be launched from the shell using the following command:

\$ i2masschroq RETURN

¹ The prompt character might be % in some shells, like *zsh*.

2 FUNDAMENTALS IN BOTTOM-UP PROTEOMICS

This chapter is an optional chapter which the reader might be referred to upon reading other part of this manual.

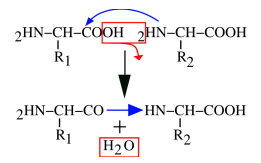
2.1 The Protein Biopolymer: Structure and Chemistry

This section introduces the basics in protein polymer chemistry. The way this topic is going to be covered is admittedly biased towards mass spectrometry and proteins. Moreover, the aim of this chapter is to provide the reader with the specialized words that will later be used to describe and explain the (inner) workings of the *i2Mass-ChroQ* program. This manual is not a "crash course" in biochemistry.

2.1.1 PROTEIN BIOSYNTHESIS

Proteins are made of amino acids. There are twenty major amino acids in nature, and each protein is made of a number of these amino acids. The combinations are infinite, providing enormous diversity to the protein realm. A protein is a polar polymer: it has a left end and a right end, and polymerization actually occurs from left to right (from N-terminus to C-terminus, see below). FIGURE 2.1, "PEPTIDIC BOND FORMATION BY CONDENSATION" shows that the chemical reaction at the basis of protein synthesis is a *condensation*. A protein is the result of the condensation of amino acids with each other in an orderly polar fashion. A protein has a left end, called *N-terminus; amino-terminal end* and a right end, called *C-terminus; carboxy-terminal end*. The left end is an amino group ($_2$ HN–) corresponding to the non-reacted α -amino group of the very first amino acid of the protein sequence. Upon condensation of a new entering amino acid onto the first N-terminal one, the amino group of the entering amino acid reacts (nucleophilic attack) with the α -carboxyl group of the N-terminal amino acid. A water molecule is released, and the formation of an amide bond between the two amino acids yields a dipeptide. The right end of the dipeptide is a carboxyl group (–COOH) corresponding to the un-reacted α -carboxyl group of the last amino acid to have been "polymerized in".

The bond formed by condensation of two amino acids is an amide bond, also called—in protein chemistry a *peptidic bond*. The elongation of the protein is a simple repetition of the condensation reaction shown in FIG-URE 2.I, "PEPTIDIC BOND FORMATION BY CONDENSATION", granted that the elongation *always* proceeds in the described direction (a new monomer arrives to the right end of the elongating polymer, and elongation is done from left to right).



The left end monomer R_1 is condensed to the right end monomer R_2 to yield a peptidic bond. A water molecule is lost during the process.

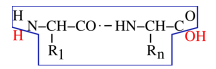
FIGURE 2.1: PEPTIDIC BOND FORMATION BY CONDENSATION



Note

Now we should point at a protein chemistry-specific terminology issue: we have seen that a protein is a polymer made of a number of monomers, called *amino acids*. In protein chemistry, there is a subtlety: once an amino acid has been polymerized into a protein, it is no more called an amino acid, but is called a *residue* instead. We may say that a residue is an amino acid less a water molecule.

From what we have seen until now, we may define a protein this way: — "A protein is a chain of residues linked together in an orderly polar fashion, with the residues being numbered starting from 1 and ending at n, from the first residue on the left end to the last one on the right end". This definition is still partly inexact, however. Indeed, from what is shown in FIGURE 2.2, "END CAPPING CHEMISTRY OF THE PROTEIN POLYMER", there is still a problem with the extremities of the residual chain: what about the amino group on the left end of a protein (the amino group sits right onto the first amino acid of the protein), and what about the carboxyl group of the right end of a protein (the carboxyl group sits right onto the last amino acid of the protein)? Because these groups lie at the extremities of the residual chain, they remained unreacted during the polymerization process. But because we are simulating a residual chain using residues and not amino-acids, we still need to put the protein polymer molecule in its "finished state": by *capping* the left end with a proton *cap* (so as to complete the carboxyl group). The capping of the residual chain extremities ensures that the polymer is in its finished state, and that it cannot be elongated anymore. The proton is the *left cap* of the protein polymer and the hydroxyl is the *right cap* of the protein polymer.



A protein is made of a chain of residues and of two caps. The left cap is the N-terminal proton and the right cap is the C-terminal hydroxyl. Altogether, the residual chain (enclosed here in the blue polygon) and both the H and OH red-colored caps do form a complete protein polymer in its finished state.

FIGURE 2.2: END CAPPING CHEMISTRY OF THE PROTEIN POLYMER

Now comes the question of unambiguously defining the structure of a protein. It is commonly accepted that the simple ordered sequence of each residue code in the protein, from left to right, constitutes an unambiguous description of the protein's primary structure (that is, its sequence). Of course, proteins have three-dimensional structures, but this is of no interest to a program like *massXpert*, which is aimed at calculating masses of polymers. To enunciate unambiguously the sequence of a protein, one would use a symbology like this:

- Using the 3-letter code of the amino acids: Ala Gly Trp Tyr Glu Gly Lys
- Using the 1-letter code of the amino acids:
 A G W Y E G K
 Alanine is thus the residue 1 and Lysine is the last residue (n = 7)

2.1.2 PROTEIN DISRUPTING CHEMISTRIES

The "polymer chain disrupting chemistry" was mentioned earlier as a complex subject that was of *enormous* importance to the mass spectrometrist. This is why that subject will be treated in a pretty thorough manner. First of all it should be noted that a chemical modification of a polymer does not necessarily involve the perturbation of the chain structure of the polymer. Here, however, we are concerned specifically with a number of chemical modifications that yield a polymer chain perturbation; *cleavages* and *fragmentations*:

Cleavages. These are chemical processes by which a cleaving agent will act directly on the protein residual chain making it fall into at least two separated pieces (the peptides).

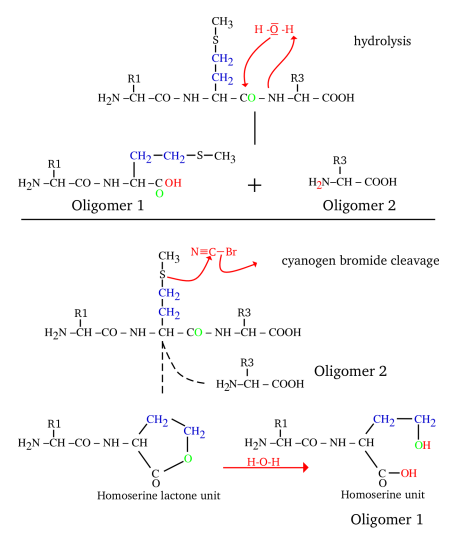
Fragmentations. These are chemical processes by which the polymer structure is disrupted into separated pieces (the *product ions*, or *fragments*) mainly because of energy-dependent electron doublet rearrangements leading to bond breakage.

2.1.2.1 PROTEIN CLEAVAGE

Upon cleavage of a protein, the cleaving molecule reacts with it, and by doing so directly or indirectly *"dissolves"* an inter-residue bond. A protein cleavage always occurs in such a way as to generate a set of *true* finished polymerization state "proteins" (smaller in size than the parent polymer, evidently, which is why they are called *oligopeptides*, or *peptides*). Indeed, let us take the example shown in FIGURE 2.3, "PROTEIN CLEAVAGE BY WATER AND CYANOGEN BROMIDE", where a tripeptide (a very little protein, containing a methionyl residue at position 2) is submitted either to a water-mediated cleavage (hydrolysis, upper panel) or to a cyanogen bromide-mediated cleavage (lower panel). The two cases presented in this figure are similar in some respects and different in others:

- In the first case the molecule that is responsible for the cleavage is water, while in the second case it is cyanogen bromide;
- In both cases the bond that is cleaved is the inter-monomer bond (in protein chemistry this is a peptidic bond);
- In both cases the Oligomer 2 has the same structure;
- The structures of the Oligomer 1 species differ, when produced using water or cyanogen bromide as the cleaving molecule.

The difference between hydrolysis and cyanogen bromide cleavage is in the generation of the Oligomer 1 species: the cyanogen bromide cleavage has a side effect of generating a homoseryl residue at the C-terminus of Oligomer 1, while hydrolysis generates a genuine methionyl residue. This is because water reverses in a very symmetrical manner what polymerization did (hydrolysis is the converse of condensation), while cyanogen bromide did some chemical modification onto the generated Oligomer 1 species.



A tripeptide is cleaved at position 1 either by hydrolysis (top) or by cyanogen bromide (bottom). Cyanogen bromide cleaves specifically on the right of a methionine monomer. Upon cleavage, the methionyl monomer gets converted into homoserine by the cyanogen bromide reagent

FIGURE 2.3: PROTEIN CLEAVAGE BY WATER AND CYANOGEN BROMIDE

Nonetheless, the reader might have noted that—interestingly—all the four oligomers do effectively have their left cap (the proton, making the N-terminal amino group) and their right cap (the hydroxyl, making the C-terminal carboxyl group). This means that in both water- and cyanogen bromide-mediated cleavages, all the generated oligomers are indeed true polymers in the sense that: 1) they are a chain of residues (modified or not) and 2) they are correctly capped (*i.e.* they are polymers in their finished polymerization state). This is important because it is the basis on which we shall make the difference between a cleavage process and a fragmentation process. Thus, our definition of a peptide might be: *a peptide is a* protein (*of at least one residue*) *in its finished polymerization state that was generated upon cleavage of a longer protein*. Of course, when we use the term "*protein*", above, we mean "protein polymer", irrespective of its size.

When the protein cleavage reaction precisely reverses the reaction that was performed for the same protein's biosynthesis, there is no special difficulty. But when the cleavage reaction modifies the substrate, then this should be carefully taken into account when using *i2MassChroQ*. This is true for any chemical modification that happens onto a protein.

Well, all this sounds reasonable. But what about the "normal" case, when the cleavage is done using water? Nothing special: the mass of the oligomer is calculated by summing the mass of each monomer in the oligomer (since the monomers are not modified, this is easily done) and the masses corresponding to the left and right caps (these are defined in the polymer chemistry definition; in our present case it would be a proton on the left end, and a hydroxyl on the right end). In this way, the oligomer complies with its definition, which states that it is a faithful polymer made of monomers and that it is in its finished state.

Yes, but then how should one calculate the mass of the modified oligomer, like our Oligomer 1 in the case of the cyanogen bromide-mediated cleavage? Simple enough: in a first step it does exactly the same way as for the unmodified oligomer. Next, each oligomer is checked for presence or absence of a methionine residue on its right end. If a methionine is found, the mass corresponding to the "-C1H2S1+O1" chemical reaction is applied. And that's it.

2.1.2.2 PROTEIN FRAGMENTATION

In a fragmentation process, the bond that is broken does not necessarily yield smaller-sized "proteins" because fragmentation does not necessarily break the inter-residue bond the same way that the hydrolysis does. Indeed, fragmentations are oft-times high energy chemical processes that can affect peptidic bonds at different locations, not necessarily between the CO-NH bond of the peptidic bond. This is one of the reasons why fragmentations do differ from cleavages.

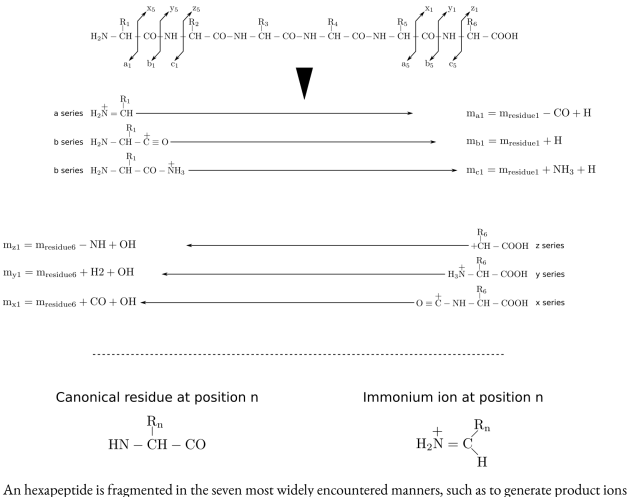
Another peculiarity of fragmentations, compared with cleavages, is the fact that there is no cleaving molecule starting the process, like water or cyanogen bromide, for example. Indeed, in the gas phase, the peptidic ions are "isolated": that is, very far one from each other. A fragmentation process is often initiated by an intra molecular electron doublet rearragement that propagates more or less in the polymer structure to eventually break it. Fragmentations are mainly a gas phase process, not some reaction that happens in solution as a result of putting in contact the polymer and some reagent. It is precisely because no cleaving molecule is involved in the fragmentation process that the obtained fragments are not necessarily capped like a normal polymer should be; and this is another really important difference between cleavage and fragmentation. The following examples should illustrate these concepts.



Tip

For the sake of completeness of this section, it must be noted that it is possible to have other "*chemi-cal/physical* entities" intervene during the gas phase fragmentation process by enacting a chemical reaction, be these entities ions, electrons or photons. In bottom-up proteomics, the intervening molecules are gas molecules (nitrogen, most often, or helium) that act as physical entities imposing collisions to the peptidic ions with the effect that the ions acquire internal energy, eventually leading to dissociation (CID, for "collisionally-activated dissociation").

There is a pretty important number of different kinds of fragments that can be generated upon fragmentation of peptides. We are going to detail the most common ones.



An hexapeptide is fragmented in the seven most widely encountered manners, such as to generate product ions of the a, b, c, x, y, z series and also immonium ions. The figure illustrates the position of the bond dissociation for each kind of fragment (exemplified using the case of the smallest fragment possible) and the mass calculation method is described for each fragment kind; consider that each fragment bears only *one positive* charge.

FIGURE 2.4: PROTEIN FRAGMENTATION PATTERNS MOST WIDELY ENCOUNTERED

As can be seen from FIGURE 2.4, "PROTEIN FRAGMENTATION PATTERNS MOST WIDELY ENCOUNTERED", the fragmentations do generate fragments of three categories: the ones that include the left end of the precursor polymer (a, b, c), the ones that include the right end of the precursor polymer (x, y, z), and finally the special case in which the fragment is an *internal fragment*, like the immonium ions. When looking at the fragmentations described in the figure, it becomes immediately clear why a fragmentation cannot be mistaken for a cleavage: the ionization of the fragment is not necessarily due to the captation of a proton by the fragment. Furthermore, we can also see that a fragmentation is not a cleavage because the fragment that is generated is *absolutely* not necessarily what we call a polymer, in the sense that the fragment might not be capped the same way as the precursor protein/peptide is (that is, the fragment is not in its finished polymerizaton state).

By looking at FIGURE 2.4, "PROTEIN FRAGMENTATION PATTERNS MOST WIDELY ENCOUNTERED", the reader should have noticed that the fragment naming scheme takes into consideration the fact that the fragment bears the N-terminal or C-terminal end of the precursor peptide (or none, also). Indeed, the numbering of fragments holding the N-terminal end of the precursor polymer sequence begins at the left end, and for fragments that hold the C-terminal end, at the right end. Thus the third fragment of series $a(a_3)$ would involve monomers $[I \rightarrow]$ and the third fragment of series $y(y_3)$ would involve monomers $[6 \rightarrow]$ (see arrows in the figure).

2.2 GENERAL OVERVIEW OF BOTTOM-UP PROTEOMICS

Bottom-up proteomics is a field of endeavour where the ultimate goal is to identify the greatest number of proteins in a given sample. This goal might also, depending on the project at hand, be doubled with another goal: characterize at the finest level possible the nature and the position of post-translational/chemical modifications beared by the proteins.

To achieve the best results, proteomics has developed over the years a number of methods and techniques that, taken together, have allowed scientists to obtain impressive results of protein identification on pretty complex samples. These are listed below:

- *Mass spectrometers:* The development of mass spectrometers of ever-greater resolution power has allowed to attain at ever-lower false discovery rates over the years. In particular, the development of the Orbitrap analyzers, along with the huge improvements of the time-of-flight (TOF) mass analyzer technology, have strongly increased the identification results reliability by allowing the downstream data processing step to be more stringent in the protein identification task (see below);
- *Chromatography:* The development of highly resolutive chromatography resins along with the elaboration of hardware (columns, chromatography setups) that yields sensitivity improvements have had their share in the way proteomics has evolved over the years;
- *Bioinformatics:* The development and refinement of software that can cope with extremely large data sets (think metaproteomics) is one major field that enabled significant advances in proteomics. Also, refinement of algorithms related to the simulation of isotopic clusters and comparison with experimental data have had their part. Likewise so for algorithms that detect the charge of ions based on the analysis of the isotopic cluster peaks. Being able to single out without error the monoisotopic peak of an isotopic cluster (whater the ion charge or m/z ratio) is a big part of the successfully tackled challenges at the root of successful proteomics data processing.

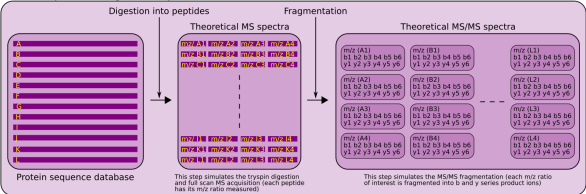
In this section, we will review the bioinformatics-based mass spectrometric data processing, as it is the core subject of this user manual. In particular, we will provide an outline of how the major software packages on the market perform protein identification on the basis of mass spectrometric analyses of biological samples.

This section will outline in not-so-rough terms how bottom-up proteomics works, from the protein sample to the protein identification list. The workflow comprises two sequential processes:

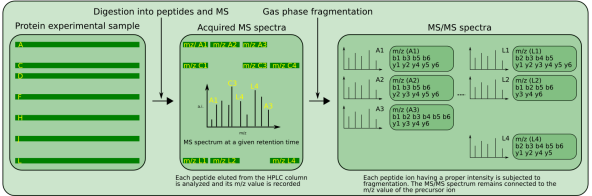
- *From the protein to the sequences of the peptides*: this initial part of the workflow is somehow doubled by having two parallel processes replicating it:
 - In silico process;
 - Experimental process.

These two processes are described in Figure 2.5, "Theoretical and experimental parallel data-producing processes".

In silico processing of the proteins



Experimental processing of the proteins



The digestion of the proteins, the analysis of the m/z of the peptides and the sequencing of the peptides are processes that exist both *in silico* and experimentally. This figure shows how the processes somehow mirror each other in the virtual and real contexts.

FIGURE 2.5: THEORETICAL AND EXPERIMENTAL PARALLEL DATA-PRODUCING PROCESSES

• Database searching using experimental data: this last part of the workflow is entirely based on bioinformatics software and involves the search for peptide *vs* mass spectrum matches and then a process called *protein inference* (see Section 2.2.5, "MATCHING FRAGMENTATION SPECTRA WITH THEORETICAL SPECTRA").

2.2.1 The First Step: Digestion of the Sample's Proteins

The very first step in the bottom-up protemics workflow is to digest all the proteins in the initial biological sample with a site-specific endoprotease: typically tryspin.

The sample is subjected to proteolysis with all its proteins unresolved. This produces a highly complex mixture of peptides, each having a constant characteristic: each peptide has one predictable end (unless it is either the protein's N-terminal or the C-terminal peptide, as detailed below), either N-terminal or C-terminal:

- *Predictable N-terminus:* when the protease cuts at the N-terminal end of the target residue. For example, EndAspN cleaves left of Asp residues, thus producing peptides that always have Asp as their N-terminal residue. The only exception is when the peptide is the protein's N-terminal peptide and the first residue is not Asp);
- *Predictable C-terminus:* when the protease cuts at the C-terminal end of the target residue. For example, the most used enzyme, trypsin, cuts right of the basic residues Lys and Arg. The generated peptides thus necessarily end with one of these two residues. The only exception is when the peptide is the protein's C-terminal peptide and the last residue is not Lys nor Arg.



Tip

One interesting feature of trypsinolyis is that it generates peptides that—for their major part—will most probably be protonated twice: on their N-terminal end (the primary NH_2 amine group ¹ and on the basic residual chain of the basic residue found at their C-terminal position (the ε -amine group for Lys and the guanidium group for Arg). Upon fragmentation of the peptide's precursor ion, both the left hand side fragment and the right hand side fragment will bear a proton and will thus be detected, thus potentially providing a better coverage of the peptide's sequence during the MS/MS experiment.

2.2.2 Chromatographic Separation of the Peptidic Mixture

One major analytic step in bottom-up proteomics is the separation of the peptides obtained by endoproteolysis of all the proteins in the sample. Indeed, analyzing all the peptides in one single injection without any prior chromatographic separation would yield catastrophic results, similar to having injected nothing in the mass spectrometer.

The typical method for resolving peptides is by separating them on a chromatographic column functionalized with a hydrophobic group (for peptides, that would be a C₁₈ reversed phase column).

The chromatographic gradient that will elute the peptides progressively according to their increasing hydrophobicity will be developed over the 5–95 % of acetonitrile (a non-protic organic solvent).



Tip

Using acetonitrile as the non-protic organic solvent has the huge benefit of not injecting protons inside the mass spectrometer as the chromatographic gradient develops.

¹ If not either converted to an amide group by acetylation or formylation or cyclised.

The eluate of the chromatographic column is directly injected into the mass spectrometer's source. The role of the mass spectrometer's source device is to ensure that the analytes are desolvated and ionized upon their entering in the core part of the mass spectrometer. Most often, that source is an electrospray source that is fed a liquid (typically, the eluate from the column). The source is designed to evaporate the solvent (analyte desolvation) and —having an electric potential applied to it— to help ionize the analytes (often the peptides are already ionized in solution, prior to desolvation). The electrically charged analytes in the gas phase are thus ions, the m/z (mass-to-charge) ratio of which can be measured by the mass spectrometer analyzer.



WARNING

There are two main sources used in the mass-spectrometry-for-biology specialty: the matrix-assisted laser desorption ionization (MALDI) source and the electrospray ionization (ESI) source. One important difference between the two is that the MALDI process mostly produces mono-charged ions ($[M+H]^+$), while the ESI process mostly produces multi-charged ions ($[M+nH]^{n+}$). This has huge implications in the mass data analysis.

The source that is mainly used in bottom-up proteomics is the ESI source.

2.2.3 Mass Spectrometric Analysis of the Peptides

Upon elution off the chromatographic column, the peptides are desolvated, ionized and drawn into the mass spectrometer using an electrical field. Once they have entered the mass spectrometer they are analyzed in the mass analyzer of the instrument.



Note

There are a variety of mass analyzers commonly used in bottom-up proteomics. In fact, one single instrument might have as many as 4 or 5 mass analyzers. However, not all the analyzers in the instrument are responsible for the m/z measurement.

Sometimes, during the whole cycle of the analysis, two different mass analyzers are used at different steps of the cycle: one analyzer selects the ion for fragmentation and another analyzer measures the m/z value of the fragments.

In bottom-up proteomics, two different kinds of mass spectrometric data are required—ideally, for each peptide eluted from the column— in order to effectively identify the proteins in the initial sample:

- The mass-to-charge ratio value (m/z) of the peptide ion;
- The m/z values of the fragments (the product ions) of the peptidic precursor ion that has undergone an MS/MS gas phase fragmentation².

These two kinds of data are necessary because the protein identification process is based on searches in protein databases using the precursion ions' m/z value and the m/z values of that ion's fragments when it is fragmented. The way the protein databases are used as the substrate of these searches is described in the next section.

2.2.4 The Protein Databases and Their Use

The previous section ended on the idea that the protein identification process, that is based on the analysis of all the peptides of a peptidic mixture resulting from the endoproteolysis of a sample containing many proteins, requires searches into protein databases.

A bottom-up proteomics experiment typically needs at least one protein database: a database listing all the known proteins of the organism from which the initial sample of proteins was prepared. That organism might be a bacterium, a Eucaryote, like a fungus, a protist, a plant, a mammalian... Optional databases might be used, like protein databases listing all known protein contaminants, for example.

The protein databases are files in the following FASTA format:

>GRMZM2G009506_P01 NP_001149383 serine/threonine-protein kinase receptor MEEQHMAGPPYRYRLQHRRLMDIAPASASDDDSGHHGSNGMAIMVSILVVVIVCTLFYCV YCWRWRKRNAVRRAQIERLRPMSSSDLPLMDLSSIHEATNSFSKENKLGEGGFGPVYRGV MGGGAEIAVKRLSARSRQGAAEFRNEVELIAKLQHRNLVRLLGCCVERDEKMLVYEYLPN RSLDSFLFDSRKSGQLDWKTRQSIVLGIARGMLYLHEDSCLKVIHRDLKASNVLLDNRMN PKISDFGMAKIFEEEGNEPNTGPVVGTYGYMAPEYAMEGVFSVKSDVFSFGVLVLEILSG QRNGSMYLQEHQHTLIQDAWKLWNEDRAAEFMDAALAGSYPRDEAWRCFHVGLLCVQESP DLRPTMSSVVLMLISDQTAQQMPAPAQPPLFASSRLGRKASASDLSLAMKTETTKTQSVN EVSISMMEPRFWADPGTSNGAATSHPATGACKKRGGQGGDRNVKDGLAARTPTHQPVARW HHDRRIVD

This format is really simple, because it only contains three information pieces, grouped in as many stanzas as there are proteins in the database:

² Most often, that fragmentation step is performed using collisionally-activated dissociation (CID). In this process, the peptidic precursor ion is first isolated in the gas phase on the basis of its m/z value and then is accelerated against a gas "fog" inside of the collision cell of the instrument. The ion hits gas molecules multiple times, acquires a lot of energy and finally breaks.

- The *unique* protein's accession id in the database (<u>GRMZM2G009506_P01</u>) that comes right after the '>' prompt that signals a new protein stanza;
- The protein description (<u>NP_001149383</u> serine/threonine-protein kinase receptor) that provides some functional data bits for the protein at hand;
- The protein sequence (the rest of the stanza above).

The first (id) and second (description) information bits are used in various places in the *i2MassChroQ* program. The protein databases are used by the protein identification software as the very first step in a bottom-up proteomics data analysis process: the proteins in the database are digested *in silico* in order to produce a list of peptides that retain a connection to the protein from which they were generated. For each one of all these peptides, the following data bits are computed (FIGURE 2.5, "THEORETICAL AND EXPERIMENTAL PARALLEL DATA-PRODUC-ING PROCESSES", top panel):

- *sequence:* The peptide's sequence;
- m/z value: The peptide's m/z value, often computed for the mono-protonated ($[M+H]^+$) ion;
- *MS/MS spectrum:* The peptide's fragmentation spectrum is nothing but an array of m/z values corresponding to the set of calculated fragments (of the b and y ion series). The m/z values of the product ions are crucial for the database search algorithm;

The next step is the establishment of a relation between the experimental MS/MS data acquired by the instrument and the theoretical MS/MS spectra computed from the protein sequences in the database. This next step is described in detail in the next sections.

2.2.5 MATCHING FRAGMENTATION SPECTRA WITH THEORETICAL SPECTRA

This section is about how the protein database searching software sets a relation between the experimental mass data and the theoretical mass data originating in the protein database. The elementary relation is between a given *experimental* MS/MS mass spectrum of a peptide's ion at a given m/z value and its *theoretical* counterpart from the database: when these two MS/MS spectra match at a sufficiently convincing level, then a *"peptide vs mass spectrum match"* was achieved (abbreviated name: PSM). The computing of a PSM is described in detail in FIGURE 2.6, "THE STEPS LEADING TO A SCORED PEPTIDE VS MASS SPECTRUM MATCH (PSM)".

We have seen in SECTION 2.2.4, "THE PROTEIN DATABASES AND THEIR USE", that two somehow similar processes are at the basis of the preparation of the data for the subsequent database searches. These processes were described in FIGURE 2.5, "THEORETICAL AND EXPERIMENTAL PARALLEL DATA-PRODUCING PROCESSES".

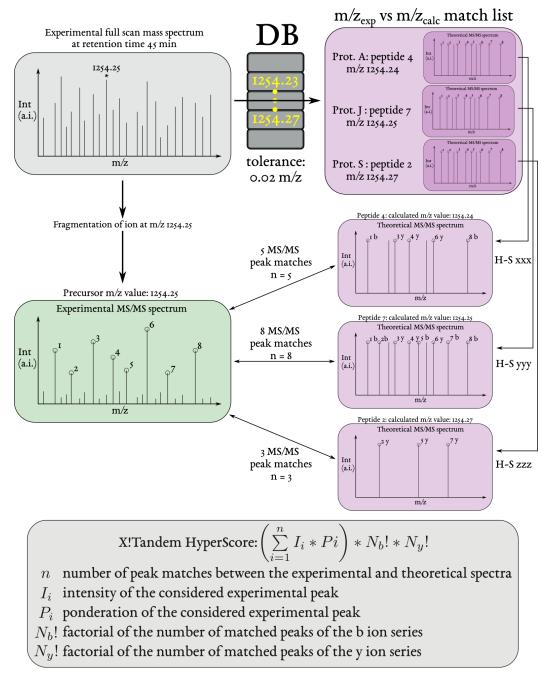
On the one hand (top panel, violet), the protein database is processed to digest *in silico* every protein it contains into a list of peptides. For each peptide arising from the digestion of a protein, the following data elements are recorded:

- The peptide's m/z value is computed. The association between that m/z value, the peptide and its originating protein is maintained;
- The peptide is fragmented into a list of peptidic fragments (product ions' m/z values, that is, the MS/ MS spectrum; typically b and y ions series). The connection with the earlier data elements above is also maintainted.

It is thus easy to determine the filiation between any given MS/MS theoretical mass spectrum, the precursor ion's m/z value, the peptidic sequence and, finally, the protein whence that peptide came.

On the other hand (bottom panel, green), the mass spectrometric data acquisition yields a huge set of the following pairs of data elements that are recorded over time:

- The m/z value of the peptidic precursor ion undergoing fragmentation (keeping a connection with the retention time at which it is recorded);
- The list of peptidic fragments (product ions' m/z values, that is, the MS/MS spectrum). The connection with the precursor ions' m/z value and with the retention time is maintained.



H-S yyy >> H-S xxx >> H-S zzz

The process starts with a full scan mass spectrum from which the mass spectrometer selects one precursor ion at a definite m/z value. That ion is fragmented and thus generates a MS/MS spectrum. During the data exploration, the software extracts from the database all the peptides having the same m/z value as that of the fragmented ion (top right, violet background). Next, the experimental MS/MS spectrum is compared in turn to each one of the MS/MS spectra of the extracted peptide list. A HyperScore is computed at each comparison. Because *izMassChroQ* uses *X!Tandem* as its preferred protein database search engine, the HyperScore calculation, as performed by *X!Tandem*, is described.

FIGURE 2.6: THE STEPS LEADING TO A SCORED PEPTIDE VS MASS SPECTRUM MATCH (PSM)

Once the acquisition of the experimental data is complete, the analysis of these data involves going through all the fragmentation data of the acquisition and performing these steps for *each* MS/MS spectrum (as evidenced in Figure 2.6, "The steps leading to a scored peptide vs mass spectrum match (PSM)":

- Get the precursor ion's m/z value;
- Compute the match m/z range. For example, if the software is configured with a m/z tolerance for the m/z matches set to 0.02 and the precursor ion's m/z value is 1254.25, then the match m/z range would be [1254.23-1254.27];
- Construct a list of all the peptides in the database that have their m/z value contained in the match m/z range;
- For *each* peptide in the list returned from the database, compare its theoretical MS/MS spectrum with the experimental one. Compute a HyperScore for comparison.

2.2.5.1 Computation of the PSM HyperScore

Of course, it is extremely rare that an experimental MS/MS spectrum matches fragment-by-fragment an identical theoretical spectrum. Most often, some theoretical product ions (MS/MS spectrum peaks) are missing from the experimental fragmentation spectrum. Also, there will almost certainly be dozens (if not hundreds) of peptides having a m/z value in the searched m/z range. Most certainly, the vast majority of these peptides are not of the right sequence (that is, do not have their MS/MS theoretical mass spectrum matching the experimental one). To make without any human scrutiny of the matches, it is necessary to compute a score that somehow assesses the extent to which both the experimental and theoretical MS/MS spectra match. That score, in *X*!*Tandem*, is called *HyperScore* and is described at the bottom of the figure.

The HyperScore computation process is relatively straightforward. First off, it is necessary to stress the fact that a HyperScore is computed each time an *experimental* MS/MS spectrum is compared to a theoretical (*calculated*) MS/MS spectrum (see m/z_{exp} vs m/z_{calc} match list in Figure 2.6, "The steps leading to a scored peptide vs MASS SPECTRUM MATCH (PSM)").

In the example, three peptides from the database have their m/z value matching the searched m/z range (the m/z value of the precursor ion with accounting for the tolerance). So, the program checks the similarity between the experimental MS/MS spectrum and each one of the three theoretical ones. Each similarity test is associated to a HyperScore value.

The HyperScore is computed by summing—for each tested fragment peak *in the theoretical MS/MS spectrum* the product of two variables described below. Once that sum is computed, it is compounded by two factorial numbers also described below:

- Ii: the intensity of the matching mass peak in the experimental MS/MS spectrum (if found);
- P_i: the ponderation factor of the matching mass peak in the experimental MS/MS spectrum. That variable can take a number of values, depending on the presence or not of this fragment peak in the experimental MS/MS spectrum (if not found, then P_i is naught and the peak is disregarded entirely). There are other values greater than naught, accounting for the physico-chemical properties of the peptidic bond that was cleaved to obtain that fragment (presence of proline will lower the P value, for example). Intuitively, the HyperScore will end up larger if there are a lot of fragment peaks in the theoretical MS/MS spectrum that are matched with experimental ones (each P_i value compounded by the I_i value is being summed into the HyperScore final value).
- N_b!: the sum computed above is then compounded by the factorial of the number of ions of the b ion series that are found in the experimental MS/MS spectrum;
- N_y!: the product computed at the previous step is then compounded by factorial of the number of ions of the y ion series that are found in the experimental MS/MS spectrum. This last compounding operation terminates the computation of the HyperScore value.

It is apparent now that the HyperScore value will tend to be greater if there are numerous fragment peaks in the theoretical MS/MS spectrum that are matched by fragment peaks in the experimental MS/MS spectrum. Also, the score value is incremented if the intensity of the matching peaks is greater and if the number of matching peaks of the two b and y ions series is greater.

This, however, cannot be all of it, because the HyperScore does not really answers the question: "*what are—if* any—, of all the PSMs found for a given experimental MS/MS spectrum, the one (or ones) that we can faithfully tell as true match(es)?". To answer that question, some more computational steps need to be carried over, that should lead to a numerical value that is truly indicative of the confidence we may have that a given PSM is a real match. In X!Tandem, that numerical value is called expectation value (abbreviation: E-value). We describe the whole process of its computation below.

2.2.5.2 Computation of the Peptide Expectation Value (E-value)

First of all, it needs stating that we describe the *peptide E-value*, not the protein E-value. A peptide E-value is obtained for a single experimental MS/MS spectrum. It is computed by looking into the HyperScore values obtained for all the MS/MS spectra comparisons described at the previous section. The HyperScore values (for example, the three values denoted *H-S xxx*, *H-S yyy* and *H-S zzz* in FIGURE 2.6, "The steps leading to a scored peptide vs mass spectrum match (PSM)") are used to perform the E-value computation. In the following text, we'll assume that there are many more PSMs than these three, for a given experimental MS/MS spectrum (which is actually the reality, with hundreds of peptides in the database that match a given searched m/z range). As illustrated in FIGURE 2.7, "COMPUTATION OF A PEPTIDIC EXPECTATION VALUE (E-VALUE)", a histogram is crafted

plotting the count of MS/MS spectral pair comparisons (let us call them "wannabe PSMs") against a number of HyperScore bins. This histogram is a good representation of the distribution of the HyperScore values among the various peptides in the m/z value-matching list (see previous section). In this example, the very best HyperScore value is 82 and the number of PSMs having that score is obviously very low! Instead, the distribution clearly shows that there are a vast majority of wannabe PSMs that have very low HyperScore values and that will not ultimately be considered as real PSMs.

In order to be able to use the distribution pattern further, the second half of the distribution's main peak is replotted by computing the natural logarithm of the count of MS/MS spectral pair comparisons, still against the HyperScore value bins. The new plot is easily fitted into a line, of which the equation is computed.

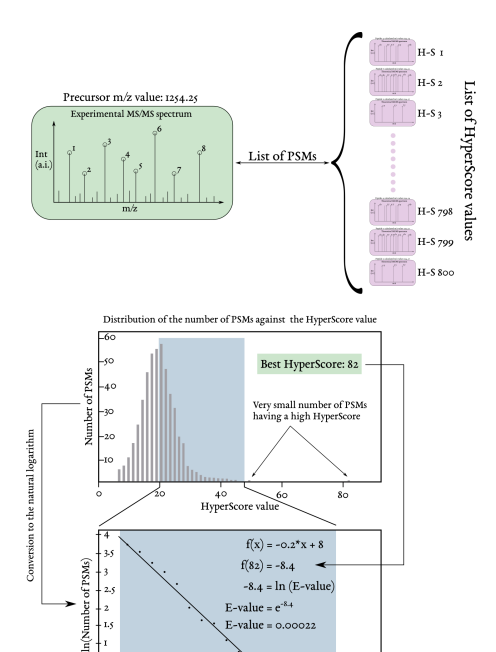
The best HyperScore value (82, in the example) is then used in the line equation to compute the corresponding ordinate (the natural logarithm of the PSMs count having that HyperScore). That value (-8.4, in the example) corresponds to the natural logarithm of the expectation value (E-value). By using the exponential function, the E-value is thus computed to be 0.00022, which a pretty low number. Since the E-value somehow gives an idea that a given PSM was obtained by chance, the very small obtained result shows that the match almost certainly was a faithful one.



Note

The expectation value is defined as the probability that the peptide sequence would match an experimental tandem mass spectrum by chance, if the trial is repeated many times. For example, if the E-value is found to be 1, then that means that the match can occur by chance or not with an equal probability. Instead, if the E-value is found to be 0.01, then that means that there is one event over 100 trials that the match has occurred by chance.

The smaller the E-value, the more confidence one has that the match is correct and that the PSM is a faithful one.



For each experimental MS/MS spectrum, gather all the peptides in the database that have a m/z value matching the precursor ion's m/z value. For each peptide sequence, compute the HyperScore. With all the HyperScore values, go on with the calculation of the expectation value for the peptide set. The peptidic E-value should be the smallest possible, as it is an indication of the possibility that the match between the experimental MS/MS spectrum and the theoretical mass spectrum occurred by chance.

50

 $-8.4 = \ln (\text{E-value})$

E-value = $e^{-8.4}$

E-value = 0.00022

40

45

FIGURE 2.7: COMPUTATION OF A PEPTIDIC EXPECTATION VALUE (E-VALUE)

2.5

2

1.5

0.5

20

25

30

35

HyperScore value



Tip

The user configures the software to only consider PSMs if their peptidic E-value is below a given threshold. Typically, that threshold is given a value of 0.05 (FIGURE 3.6, "CONFIGURATION OF THE LOADING OF THE IDENTIFICATION RESULTS").

When a reliable match between an experimental MS/MS spectrum and a theoretical MS/MS spectrum is found (that is, a true PSM), the software reports the following set of data elements:

- *m/z:* the m/z value of the precursor peptidic ion that underwent fragmentation;
- *sequence:* the sequence of the peptide that was matched in the present PSM;
- *protein name:* the protein accession number that produced the matched peptide upon enzymatic digestion of the sample;
- *E-value:* the peptide expectation value, as described above.

2.2.5.3 Computation of the Protein Expectation Value (E-value)

The last step in the computation of values that help the software and the user determine if identifications are faithful (for peptides and for proteins) is the computation of the protein expected value. This value is very easily computed: it is the product of the E-values of all the peptides that participated in the identification of the protein. By necessity, then, the protein E-value will be less than the threshold peptide E-value (since that last value is below 1). By default, the protein E-value is set to 0.01 (FIGURE 3.6, "CONFIGURATION OF THE LOADING OF THE

IDENTIFICATION RESULTS").

2.2.5.4 PROTEIN INFERENCE: FROM PSMs to PROTEIN IDENTITIES

One remaining critical question is: "— *How is the list of protein identifications returned by the database searching software verified and modified?*" Indeed, there are a number of situation where the proteomics data user may want to tweak the identification results. But also, the protein identification list returned by the database software may not be as perfect as one would expect. Bioinformaticians working in proteomics have come up with a number of algorithms to better the reliability of the identification results returned by database searching software.

In *i2MassChroQ* we use an algorithm that is impinged on the concept of *parcimony*. That algorithm is detailed in an article describing *X!TandemPipeline-Java* that was published in *The Journal of Proteome Research* in 2017 by Olivier Langella and Colleagues. The general concepts are presented here for the sake of completeness of this user manual.

Spectrum Peptide number number	Identified proteins	ldentified proteins with identifying peptides	Sub-groups of proteins	Groups of proteins
Ma → 1 Ma → 2 Ma → 3	A B C A B	A 1 2 3 4 6	Sub-group 1 A 1, 2, 3, 4	Group 1 Sub-group 1 A 1 2 3 4
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	B	B 1 2 3 5 7	Sub-group 2 Sub-group 3 C.1. Sub-group 3 C.1. Sub-group 3 C.1. Sub-group 3 C.1. Sub-group 3 C.1. Sub-group 3 C.1. Sub-group 3 Sub-group 3 Sub-grou	Sub-group 2 Sub-group 3 Sub-group 3 Sub-group 3 Sub-group 3 Sub-group 3 Sub-group 3 Sub-group 4 Sub-group 4 Sub-group 5 Sub-group 5 Sub-group 5 Sub-group 7 Sub-group 7 Sub-g
$\begin{array}{cccccccccccccccccccccccccccccccccccc$			Sub-group 4	Group 2 (Sub-group 4
Ma < ∰ Ma → ™ Ma → ™		D 8 9a 10 11 12 E 8 9b 10 11 12	b 93 10 11 12	B 9a 10 11 12 E 8 9b 10 11 12
$M^2 \rightarrow \mathbb{I}$		F 10 11 12		

The process of establishing a consolidated protein identity list from the results reported by the database searching software is illustrated (see text).

FIGURE 2.8: PROTEIN INFERENCE: CONSTRUCTING A CONSOLIDATED PROTEIN IDENTIFICATIONS LIST

The protein inference process, depicted in FIGURE 2.8, "PROTEIN INFERENCE: CONSTRUCTING A CONSOLIDATED PROTEIN IDENTIFICATIONS LIST", is a multi-step one. The starting point is the huge list of PSMs that are reported by the database searching software. These PSMs are displayed in the figure as the two columns on the left hand side: one *experimental* MS/MS spectrum (*Spectrum number*) has provided a convincing PSM and thus allowed the identification of a peptide (MS/MS I \rightarrow Pep I, *Peptide number*). Of course, a given peptide (Pep I) might have allowed the identification of multiple proteins (for example, homologous proteins that share the same peptidic sequence). Thus, Pep I is found in proteins A, B and C (column *Identified proteins*). The structure of the identified proteins can thus be partially reconstructed, and that is shown in column *Identified proteins with identifying peptides*. All the other PSMs are listed below that first one.

The general concept of the algorithm is that, by going through all the PSM data it is possible to check if some form of degraded redundancy allows pruning off some proteins from the list. This pruning off of some proteins is meant to increase the confidence that the identifications are reliable. That might be at the cost of having a smaller number of identified proteins, but with an improved false discovery rate (that is, a reduced FDR). As described below, the pruning off of proteins from the protien identifications list occurs at two different steps in the inference process.



Note

The FDR is commonly computed as the ratio between the number of PSMs matching the decoy database over the number of PSMs matching the target database: FDR = (#decoy / #target).

The first step is the creation of sub-groups of the identified proteins. In this step, all the proteins that could be identified thanks to the exactly same set of peptides are gathered into a sub-group. In the example, the subgroup that contains more than one protein happens to be sub-group 4. Note how protein F in this sub-group is identified by a set of three peptides. This is two peptides less than the number of peptides that identified the other two proteins (D and E) in the sub-group. The principle of parcimony allows thus to remove Protein F as that protein is not justified *per se*, that is, it is unnecessary to explain the presence of the three peptides.

The second step is the creation of groups that gather all the sub-groups that share at least one peptide. Thus, group 1 contains sub-groups 1, 2 and 3, while group 2 contains the sub-group 4. According to exactly the same philosophy as for the previous step, the sub-groups that contain proteins identified only by peptides also shared by proteins present in other sub-groups are pruned off.

The whole process described here is dubbed "*protein grouping*" in the *iaMassChroQ* language. The ouptput of this protein grouping process is displayed in the protein identification window, to be described below.

2.2.6 Phospho-Proteomics

In this section, the typical procedures involved in phospho-proteomics projects are described, from the sample handling to the post-translational modification data exploration.

2.2.6.1 HANDLING PHOSPHO-PROTEOMICS SAMPLES

iaMassChroQ is able to cope with phospho-peptides. The mass spectrometric data are acquired exactly as usual with the mass spectrometer, but the sample preparation goes along theses steps:

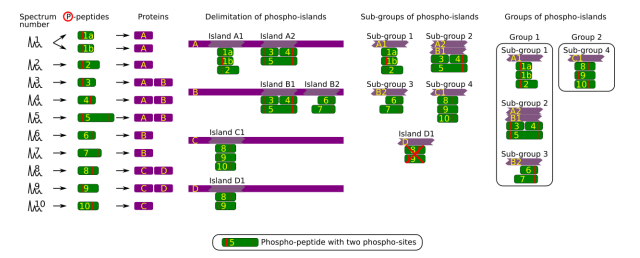
- Separate digestion of the samples (when there are more than one);
- Labeling of the peptides, each sample gets a different label;
- Pool of the whole set of peptides into a single mixture;
- Separation of the peptides on a strong cation exchange (SCX) resin, collection of the fractions;
- Phospho-peptide enrichment using IMAC³ for each SCX fraction. The SCX fraction is loaded onto the IMAC resin and, following a wash step, the phospho-peptides are eluted (pH-based elution). There is thus a one-to-one relation between a SCX fraction and an IMAC-based purification fraction.
- Mass spectrometric analysis of each IMAC-based phospho-peptide-enriched fraction.

X!Tandem needs to be configured in such a manner that it can generate all the theoretical peptides (and fragments) that might bear the phosphoryl group. This process is described in the section below.

³ Immobilized-metal affinity chromatography.

2.2.6.2 PROTEIN IDENTIFICATION IN PHOSPHO-PROTEOMICS PROJECTS

An analogous algorithm as the one used for protein inference is at play when *i2MassChroQ* is handling phospho-proteomics data. That algorithm is described below and in FIGURE 2.9, "PHOSPHO-SITE INFERENCE: CON-STRUCTING A CONSOLIDATED PHOSPHO-SITE LIST".



The process of establishing a consolidated phospho-site list from the results reported by the database searching software is illustrated (see text).

FIGURE 2.9: PHOSPHO-SITE INFERENCE: CONSTRUCTING A CONSOLIDATED PHOSPHO-SITE LIST

The phospho-island inference process, depicted in FIGURE 2.9, "PHOSPHO-SITE INFERENCE: CONSTRUCTING A CONSOLIDATED PHOSPHO-SITE LIST", is a multi-step one, most similarly to what was described in SECTION 2.2.5.4, "PROTEIN INFERENCE: FROM PSMS TO PROTEIN IDENTITIES". The starting point is the list of peptides that were identified and determined to bear one or more phospho-sites (thus called phospho-peptides; see the red vertical bar in the figure). Two difficulties here are, on the one hand, the fact that phospho-sites may be shared by more than one peptide and, on the other hand, the fact that more than one phospho-site might be determined on the *same* peptide. These are the reasons that the concept of *phospho-island* was elaborated: it is a protein region that bears at least one phospho-site, in turn beared by one or several overlapping phospho-peptides. It is important to note that the position and number of phospho-sites are not necessarily the same in all of the overlapping phospho-peptides.

In this inference process, the analogy with the previously described one is the following:

- Peptides are replaced by phospho-peptides;
- Proteins are replaced by phospho-islands.

In the first step, the phospho-islands are delimited on the phosphorylated proteins. In the second step, subgroups of phospho-islands are created using all the phospho-islands identified in different proteins and that share exactly the same set of phospho-peptides. At this step, any remaining phospho-island defined by a subset of phospho-peptides only partially defining a sub-group is disregarded. In the example, phospho-island D1 is defined by two phospho-petides, 8 and 9, that also are part of a sub-group defined by these two peptides but also by phospho-peptide 10. Phospho-island D1 is thus disregarded.

In the third step, all the sub-groups that contain phospho-islands beared by the same protein are gathered in a group.

3 The main program window

Proteomics data explorations, with *i2MassChroQ*, entail, for a large part, the following steps:

- Configuration of the *X!Tandem* external software that runs the database searches (producing peptide vs mass spectrum matches—PSMs—, leading to the peptide identifications and ultimately to protein identifications);
- Configuration of the protein database files (both the organism-specific protein databases and optional contaminant-containing databases);
- Loading of the mass spectrometry data acquisition files (the mzML format is recommended);
- Running X!Tandem from inside of i2 MassChroQ;
- Loading of the identification results produced during the previous step;

Note

i2MassChroQ can also handle peptide vs spectrum matches data (peptide identification data) from other software with the following formats:

- mzIdentML;
- pepXML;
- Mascot DAT files
- Relentless scrutiny of the peptide identification results. Optional modification of the results;
- Protein inference, that is, protein identification on the basis of the peptide identifications. *i2MassChroQ* implements a protein grouping algorithm, as described in FIGURE 2.8, "PROTEIN INFERENCE: CONSTRUCT-ING A CONSOLIDATED PROTEIN IDENTIFICATIONS LIST", that leads to consolidated protein identifications. The program has an interface geared towards the tweaking of the protein grouping process so as to let the user in full control of the stringency with which the protein identifications list is ultimately generated.

In this chapter, *i2MassChroQ*'s main window's user interface is described in detail, in particular in the way it is a starting point for the main tasks briefly mentioned above.

3.1 Starting a new *i2MassChroQ* working session

To start a session, run *i2MassChroQ* and the main program windows shows up as described in FIGURE 3.1, "MAIN **PROGRAM WINDOW**".

				X	Tan	der	nP	ipe	elin	e O	.4.3	D			~	,	^	0
F <u>i</u> le	<u>E</u> dit	0																
				Rur	ז X!T	Tand	den	n id	den	tific	atio	ns						
	Lo	ad ider	ntifica	ation	res	ults	(m	nzIo	den	tML	., pe	рХN	1L, X	!Tar	nder	m)		
			L	oad	an X	(!Tar	nde	eml	Pip	elin	e pr	ojec	t					

The main program window contains three buttons described in detail in the text.

FIGURE 3.1: MAIN PROGRAM WINDOW

The main program window contains three buttons that start the following main tasks:

- Run X!Tandem identifications. See Section 3.2, "RUNNING X!TANDEM IDENTIFICATIONS".
- Load identification results (mzIdentML, pepXML, Mascot, X!Tandem). See Section 3.4, "Loading the Protein Identification Results".
- Load an i2 MassChroQ project. See Section 3.5, "Loading 12 MassChroQ projects".

3.2 RUNNING X! Tandem IDENTIFICATIONS

To run *X!Tandem*-based identifications, click onto the *Run X!Tandem identifications* button. This triggers the opening of the window pictured in FIGURE 3.2, "X!TANDEM-BASED IDENTIFICATION CONFIGURATION".

*	XI	landem run setting	s	? ~ ^
	✓ Confid	gure the <u>X</u> !Tandem	execution	
		_		
Program path:	/usr/bin/tandem			Change 🗖
	X!Tan	dem Alanine 2017.	2.1.4	
	🖌 <u>R</u> un X	(Tandem through H	HTCondor	
X!Tandem mei	nory requirements (N	MB) 5000		
		Choose presets		
Qexactive_pro	eomics_analysis_FDF	<pre>.nosemi</pre>	~	🖉 Edit
	Cł	noose database file	25	
	***	•	Clear list	+ Add files
	Choose	MS data files to p	rocess	
/home/rusconi /home/rusconi /home/rusconi /home/rusconi	'devel/xtpcpp/doc-da 'devel/xtpcpp/doc-da 'devel/xtpcpp/doc-da 'devel/xtpcpp/doc-da 'devel/xtpcpp/doc-da	ta/proteomics/201 ta/proteomics/201 ta/proteomics/201	20906_proteomics 20906_proteomics 20906_proteomics	_extract_1_A05_u _extract_1_A09_u _extract_1_B01_u _extract_1_B05_u
<	Clear list	Add Bruker	timsTOF folders	+ Add files
		Output directory		
/home/rusconi/	level/xtpcpp/doc-data		ut	Choose

The configuration of a X!Tandem run is performed in this configuration window (see text for details).

FIGURE 3.2: X!TANDEM-BASED IDENTIFICATION CONFIGURATION

The configuration of an *X*!*Tandem* run entails defining the following:

- *Configure the X!Tandem execution*: This setting allows one to specify the path to the *X!Tandem* software program. The version of the program, if found, is displayed below (in this case, *Alanine 2017.2.1.4*). This feature is useful when the user wants to test multiple versions of the *X!Tandem* software.
- *Run X!Tandem through HTCondor*: Only check the box if running *X!Tandem* over the network on a server supporting *HTCondor*¹.
- Choose presets: This setting defines the parameters that X!Tandem must use. Either load already known presets from the drop-down list widget or edit them (or create a new set) by clicking onto the *Edit* button. Note that to load an existing presets file, it might be necessary to point *i2MassChroQ* to the directory that contains the presets file. Use the folder icon for this, as visible in FIGURE 3.3, "X!TANDEM PRESETS CONFIGURATION WINDOW (SPECTRUM TAB)".
- *Choose database files*: Add protein database files in the FASTA format. There must be at least one protein database that contains all the known proteins for the organism of interest (there might be as many such database files as necessary) and optionally protein databases containing known contaminant proteins (there might be as many such database files as necessary). Click onto the *Clear list* button to clear the database files list and start anew if an error occurred (it is not possible to remove files one at a time).
- *Choose MS data files to process*: Add the mass spectrometry data files (mzML or mzXML format) to be processed by the *X!Tandem* software. As many files as necessary might be added in the list.



Tip

When using Bruker timsTOF data, click onto the *Add Bruker timsTOF folders* button to select folders containing this kind of data. Bruker timsTOF data come as two files that must sit in the same directory.

- *Output directory*: This setting specifies the directory into which new files output by the *X*!*Tandem* process need to be created. *X*!*Tandem* produces identification results in files in an XML format that *i2MassChroQ* reads during a later step.
- *Number of threads*: This setting defines the maximum number of execution threads that *X*!*Tandem* might be using during its run.



Tip

Although *i2MassChroQ* sets that number of execution threads to 1, it is beneficial to set that number to the highest value possible.

I See HTTPS://RESEARCH.CS.WISC.EDU/HTCONDOR/ ₽.

3.3 SETTING THE X! Tandem RUN PRESETS

The *Edit* button of the *Choose presets* group box described above triggers the opening of a dialog window where the user might configure in the most detailed way the *X!Tandem* parameters. That dialog window is pictured in FIGURE 3.3, "X!TANDEM PRESETS CONFIGURATION WINDOW (SPECTRUM TAB)". Only the *Spectrum* tab is shown, but the interface is similar for all the other ones.

		sis_FDR_nosen					✓ 1 Load + N
pectrum	Protein	Residue	Scoring	Refine	Output		Qexactive_proteomics_analysis_FDR_nosemi loaded
pectrum	Frotein	Residue	scoring	Renne	Output		spectrum, parent monoisotopic
Precursor io	n [M+H]+ ma	ss tolerance w	indow unit:	C	Daltons •	ppm 💽	mass error units
Precursor io	n [M+H]+ ma	ss tolerance lo	wer window:	1	0	0	Syntax
Precursor io	n [M+H]+ ma	ss tolerance u	pper window:	1	0		The value for this parameter is an ASCII string, with two possible value Daltons or ppm
Anticipate ca	arbon isotope	parent ion as	signments erro	rs: 💽	yes 🔾 no	0	Notes
Set the maxi	mum precurs	sor charge to b	e scored:	4			1. If this parameter is not present or set to a unexpected value, a
Jse chemica	al average or	monoisotopic	mass for fragm	ent ions:	nonoisotopic	<u> </u>	default value is substituted. 2. No error message is generated when a bad value is used for this
Units for fra	gment ion ma	ass tolerance (i	monoisotopic):	•	Daltons	ppm 🕜	parameter.
- ragment io	n mass tolera	ince (monoisot	topic):	0	.02		Description
Control the u	use of the neu	utral loss windo	w:	•	yes 🔵 no	0	The value for this parameter alters the interpretation of the value for the parameters named spectrum, parent monoisotopic mass error plus and spectrum, parent monoisotopic mass error minus. The interpretation of the spectrum is a spectrum of the spec
Neutral loss	mass:			1	8.01057		two values are described below. 1. Da: If this value is set, then the value of spectrum, parent
Neutral loss	window (Dalt	ons):		0	.02	0	1. Due in this value is set, then the value of spectrum, parent monoisotopic mass error plus (T_{+p}) is simply interpreted as a ma and the condition for a parent ion to be considered found is giver $0 < m \cdot p < T_{+p}$
Set the dyna	mic range to	score spectra:		1	00.0		where m is a parent ion in a spectrum and p is the calculated ma of a peptide. The interpretation for spectrum, parent monoisotop
Maximum ni	umber of pea	ks to use in a s	pectrum:	1	00		mass error minus (T_{p}) is given by: $0 > m - p > T_{p}$
Minimum nu	mber of peal	ks for a spectru	um to be consid	ered: 1	5	•	 ppm: If this value is set, then the value of T_{+p} is interpreted as representing parts-per-million (commonly used in mass
Minimum m	/z fragment t	o be considere	d:	1	50.0		spectrometry to express precision), altering the condition for a fragment ion to be as follows: $0 < m \cdot p < m(T_{+e} \times 10^6)$
Minimum pr	ecursor ion [l	N+H]+ to be co	onsidered:	5	00.0		The interpretation for spectrum, parent monoisotopic mass error minus (T _{-p}) is given by: 0 > m - p > m(T _{-p} x 10 ⁴)
Alter how pr	otein sequen	ces are retriev	ed from FASTA	files: 1	000		
Use contrast	t angle:			C) yes 💿 no	0	see also: spectrum, parent monoisotopic mass error plus spectrum, parent
Set number	of threads:			1			monoisotopic mass error minus XI TANDEM API description project
				_			A: IANDER API description project

The configuration of the *X*!*Tandem* presets is performed in this configuration window. This window has its *Spectrum* tab selected. Each parameter is associated to a manual page² that can be displayed by clicking on the interrogation mark button next to it. It is possible to load existing presets from file or to create brand new ones.

FIGURE 3.3: XITANDEM PRESETS CONFIGURATION WINDOW (SPECTRUM TAB)

² The help texts are extracted automatically from the *X*!*Tandem* software documentation.

3.3.1 LOADING EXISTING PRESETS CONFIGURATIONS FROM FILE

It is possible to load existing X!Tandem presets (which is useful in particular if the samples most often come from the same instrument using the same configuration). To this end, first point *i2MassChroQ* to the right directory that contains the presets file of interest (click onto the folder icon at the top right corner of the window shown in FIGURE 3.3, "X!TANDEM PRESETS CONFIGURATION WINDOW (SPECTRUM TAB)"). The presets files in the chosen directory are automatically detected and listed in the drop-down list widget. At this point, select from that list the file of interest and click onto the *Load* button.



WARNING

It is compulsory to click onto the *Load* button to confirm loading of the presets file contents, because these are not updated upon choosing the file name from the drop-down list only.

3.3.2 CREATING NEW PRESETS CONFIGURATIONS

It is possible to create a new presets file by clicking onto the *New* button. This opens an input dialog window for the user to provide a new file name (the edit widget is preset with the currently loaded file's name suffixed with *_copy*).



Tip

One interesting feature of the new presets file creation process is that, if presets are already loaded, *i2Mass-ChroQ* copies the currently displayed settings to the new file. From there, it is possible to create a variant *X!Tandem* presets file, which eases the exploration of the right *X!Tandem* parameters for a given sample data set.

3.3.3 ACTUAL X! Tandem PRESETS CONFIGURATION

The dialog window pictured in FIGURE 3.3, "X!TANDEM PRESETS CONFIGURATION WINDOW (SPECTRUM TAB)" contains a number of tabs where various aspects of the *X*!*Tandem* run settings are handled. Each parameter's documentation can be seen on the pane on the right hand side of the window by clicking onto the question mark button next to it. These manual pages are authoritative because they are taken from the *X*!*Tandem* software package with no transformation whatsoever.

Once the configuration has been performed, click onto the *OK* button. If the parameters were modified, *i2Mass-ChroQ* asks if they should be stored in the file.

3.3.4 RUNNING A PROPERLY CONFIGURED X! Tandem PROCESS

Once the *X*!*Tandem* settings configuration dialog window has been closed, it is possible to run *X*!*Tandem* from inside *i2MassChroQ* by clicking onto the *Run* button at the bottom of the window pictured in FIGURE 3.2, "X!TANDEM-BASED IDENTIFICATION CONFIGURATION".

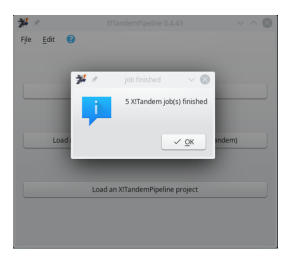
While the computation is carried over, the program shows the feedback dialog window pictured in Figure 3.4, "X!TANDEM RUN FEEDBACK TO THE USER".

[*	Running XITandem	~ ^
	on /home/rusconi/devel/xtpcpp/doc-data/p 6_proteomics_extract_1_A01_urnb-1.mzML	roteomics/
	93%	
Converting /home/rusconi/devel/xtpcpp/doc-c xtpwrp.trmXXQ/msdata.mzxml Running XITandem XI TANDEM Alanine (2017.2.1.4) Loading spectra (mzXML) loaded. Spectra matching criteria = 11892 Starting threads started. Computing models: Spectrum-to-sequence match	lata/proteomics/20120906_proteomics_extract_1_A01_urnb-1.mzML1	o mzXML /tmp/
		Stop

The text in this feedback dialog window is getting incrementally printed all along the computation.

FIGURE 3.4: X!TANDEM RUN FEEDBACK TO THE USER

Once the computation is finished, the feedback dialog window closes and the user is returned to the main program window (FIGURE 3.I, "MAIN PROGRAM WINDOW") albeit with a message shown in FIGURE 3.5, "X!TANDEM RUN FINISHED MESSAGE TO THE USER".



The *X*!*Tandem* run is now finished. Click the *OK* button to access the main program window.

FIGURE 3.5: XITANDEM RUN FINISHED MESSAGE TO THE USER

From the main program window, it is possible to open the *X!Tandem* results file(s) located in the output directory configured above. There are as many output files (XML-based format, and xml extension) as there were mass spectrometry data files to process. The loading of the results files is carried over by first clicking the button labelled *Load identification results (mzIdentML, pepXML, X!Tandem)*. The process is described in SECTION 3.4, "LOADING THE PROTEIN IDENTIFICATION RESULTS".

3.4 LOADING THE PROTEIN IDENTIFICATION RESULTS

The loading of identification results comes with a minimal set of configuration required to instruct *i2MassChroQ* on the way to handle contaminant proteins, for example. This process is pictured in FIGURE 3.6, "CONFIGURATION OF THE LOADING OF THE IDENTIFICATION RESULTS" and is described in the following section.

sults handling mode		
• Com <u>b</u> ine	O Indi <u>v</u> idual	
oose result files		
	utput.d/20120906_balliau_extract_1_A01_urnb-1.xml utput.d/20120906_balliau_extract_1_A02_urzb-1.xml	
Number of files: 2	Clear list + Add	d files
Contaminants		
Contaminants file	O Contaminant <u>r</u> egular expression	
Contaminant removal mode	Clear list Add f Groups	îles
		îles
O <u>P</u> rotein list		īles
• <u>P</u> rotein list ptide and protein filters	● <u>G</u> roups	iles
Protein list ptide and protein filters Peptide threshold on:	<u>G</u> roups <u>Evalue</u> <u>FDR</u>	
Protein list ptide and protein filters Peptide threshold on: Peptide Evalue:	<u>G</u> roups <u>Evalue</u> <u>EDR</u> <u>0.050000</u>	
Protein list ptide and protein filters Peptide threshold on: Peptide Evalue: Peptide FDR:	<u>G</u> roups <u>Evalue</u> <u>FDR</u> <u>0.050000</u> <u>1.0%</u>	
Protein list ptide and protein filters Peptide threshold on: Peptide Evalue: Peptide FDR: Number of peptides per protein:	 <u>G</u>roups <u>Evalue</u> <u>EDR</u> 0.050000 1.0% 2 	
Protein list ptide and protein filters Peptide threshold on: Peptide Evalue: Peptide FDR: Number of peptides per protein: Overall samples:	● <u>G</u> roups ● <u>Evalue</u> <u>EDR</u> 0.050000 1.0% 2 ✓	

Loading identification results comes with some configuration that is described in the text.

FIGURE 3.6: CONFIGURATION OF THE LOADING OF THE IDENTIFICATION RESULTS

3.4.1 IDENTIFICATION DATA LOADING CONFIGURATION

- *Results handling mode*: there are two possibilities:
 - *Combine*: in this mode, all the identification results coming from different identification results files are merged into a single set. That single set is the basis for the protein inference step and the identified proteins are listed into a single protein list window.

¥ *	new_project - X!Tand	emPipeline 0.4	4.43 🗸 🗸	$\sim \otimes$
F <u>i</u> le <u>E</u> dit 🕜				
Identifications	Filter parameters	FDR	Mass precision	
	View protein list	View MS	identification list	
	Number of samples	: 5		- I
	Number of groups	1288		
	Number of subgroups	1650		
Numb	per of grouped proteins	2860		
Number of gr	ouped distinct peptides	8259		
View PTM is	ands			

When loading multiple identification results files in *Individual* mode, the selection of any given identification results file is performed by selecting its name from the drop-down list widget *and* by clicking onto the *View protein list* button. Note that some metadata about the identifications are updated beneath the drop-down list widget.

FIGURE 3.7: SELECTING A PARTICULAR IDENTIFICATION RESULTS FILE'S DATA SET

• *Individual*: in this mode, the identification results coming for various files are kept separated. Thus, the identification results coming from each file are used for a separate protein inference step. The identified proteins list is thus displayed for *each single file* in turn. The selection of the file for which the protein list needs to be displayed is done via the main program window that changes its appearance:



When loading multiple identification results files in *Individual* mode, the selection of any given identification results file is performed by selecting its name from the drop-down list widget *and* by clicking onto the *View protein list* button. Note that some metadata about the identifications are updated beneath the drop-down list widget.

FIGURE 3.8: SELECTING A PARTICULAR IDENTIFICATION RESULTS FILE'S DATA SET

Right after having selected an identification results file, click onto the *View protein list* to display the protein identifications list. That list has been obtained by performing the protein inference on the file's protein identification results (see Section 2.2.5.4, "PROTEIN INFERENCE: FROM PSMs TO PROTEIN IDENTITIES"). The window that opens up will be described later (see Section 4.1, "The PROTEIN LIST WINDOW").



It is possible to open multiple protein list windows, each showing the identifications from a different file: maintain the Ctrl keyboard key pressed while clicking onto the *View protein list* button.

• *Choose results files*: by clicking onto the *Add files* button, the user is provided a file selection dialog window from which any number of protein identification results files might be selected for loading.

Note that it is possible to list all the opened protein identification results files by clicking onto the *View MS identification list* button. The window that opens up will be described later (see SECTION 3.4.2, "DIS-PLAYING THE MS IDENTIFICATIONS LIST").

- *Contaminants*: there are two possibilities here.
 - *Contaminants files*: when this radio button widget is selected, the list of contaminant proteins will be loaded from the files selected by clicking onto the *Add files* button.
 - *Contaminant regular expression*: when this radio button widget is selected, a text edit widget is shown, replacing the widget listing the contaminants database files. In this text edit widget, the user may enter a regular expression to match the accession number field of the protein databases that were used for the protein identification step. In this situation, the user must use specially crafted protein databases in which the contaminant proteins were tagged on the accession number using a particular text pattern. That particular text pattern is then matched against the *Contaminant regular expression* that the user enters in the text edit widget.
- *Contaminant removal mode*: there are two possibilities. The contaminant removal is the process by which, when identified proteins match proteins in the contaminants realm (either from the contaminants dabase files or as determined using the regular expression), they are disregarded for the later protein visualization steps.
 - *Protein list*: in this mode, as soon as a protein identification loaded from a protein identification results file matches a contaminant protein, it is disregarded.
 - *Groups*: in this mode, the protein inference process goes all the way through to the determination of the protein groups (see FIGURE 2.8, "PROTEIN INFERENCE: CONSTRUCTING A CONSOLIDATED PROTEIN IDENTIFICATIONS LIST"). When protein groups have sub-groups that contain a contamimant protein, then the whole group is disregarded. This might appear drastic, but our experience is that most often, the sub-groups in a group do identify proteins belonging to the same family. Therefore, if one protein is contaminant, all the other proteins in the group are supposed to be such also.
- *Peptide and protein filters*: this group box widget holds some parameters that configure the way protein inference is to be performed.

- Peptide threshold on: there are two possibilities:
 - *E-value*: all the PSMs having an expectation value higher than that value are disregarded. Enter the value in the spin box widget labelled *Peptide E-value*. A typical value for the *X!Tandem* engine is 0.05. When more stringent results are desirable, setting 0.02 should yield satisfactory results. See SECTION 2.2.5.2, "COMPUTATION OF THE PEPTIDE EXPECTATION VALUE (E-VALUE)" of a detailed explanation of the E-value computation.
 - *FDR* (false rate discovery): the PSMs are disregarded if their FDR value does not match this parameter. Enter the value in the spin box widget labelled *Peptide FDR*. A typical setting is *1%*.

😰 Тір

Using *FDR* is most useful when the identification results come from a database searching engine that does not compute an E-value. However, it does only work if the searching step was performed also on a decoy database. In *X*!*Tandem* the decoy database is crafted by reversing the peptide sequences. In this case, when proteins are identified on the basis of the reversed peptide PSM, then the protein identity is tagged with the "reversed" string, which might be used with the *Contaminant regular expression* setting defined earlier.

- *Number of peptides per protein*: this is the minimal required number of peptides that must be identified as belonging to a given protein in order to consider that protein identity as a valid one. These peptides have to be from non-contaminant proteins, of course.
- Overall samples: when checked and if multiple identification results files are to be loaded, then the *Number of peptides per protein* requirement might be fulfilled by looking for peptides in all the loaded files. For example, if one results file provides one peptide for a protein identification and another file provide another peptide (different from the first one) to identify the same protein, and if the *Number of peptides per protein* is 2, then the protein is considered as a valid protein. If not checked, that number of peptides requirement must be fulfilled by looking into each results file separately. This last setting is more stringent. A typical value for this setting is 2.



This setting needs to be checked in at least one case: when a complex peptidic mixture is separated by ion chromatography (typically on an SCX—strong cation exchange—resin) and the different fractions are analyzed by bottom-up proteomics. The peptides coming from a given protein might be located in different fractions, and thus in different protein identification results files!

- *Protein Evalue*: threshold above which a protein identification is disregarded (see Section 2.2.5.3, "Computation of the Protein Expectation Value (E-value)").
- *Protein Evalue (log10)*: convenience spin box widget for the user to easily set the protein E-value.
- *Pep repro*: if set to 1, a peptide, to be accounted for, needs to be found in one protein identification results file. If set to a greater number, then that peptide needs to be found in that number of results files. This setting sets more stringent protein identification conditions each time it is incremented.

3.4.2 DISPLAYING THE MS IDENTIFICATIONS LIST

Alignments Groups	i dentification Run II 🗸	Identification result file	Ms Run File	Identification engine	Fasta file(
elect Group	identa0	20120906_proteomics_extract_1_A01_urnb-1.xml	20120906_proteomics_extract_1_A01_urnb-1.mzML	🖄 X!Tandem 2017.2.1.4	2 fasta use
elect Group	identa1	20120906_proteomics_extract_1_A05_urnb-2.xml	20120906_proteomics_extract_1_A05_urnb-2.mzML	🖸 X!Tandem 2017.2.1.4	2 fasta use
elect Group	identa2	20120906_proteomics_extract_1_A09_urnb-3.xml	20120906_proteomics_extract_1_A09_urnb-3.mzML	🗹 X!Tandem 2017.2.1.4	2 fasta use
elect Group	identa3	20120906_proteomics_extract_1_B01_urnb-4.xml	20120906_proteomics_extract_1_B01_urnb-4.mzML	🖸 X!Tandem 2017.2.1.4	2 fasta use
elect Group	identa4	20120906_proteomics_extract_1_B05_urnb-5.xml	20120906_proteomics_extract_1_B05_urnb-5.mzML	🗹 X!Tandem 2017.2.1.4	2 fasta use

This window displays a list of all the files that were involved in the X!Tandem run (first columns).

FIGURE 3.9: DISPLAYING THE MS IDENTIFICATIONS LIST (FIRST COLUMNS)

Total spectra u	used 🗄 Total spectra assigned	Total unique assigned	Assignment %	Sample ID	Total MS1	Total MS2	Total MS3	TIC MS1	TIC MS2	TIC MS
1892	7202	6167	60.56 %	msruna1	4614	11962	0	6.06742e+12	1.47302e+11	0
2008	7341	6233	61.13 %	msruna2	4644	12080	0	6.35385e+12	1.52904e+11	0
2004	7307	6194	60.87 %	msruna3	4621	12113	0	6.27244e+12	1.56102e+11	0
2083	7357	6229	60.89 %	msruna4	4652	12139	0	6.79543e+12	1.67761e+11	0
2058	7315	6164	60.67 %	msruna5	4610	12128	0	6.35616e+12	1.58973e+11	0

This window displays a list of all the files that were involved in the *X*!*Tandem* run (last columns).

FIGURE 3.10: DISPLAYING THE MS IDENTIFICATIONS LIST (LAST COLUMNS)

3.4.3 SAVING *i2MassChroQ* projects

Once exploration and optional modication of the identification data have been performed, the user can save the resulting data set into a *i2MassChroQ* project by selecting the *Save project* menu item of the *File* menu in the main program window (the extension of the file name typically should be <u>xpip</u>). See <u>Section 3.5</u>, "Loading I2MassChroQ projects" for loading such a project.

3.5 LOADING *i2MassChroQ* PROJECTS

Loading of *i2MassChroQ* project files (file of xpip) extension) is only possible if the user has previously

- Loaded identification results;
- Saved the data to an *i2MassChroQ* project file using the *Save project* menu item of the *File* menu in the main program window.

4 Exploring identification data

This chapter describes in detail all the steps that the user accomplishes in their data exploration session. The general workflow is to start by looking at a protein identification results window and then by going into the details of the various identifications listed in it. This latter task entails looking into the peptides that provided the protein identification and then looking at the mass spectrum that provided the peptide identification. The mass spectrum, that is, the MS/MS spectrum, has features aimed at allowing the user to make an informed opinion on the validity of the peptide *vs* mass spectrum match (PSM) at hand. At each moment, it is possible to invalidate a PSM and the identification results are recomputed automatically by taking into account the modification entered by the user.

4.1 The Protein List Window

When identification results files are loaded, *i2MassChroQ* automatically performs the protein inference process by using the configuration settings described in Section 3.4.1, "IDENTIFICATION DATA LOADING CONFIGURA-TION".

4.1.1 THE PROTEIN LIST TABLE VIEW

When the protein inference process is finished, *i2MassChroQ* displays the protein identifications list in a table view, as pictured in FIGURE 4.I, "THE PROTEIN LIST WINDOW".

port Columr	is Show only				untitled - Pro	cein lisc							~
checked *	group	accession	description	log(Evalue)	Evalue	spectra	specific spectra	sequences	pecific sequence:	coverage	MW	PAI	emPA
V	a1.a1.a1	GRMZM2G083841_P01	P04711 Phosphoenolpyruvate	-436.204	0	269	251	58	54	56.49 %	109272	1.84615	69.1704
• •	c117.a1.a1	GRMZM2G033841_101	NP_001152746 ascorbate	-71.1121	7.72497e-72	32	7	9		52.00 %	27353.9	1.33333	20.5443
• •	c117.a2.a1	GRMZM2G054300_P01	NP_001150192 APx1 - Cytosolic	-59.8189	1.5174e-60	27	2	8	1	44.80 %	27290.8	1.16667	13.678
- -	c117.a2.a1	GRMZM2G054300_P04	NP_001150192 APx1 - Cytosolic		1.5174e-60	27	2	8	1	37.46 %	32330.3	1	9
• •	C117.02.02	GRMZM2G054300_P02	NP_001150192 APx1 - Cytosolic		7.66366e-43	17	2	5		35.94 %	20841.5	1.125	12.3352
v V		GRMZM2G054300_P02	NP_001150192 APx1 - Cytosolic	-42.1156	7.66366e-43	17		5		31.80 %	23404.6	0.9	6.94328
• •	c407.a1.a1	GRMZM2G046841_P01	B6SIF9 Histone H2B	-24.4482	3.56257e-25	8	3	5		37.33 %	16133.9	0.714286	4.17947
- J	c407.a1.a2	GRMZM2G119071_P01	P30756 Histone H2B.2	-24.4482	3.56257e-25	8	3	5		37.33 %	16163.9	0.714286	4.17947
• •	b36.a1.a1	GRMZM2G449496_P01	NP_001169327 hypothetical	-148.005	9.88939e-149	37	37	19	19	49.19 %	51820.9	0.88	6.58578
• •	b78.a1.a1	GRMZM2G027995_P01	Q41741 Eukaryotic initiation	-102.438	3.64897e-103	29	2	12	1	36.47 %	46952.9	0.9	6.94328
• •	b78.a1.a2	GRMZM2G027995_P02	Q41741 Eukaryotic initiation	-102.438	3.64897e-103	29	2	12	1	36.47 %	46952.9	0.9	6.94328
- -	b78.a2.a1	GRMZM2G116034_P02	NP_001104874 translation	-97.3364	4.60922e-98	29	2	12	1	36.47 %	46922.9	0.85	6.07946
 ✓ 		GRMZM2G116034_P01	NP_001104874 translation	-91,1493	7.09111e-92	28	-	11		33.58 %	46663.8	0.8	5.30957
v	b60.a1.a1	GRMZM5G845611_P01	B4F8L7 Glyceraldehyde-3	-98.5797	2.63223e-99	52	36	14		37.64 %	47150.2	0.954545	8.00628
 ✓ 	b60.a2.a1	GRMZM2G337113_P02	P09315 Glyceraldehyde-3	-80.9409	1.14578e-81	41	25	10		33.25 %	42830	1.14286	12.895
~ V		GRMZM2G129246_P01	NP_001146005 hypothetical	-73.2632	5.45513e-74	24		12		32.02 %	53278.8	0.576923	2.77505
✓	b82.a1.a1	GRMZM2G129246_P02	NP_001146005 hypothetical	-86.6197	2.40026e-87	25	25	13	13	46.07 %	40021.1	0.842105	5.95193
v		GRMZM2G129246_P05	NP_001146005 hypothetical	-60.7567	1.75125e-61	20		10		40.45 %	32985.5	0.8125	5,49382
✓		GRMZM2G129246_P03	NP_001146005 hypothetical	-34.8504	1.41117e-35	10		7		38.89 %	24162.3	0.538462	2.45511
v		GRMZM2G129246 P04	NP_001146005 hypothetical	-34.8504	1.41117e-35	10		7		38.89 %	24162.3	0.538462	2.45511
~ •	c117.a3.a1	GRMZM2G140667_P01	NP_001105500 ascorbate	-24.1051	7.85039e-25	10	8	5		24.13 %	30900.7	0.357143	1.27585
v	c117.a3.a2	GRMZM2G140667_P02	NP_001105500 ascorbate	-24.1051	7.85039e-25	10	8	5		24.47 %	30482.5	0.384615	1.42446
v		 GRMZM2G140667_P04	- NP_001105500 ascorbate	-20.5201	3.01938e-21	8		4		31.41 %	20781.4	0.5	2.16228
v		GRMZM2G175867_P01	NP_001130422 hypothetical	-1.60206	0.025	1		1		2.12 %	66514.1	0.0454545	0.110336
/		GRMZM2G175867_P02	NP_001130422 hypothetical	-1.60206	0.025	1		1		3.61 %	38959.6	0.0588235	0.145048
✓		GRMZM2G153969_P01	NP_001149564 OB-fold nucleic	-2.52288	0.003	2		1		8.43 %	18248.5	0.142857	0.389495
✓		GRMZM2G174479_P01	B6TM56 Chloroplast outer	-3.85387	0.00014	2		1		11.40 %	12394.1	0.166667	0.467799
v		GRMZM2G167698_P01	seq=translation;	-2.5376	0.0029	2		1		2.48 %	58821.2	0.04	0.0964782
✓		GRMZM2G009232_P01	NP_001141257 hypothetical	-2.5376	0.0029	2		1		2.49 %	58693.1	0.0416667	0.100694
v		GRMZM2G009232_P02	NP_001141257 hypothetical	-2.5376	0.0029	2		1		3.95 %	37105.2	0.0769231	0.193777
v		GRMZM2G009232_P03	NP_001141257 hypothetical	-2.5376	0.0029	2		1		3.94 %	37454.5	0.0769231	0.193777
v	c141.a1.a1	GRMZM2G318635_P01	seq=translation;	-66.2764	5.29139e-67	16	16	10	10	9.69 %	190929	0.136986	0.370839
v	c530.a1.a1	GRMZM2G157462_P01	NP_001152484 dynamin-2A	-20.5331	2.93026e-21	7	7	4	4	4.93 %	99589.6	0.0851064	0.216484

proteins all:6814 valid:3895 valid&checked:3895 grouped:2481 displayed:6814

The protein identifications list window displays the proteins assembled into groups. A number of metadata about the identifications are shown in a number of colums, the contents of all of which are described in detail in the text.

FIGURE 4.1: THE PROTEIN LIST WINDOW

The columns that make the protein list table view are detailed below:

- *Checked*: if checked, the identified protein listed on the table row is set to an "accepted" state. By default, all proteins are set to this accepted state. Unchecking a protein determines the protein inference reprocessing, because disregarding a protein modifies the whole protein identifications results set;
- *group*: the group the protein belongs to;
- *accession*: the accession number field of the protein database;
- *description*: the description field in the protein database;
- *log(E-value)*: the Log10 of the protein E-value;
- *E-value*: the protein E-value;
- *spectra*: the number of spectra that identified the protein;

- *specific spectra*: the number of spectra that identified *only* this protein;
- *sequences*: the number of peptidic sequences that can be assigned to this protein;
- *specific sequences*: the number of peptidic sequences that can be assigned *only* to this protein;
- *coverage*: the percentage of the protein sequence covered by the peptides that identified it;
- *MW*: the molecular weight of the protein (M_r);
- PAI: "Protein abundance index". This index was defined as the "number of peptides identified divided by the number of theoretically observable tryptic peptides". See HTTPS://WWW.NCBI.NLM.NIH.GOV/PMC/ ARTICLES/PMCI86633/ ♂;
- emPAI: "Exponentially modified protein abundance index". This index was defined as emPAI = 10^{PAI} − 1. See https://pubmed.ncbi.nlm.nih.gov/15958392/ 7.

It is possible to select the columns that must be displayed in the table by checking or unchecking the corresponding item in the *Columns* menu.

The Show only menu allows one to select the kind of protein items to be shown:

- *Valid proteins*: when checked, the program only shows valid proteins, that is, protein identifications that fullfill the restriction parameters, like protein E-value, for example. These parameters were set at protein identification results loading time but can be modified later;
- *Checked proteins*: show only the proteins that were checked. This setting is useful when the user has unchecked a number of proteins and that they want to regularly keep an eye on them. When proteins are unchecked, the protein inference process is run anew to compute a new grouping by taking *not* into account the proteins that were disregarded;
- *Grouped proteins*: only show the proteins that belong to a group.

The protein identifications list table view above shows greyed protein identities. These are proteins that, by current filter parameters (E-value threshold, for example), are considered *not* valid.

4.1.2 Operations in the Protein List Window

The *Protein list* window houses a number of useful features that let the user scrutinize the protein identifications and also modify the results to suit either more or less stringent filtering parameters.

Searching data in the table view. One interesting feature of the *Protein list* window is the ability to search through the table's contents using the *Search* item at the bottom of the window. A number of fields of the protein record, that is, columns in the table view, might be searched.

Dynamic setting of the filter parameters. *i2MassChroQ* provides a rather high level of flexibility: once a protein identification results set of files has been loaded and that the protein inference process is achieved, the resulting protein groups are displayed in the *Protein list* window. At this time, the grouping was performed using the parameters set as pictured in Section 3.4.1, "IDENTIFICATION DATA LOADING CONFIGURATION". It is nonetheless possible to modify these parameters on the fly using the main program window's *Filter parameters* tab, as pictured in FIGURE 4.2, "PROTEIN IDENTIFICATION FILTER PARAMETERS TAB OF THE MAIN WINDOW".

Identifications Filter parameters	FDR Mass precision					
Peptide threshold on:	Evalue <u>F</u> DR					
Peptide Evalue:	0.050000					
Peptide FDR:	1.0%					
Number of peptides per protein:	2 *					
Overall samples:						
Protein Evalue:	0.01000000					
Protein Evalue (log10):	-2.00					
Pep repro:	1					
Contaminants O Contaminants file	Contaminant <u>r</u> egular expression					
^contaminant.*						
Contaminant removal mode						
O <u>P</u> rotein list	• <u>G</u> roups					
	✓ Apply					

The filter parameters in this dialog box window do mirror the ones that one can set prior to loading protein identification results files. When modified, these parameters elicit a complete run of the protein inference process.

FIGURE 4.2: PROTEIN IDENTIFICATION FILTER PARAMETERS TAB OF THE MAIN WINDOW

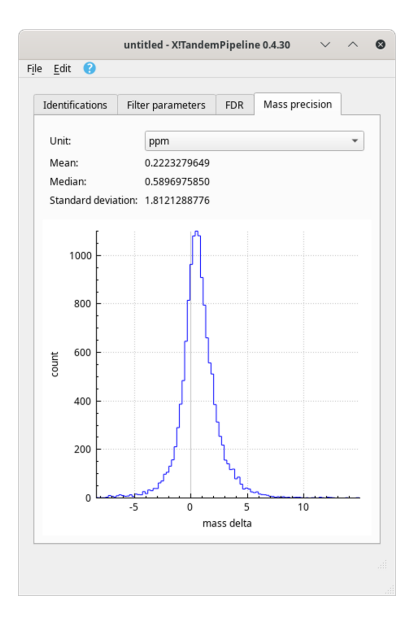
Real time update of the false discovery rate. The false discovery rate (FDR) is recalculated at each protein inference process. The data regarding this quality assessment criterion are shown in FIGURE 4.3, "FALSE DISCOVERY RATE (FDR) DATA AFTER A PROTEIN INFERENCE PROCESS IS RUN".

		untitled - X!Tan	demPipelin	ne 0.4.30	\sim	~ Ø		
ile	<u>E</u> dit 🕜							
1	Identifications	Filter paramete	rs FDR	Mass pre	cision			
F	False discovery r	ate (FDR)						
	FDR on psm:		0.2101 %					
	FDR on peptide	25:	0.2445 %					
	FDR on protein	IS:	0.0000 %					
ľ	Decoy settings							
	O Decoy file		• Decoy <u>r</u> egular expression					
	.*reversed\$							
			O Comput	te cumulate	✓ App d FDR on I			

The various data bits about the false discovery rate that is computed each time a protein inference process is run. Note that it is possible to modify the *Decoy settings*, after which the *Apply* button triggers the recalculation of the FDR.

FIGURE 4.3: FALSE DISCOVERY RATE (FDR) DATA AFTER A PROTEIN INFERENCE PROCESS IS RUN

Distribution of mass errors on PSMs plotted in a histogram. It is possible to visualize the distribution of the mass errors over the whole dataset, as pictured in FIGURE 4.4, "MASS PRECISION QUALITY ASSESSMENT". The histogram plots the number of mass spectra that could achieve a PSM against the mass error (mass delta), that is, the difference between the experimental peptide mass and the calculated peptide mass. FIGURE 4.4, "MASS PRECISION QUALITY ASSESSMENT".



The histogram plots the number of PSMs against the mass error calculated between the experimental mass of the peptide and the calculated mass.

FIGURE 4.4: MASS PRECISION QUALITY ASSESSMENT

The mass delta calculation involves only the peptides that successfully identifed proteins that are currently checked in the protein identification list and that satisfy the filter parameters. The proteins identified in the decoy database are not processed. The unit of the mass delta may be selected using the *Unit* drop-down list. Two units are available: ppm (for part-per-million) or Dalton.

Exporting the final protein identifications list to a spread sheet. Once all the proteins in the identifications list have been properly checked, the user might export the data set to an OpenDocumentFormat (ODF) spread sheet file using the *As ODS file* menu item of the main window's *Export* menu.

4.1.3 Delving Inside the Protein Identification Data

The protein list table view, as pictured in FIGURE 4.1, "THE PROTEIN LIST WINDOW" is actually an active matrix in which the user can easily trigger the exposition of the data that yielded any protein identification element of the table. This is simply done by clicking onto any cell of the table at the row matching the protein for which scrutiny of the data is desired.

Depending on the column at which the mouse click happens, there might be two different windows showing up:

• The *Protein details* window, showing the sequence of the protein, the matching peptides and other informational data bits, as pictured below:

	untitled - Protein details	~ ^ 0
B4F8L7 G ; seq=trar parent_tra MATHAAL KVAINGF PWGELGI EEFGIVK VKTGITA	G845611_P01 lyceraldehyde-3-phosphate dehydrogenase BPutative uncharacterized protein islation; coord=1:64104616413123:1; anscript=GRMZM5G845611_T01; parent_gene=GRMZM5G845611 AASRIPAGARLHSRAPASSRHGVQRLDFADFSGLRPGSCSVSAAAREASFSDVLGAQLVAKA GRIGRNFLRCWHGREDSPIDVVVVNDSGGVRNASHLLKYDSMLGTFKADVKIVDDTTISVDG DIVIEGTGVFVDGPGAGKHIQAGAKKVIITAPAKGADIPTYVVGVNEGDYDHSVADIISNAS GTMTTTHSYTGDQRLLDASHRDLRRARAAALNIVPTSTGAAKAVALVLPQLKGKLNGIALRV DDVNAAFRKAAAGPLQGILEVCDVPLVSVDFRCSDVSCTIDASLSMVMGDDMVKVVAWYDNE AAAGSGDPLEDFCKDNPETDECKVYEA	KPITVVSSRDPLKL CTTNCLAPFVKILD PTPNVSVVDLVINT
coverage MW	37.6392 % 47.1502 kDa	📴 get PDF
E-value	2.63223e-99 (log10: -98.5797)	

When one cell in the *Accession*, *Description* or *Coverage* column is clicked, this window shows up and displays the sequence of the protein, the coverage of the peptides and other useful data.

FIGURE 4.5: PROTEIN DETAILS WINDOW

• When one cell in any one of the remaining columns is clicked, the window that shows up is the *Peptide list* window showing a list of all the peptide identifications, to be described in the next section.



Tip

When clicking one cell in one column and one given row, the corresponding window shows up, if one was not already open. If one window is already open, no other window shows up, but the existing window has its data updated to match the new protein row being clicked on.

It is possible to have multiple windows opened at a time by clicking a new row while maintaining the Ctrl key pressed.

4.2 The Peptide List Window

The *Peptide list* window displays all the data in a table view similar to the one used to display the protein list described in the previous sections.

4.2.1 The Peptide List Table View

The *Peptide list* table view has a pretty large number of columns to display all the data about each peptide that identified a given protein. These columns are described in the following figures.

					unditied -	Peptide list					~ ^
<u>kport</u> <u>C</u> olum	ns S <u>h</u> ow only										
RMZM2G1190	71_P01										
30756 Histone	H2B.2 seq=translation	n; coord=4:630925	7163093826:-1; parent_transcript=GRMZM2	G119071_T01; pare	nt_gene=GRMZN	12G119071					
Checked	 Peptide ID 	Engine	Sample	Scan	Index	RT (min)	RT (s)	Charge	Obs. m/z	N-term	Sequence
v	pepc407a1	X!Tandem	20120906_balliau_extract_1_A01_urnb-1	15581		46.16'	2769.40"	2	880.42310	к	AMSIMNSFINDIFEK
v	pepc407a1	X!Tandem	20120906_balliau_extract_1_A02_urzb-1	16007		46.19'	2771.48"	2	880.42383	к	AMSIMNSFINDIFEK
v	pepc407a2	X!Tandem	20120906_balliau_extract_1_A01_urnb-1	9412		30.56'	1833.81"	2	470.29733	R	LVLPGELAK
v	pepc407a2	X!Tandem	20120906_balliau_extract_1_A02_urzb-1	9642		30.21'	1812.83"	2	470.29752	R	LVLPGELAK
v	pepc407a3	X!Tandem	20120906_balliau_extract_1_A01_urnb-1	2296		12.48'	748.85"	3	579.31262	к	KPAAKKPAEEEPAAEK
✓	pepc407a4	X!Tandem	20120906_balliau_extract_1_A01_urnb-1	2038		11.43'	685.96"	3	541.61798	к	KPAEEEPAAEKAPAGK
v	pepc407a4	X!Tandem	20120906_balliau_extract_1_A02_urzb-1	1971		10.86'	651.38"	3	541.61755	к	KPAEEEPAAEKAPAGK
√	pepc407a6	X!Tandem	20120906_balliau_extract_1_A02_urzb-1	4898		19.07'	1144.02"	2	590.81934	к	QVHPDIGISSK
(

The *Peptide list* table view has many columns (first columns).

FIGURE 4.6: THE PEPTIDE LIST WINDOW (FIRST COLUMNS)

xport <u>C</u> olumns Show only													
RMZM2G119071_P01													
30756 Histone H2B.2 seq=translatio	on; coord=4:6309257163	093826:-1; par	ent_transcript=G	RMZM2G11907	1_T01; parer	nt_gene=GRMZM2G	119071						
Sequence	C-term	Modifs	Start	Length	Used	Subgroups	E-value	Cumulated FDR	Obs. [M+H]+	Theor. [M+H]+	Delta [M+H]+	Delta (ppm)	HyperScore
AMSIMNSFINDIFEK	L		84	15	2	c407.a1 c407.a2	4.2e-07		1759.83892	1759.83936	-0.000440704	-0.250423	41.1
AMSIMNSFINDIFEK	L		84	15	2	c407.a1 c407.a2	4.6e-07		1759.84038	1759.83936	0.0010243	0.582039	41.2
LVLPGELAK	н		126	9	2	c407.a1 c407.a2	0.021		939.58739	939.58734	4.55356e-05	0.0484634	27.5
LVLPGELAK	н		126	9	2	c407.a1 c407.a2	0.031		939.58776	939.58734	0.000411536	0.437996	27.5
KPAAKKPAEEEPAAEK	A	1K:42.01	8	16	1	c407.a1	4.4e-08		1735.92331	1735.92249	0.000823266	0.474253	47.3
KPAEEEPAAEKAPAGK	к		13	16	1	c407.a1	1.8e-05		1622.83939	1622.83843	0.000964245	0.594172	35.1
KPAEEEPAAEKAPAGK	к		13	16	1	c407.a1	3.4e-05		1622.83811	1622.83843	-0.000317755	-0.195802	35.5
QVHPDIGISSK	A		73	11	2	c407.a1 c407.a2	5.1e-05		1180.63140	1180.63206	-0.000666812	-0.564792	44.9
(

The *Peptide list* table view has many columns (last columns).

FIGURE 4.7: PEPTIDE LIST WINDOW (LAST COLUMNS)

The table's contents are well described by the column headers that are self-explanatory. When hovering over a column header with the mouse cursor, a tool-tip explanatory text is displayed.

It must be noted that more columns might make the table view depending on the protein identification data that were loaded. Indeed, depending on the database searching engine that was used for the protein identification, the data to be displayed vary. The whole list of columns that might be displayed in the table view are pictured in FIGURE 4.8, "COLUMNS THAT POPULATE THE PEPTIDE LIST TABLE VIEW"

✓ Checked	✓ Inter prophet prob.
✓ Peptide ID	✓ HyperScore
✓ Engine	✓ Mascot score
✓ Sample	✓ Mascot E-value
✓ Scan	✔ OMSSA E-value
✓ Index	✔ OMSSA p-value
🗸 RT (min)	✔ MS-GF raw score
✓ RT (s)	✓ MS-GF de novo
✓ Charge	✓ MS-GF energy
✓ Obs. m/z	✔ MS-GF spectral E-value
✓ N-term	✓ MS-GF E-value
✓ Sequence	✔ MS-GF isotope error
✓ C-term	✓ Comet XCorr
✓ Modifs	✔ Comet DeltaCn
✓ Label	 Comet DeltaCnStar
✓ Start	✔ Comet SpScore
✓ Length	✔ Comet SpRank
✓ Used	✔ Comet E-value
✓ Subgroups	 DeepProt matched peaks
✓ E-value	 DeepProt fitted peaks
 Cumulated FDR 	 DeepProt match type
✔ Obs. [M+H]+	 DeepProt status
✔ Theor. [M+H]+	✓ DeepProt mass delta
✔ Delta [M+H]+	✓ DeepProt mass delta pos.
🗸 Delta (ppm)	
✓ Prophet prob.	

Depending on the provenience of the protein identifications (the database search engine), the columns that are part of the table view differ. This full list is displayed when selecting the *Columns* menu.

FIGURE 4.8: COLUMNS THAT POPULATE THE PEPTIDE LIST TABLE VIEW

4.2.2 Operations in the Peptide List Window

The *Peptide list* window houses a number of pretty interesting features that let the user scrutinize the peptide details.

Searching data in the table view. One interesting feature of the *Peptide list* window is the ability to search through the table's contents using the *Search* item at the bottom of the window. A number of fields of the protein record, that is, columns in the table view might be searched.

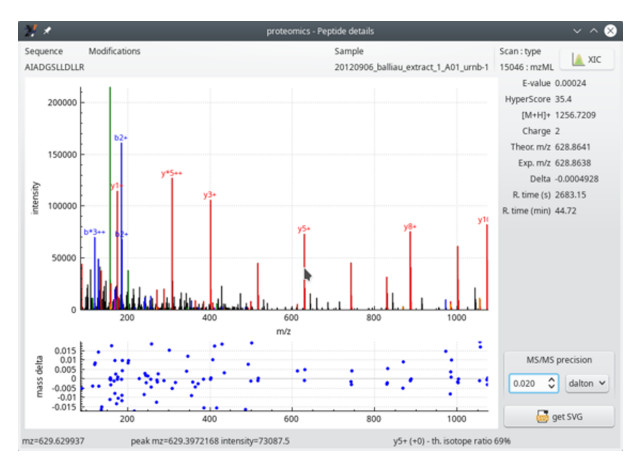
Exporting the final protein identifications list to a spread sheet. Once all the peptides in the identifications list have been properly checked, the user might export the data set to an OpenDocumentFormat (ODF) spread sheet file using the *As ODS file* menu item of the main window's *Export* menu.

4.2.3 Delving Inside the Peptide Identification Data

The *Peptide list* table view, as pictured in FIGURE 4.6, "THE PEPTIDE LIST WINDOW (FIRST COLUMNS)" is actually an active matrix in which the user can easily trigger the exposition of the data that yielded any peptide identification element of the table. This is simply done by clicking onto any cell of the table at the row matching the peptide for which scrutiny of the data is desired.

4.2.3.1 The Peptide Details Window

When clicking any one of the cells of the peptide list table view, one window shows up that details the various data elements for the peptide documented in the table row. The window is pictured in FIGURE 4.9, "PEPTIDE DETAILS WINDOW".



This window displays the MS/MS spectrum that allowed identifying a peptide (that is, a PSM). A number of informational data bits are displayed, like the MS/MS scan number, the E-Value for the peptide, along with its Hyperscore, for example (see text below for a thorough description).

FIGURE 4.9: PEPTIDE DETAILS WINDOW

In FIGURE 4.9, "PEPTIDE DETAILS WINDOW", the two graphs show the following:

- The top graph displays the mass spectrum of this PSM. This MS/MS spectrum has its recognized peaks in the *b* and *y* ion series labelled in blue and red respectively. When the mouse cursor hovers over a mass peak, the details of that mass peak are printed in the status bar of the window (bottom line). Navigating the spectrum is straightforward: to zoom/unzoom in a given area of the spectrum, point the mouse cursor at the peak of interest and use the mouse wheel to zoom/unzoom. To modify the ordinate intensity scale, click onto the axis and drag the mouse upwards or downwards.
- The bottom graph plots—for each matching MS/MS peak (that is, *b* and *y* ion series)—the mass difference (mass delta) between the ion's measured mass and the theoretical mass. In this example, we see that the *y* ion series is moderately matched (large error range).

It is possible to set the *MS/MS precision* to a determinate value and unit (Dalton, ppm or res). The value entered in the spin box widget modifies the assignment of the fragmentation peaks.



The MS/MS spectrum mass peaks are annotated using the following naming convention:

- *: neutral NH₃ loss;
- *o*: neutral H₂O loss;

The ion charge is displayed in the form of "+" or "++" text strings.

The right hand side margin of the window provides a number of data about the PSM, like the peptide E-value, the HyperScore, the ion charge, the theoretical and experimental masses, the difference between the two, the retention time at which this ion was detected... These informational data bits are self-explanatory.

The *XIC* button at the top right corner of the window triggers the calculation of the extracted ion current chromatogram, as described in the section below.

4.2.3.2 The XIC Viewer Window for the Peptide Details

One interesting feature of the *Peptide details* window, is the *XIC* button (top right) that triggers the calculation of an extracted ion current chromatogram, as pictured in FIGURE 4.10, "THE EXTRACTED ION CURRENT (XIC) CHROMATOGRAM VIEWER WINDOW".



TIP: WHAT IS A XIC CHROMATOGRAM?

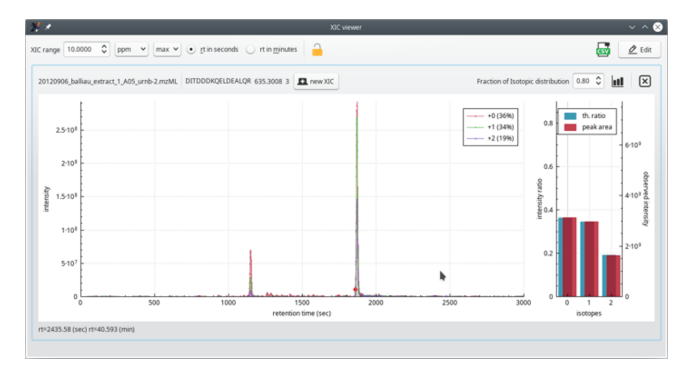
The notion of *extracted ion current* chromatogram is best explained by describing the computation that yields that chromatogram.

The user defines the m/z value for which the chromatogram is to be determined. The program iterates in each MS spectrum (that is, full scan spectrum) and looks if an ion by that m/z value was encountered. If so, a variable holding the cumulated intensity of that ion is incremented for the retention time at which the mass spectrum was acquired. For example, if m/z value 1254.25 is searched for, and an ion of that m/z value is found in the mass spectrum acquired at retention time 2.5 min, then a tuple variable is stored like this: (2.5, intensity). Then, another mass peak by that m/z value is found in mass spectrum acquired at retention time 47 min, for which another tuple is created: (47, intensity).

If the data are from ion mobility—mass spectrometry (IM-MS) experiments, there might be a large number of spectra acquired at a given retention time. For example, data from the *Waters SynaptzTM* instrument have 200 spectra acquired for any given retention time value (the spectra are drift-related spectra). In *Bruker timsTOFTM* data, there are more than 700 spectra acquired at any given retention time. Thus,

the searched m/z value might be found more than once for a retention time value. In this case, the tuple's intensity value is incremented by the intensity of the new peak of the m/z value at that specific retention time value.

When the program has finished iterating in all the mass spectra of the acquisition, it plots the XIC chromatogram as intensity=f(retention time). This is the reason why it is considered a chromatogram.



The extracted ion current (XIC) chromatogram viewer is useful to scrutinize the mass data at the very origin of a PSM. It is routinely used to ensure that the PSM is faithful. If not, the corresponding peptide can be unchecked from the peptide identifications list table view. As a response, the protein inference process is run anew.

FIGURE 4.10: THE EXTRACTED ION CURRENT (XIC) CHROMATOGRAM VIEWER WINDOW

The XIC viewer window displays the "guts" of the of MS spectrum of the precursor ion that was fragmented and that yielded a PSM. The XIC chromatogram (left plot panel) is actually a set of XIC chromatograms that are superimposed in the plot widget (see FIGURE 4.11, "THE EXTRACTED ION CURRENT (XIC) CHROMATOGRAM VIEWER WINDOW (ZOOMED VIEW)"). One of the traces (legend +o) is for the first peak of the isotopic cluster of the searched ion; the second trace (legend +i) is for the second peak of the isotopic cluster. Likewise for the third trace. In the typical informatics-oriented style of numbering, the first isotopic peak (only light isotopes enter in the composition of the peptidic ion), is "isotope o"; the second isotopic peak (one light isotope is substituted with a heavy one) is "isotope 1" and, finally, the third isotopic peak (two light isotopes were replaced by heavy ones) is "isotope 2". The right panel is a bar plot showing —for each one of the isotopes— a comparison between the experimental peak area and the computed probability of the corresponding isotope peak. In the example, the match between the experimental and the theoretical cluster shape is perfect. This scrutinization of the data is very useful when one wants to double-check the quality of a protein identification on the basis of a given PSM.



Note

The theoretical isotopic cluster peaks are calculated using the formula of the peptide that has been identified in the PSM for which the XIC chromatogram is being requested.



Zoomed in view of the three XIC chromatogram plots in the left hand side plot widget.

FIGURE 4.11: THE EXTRACTED ION CURRENT (XIC) CHROMATOGRAM VIEWER WINDOW (ZOOMED VIEW)

Another interesting bit of information is the *Fraction of Isotopic distribution* (spin box widget top right corner of the window). This one needs some background. When one has a peptide formula and the peptidic ion charge, one can calculate the theoretical isotopic cluster corresponding to that specific ion. The calculation is CPU-intensive and sometimes one would like to limit its duration. This is possible by indicating that one is interested, for example, in only the 80 % of the total isotopic peaks that one would effectively find (even in minute amouts) in nature. This value tells exactly that. The calculation displayed in the window, encompassing only 80 % of the whole natural span of the isotopic cluster, yields a calculated cluster made of only three isotopic peaks. If the user had set the value to 99 %, then, most probably, numerous other isotopic peaks of very low intensity would have been calculated on the right hand side of the isotopic cluster (heavier ions because more heavy isotopes are included in the computation).

Remember to click onto the "*Histogram plot*" button next to the spin box widget for the new *Fraction of Isotopic distribution* value to take effect.

As for the previous MS/MS spectrum plot, to zoom in/out regions of the XIC chromatogram plot widget, hover the mouse cursor over the region of interest and rotate the mouse wheel.

4.3 HANDLING PHOSPHO-PROTEOMICS DATA

i2MassChroQ is able to cope with phospho-peptides. The mass spectrometric data are acquired exactly as usual with the mass spectrometer, but the sample preparation goes along theses steps:

- Separate digestion of the samples (when there are more than one);
- Labeling of the peptides, each sample gets a different label;
- Pool of the whole set of peptides into a single mixture;
- Separation of the peptides on a strong cation exchange (SCX) resin, collection of the fractions;
- Phospho-peptide enrichment using IMAC¹ for each SCX fraction. The SCX fraction is loaded onto the IMAC resin and, following a wash step, the phospho-peptides are eluted (pH-based elution). There is thus a one-to-one relation between a SCX fraction and an IMAC-based purification fraction.
- Mass spectrometric analysis of each IMAC-based phospho-peptide-enriched fraction.

X!Tandem needs to be configured in such a manner that it can generate all the theoretical peptides (and fragments) that might bear the phosphoryl group. This process is described in the section below.

¹ Immobilized-metal affinity chromatography.

5 EXPLORING POST-TRANSLATIONAL MODIFICATION DATA

This chapter describes in detail all the steps that the user accomplishes in their post-translational modification (PTM) data exploration session.

5.1 SETTING THE X! Tandem RUN PRESETS FOR PHOSPHO-PROTEOMICS

The very first step in starting a phospho-proteomics-based protein identification run is to configure the run so that the database search engine can model phopshorylated peptides on the basis of the sequence of the peptides in the database. That configuration step is started as described in FIGURE 5.1, "X!TANDEM SETTINGS WINDOW FOR A PHOSPHO-PROTEOMICS PROJECT".



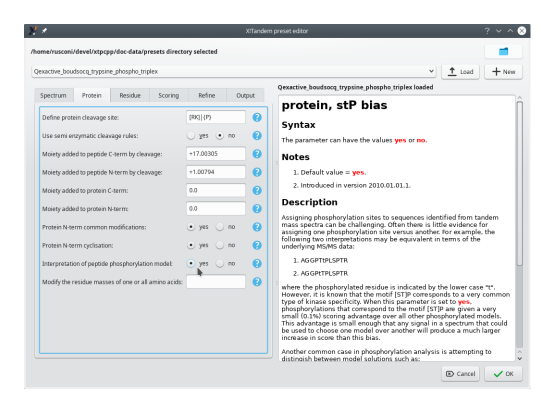
The *Choose presets* option in this window allows the user to select an *X*.*Tandem* presets file to suit the protein identification run. In this example, the chosen presets file contains a number of configuration bits specific for a phospho-proteomics project.

FIGURE 5.1: XITANDEM SETTINGS WINDOW FOR A PHOSPHO-PROTEOMICS PROJECT

The configuration of *X*!*Tandem* needs to be performed by using the presets method, described in **SECTION 3.3**, **"SETTING THE X**!**TANDEM RUN PRESETS"** and following sections. The *Edit* button next to the drop down list allows one to edit the presets that configure the handling of the database by the *X*!*Tandem* database search en-

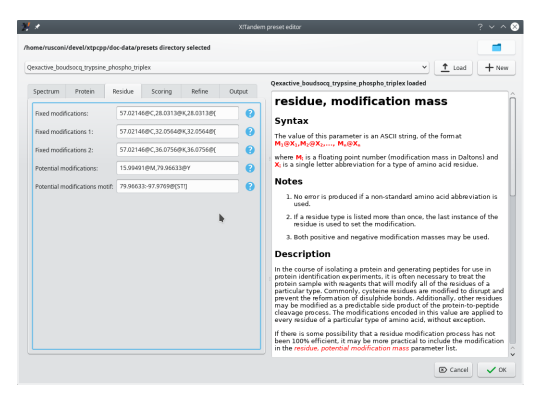
gine. The window that opens up upon clicking onto that button has two tabs that require the user's attention, as shown in Figure 5.2, "Setting the project to be a phospho-proteomics project" and Figure 5.3, "Con-Figuring the phosphorylated residues".

As described in SECTION 3.3.1, "LOADING EXISTING PRESETS CONFIGURATIONS FROM FILE", it is possible to load existing presets in case these were defined already and might be reused for a repeated data exploration session on the same data set.



The major bit that need the user's attention in the *Preset editor* window's *Protein* tab is the *Interpretation of peptide phosphorylation model* to *yes*. The question mark icon on the side of that configuration option displays explanatory text on the right hand side of the window.

FIGURE 5.2: SETTING THE PROJECT TO BE A PHOSPHO-PROTEOMICS PROJECT



The phosphorylation events are not *Fixed modifications* because it is not possible to predict that they will occur systematically. This is the reason why the phosphorylation events are configured as *Potential modifications*.

FIGURE 5.3: CONFIGURING THE PHOSPHORYLATED RESIDUES

FIGURE 5.3, "CONFIGURING THE PHOSPHORYLATED RESIDUES" shows how to configure the potential phosphorylation of selected residues (tyrosine, threonine and serine). Setting the *X!Tandem* parameters for phosphoproteomics analyses involves specifying the mass difference between unmodified and modified residues and the nature of the residue that might bear the modification. There are two different settings available:

- *Potential modifications*: the *Y* residue might be phosphorylated, with a net mass increment of *79.96633* Da.
- *Potential modifications motif*: the S and T residues might be phosphorylated (net mass increment of *79.96633* Da) or be subject to a neutral phosphoric acid loss (net mass loss of *97.9769* Da).

The reason why the potential phosphorylation of tyrosine (Y) is not mentioned along with the S and T modifications in the motif setting is that phosphorylated tyrosine residues do not suffer from phosphoric acid neutral loss upon collisionally activated dissociation (CID). Phosphorylated serine and threonine residues are readily dephosphorylated upon CID. Hence, the requirement to configure both the phosphorylation and the dephosphorylation events as a PROSITE motif (see the question mark help).



Tip

The loss of a phosphoric acid molecule *(not ion)* is called a "neutral loss". By essence, the lost molecule cannot be detected, because it bears no charge. However, the search software may detect that there might be a negative mass delta between calculated fragments bearing a phoshoryl group and the measured mass of product ions. In this eventuality, the software may deduce that the fragment was phosphorylated before the fragmentation occurred.

The mass spectrometer might be configured to monitor neutral phosphoric acid loss, or not. In some instruments, that workflow is not available; however, in these instruments a higher energy collisional dissociation¹ process elicits two fragmentation events: loss of a phosphoric acid molecule and peptide backbone dissociation. In this case, the database searching engine (*X!Tandem*, for us) is instructed to monitor the loss of phosphoric acid (that is, a neutral loss) on the product ions of the y ion series. In the best cases (best sequence coverage by the product ions), it is thus possible to locate the phosphoryl group on the peptide.

CAUTION: PHOSPHO-PROTEOMICS PROJECTS OFTEN INVOLVE LABEL-BASED QUANTIFICATION

The *Residue* tab of the *X!Tandem preset editor* window shown above lists a number of fixed modifications that need an explanation, because we'll find them later in other figures. The 57.02146@C modification is the carbamidomethylation of cysteine residues. The various 28, 32 and 36 mass increments on lysine residues are the di-methylation modifications with (32, 36) or without (28) heavy isotopes². The exact same mass increments labelled with the "@[" notation are the same modifications occurring on the N-terminus of the peptide.

Once all the settings have been validated, click the *Run* button, in the same manner as described at Section 3.3.4, "Running a properly configured X!Tandem process", to actually start the database search process.

I In Orbitrap analyzer-based instrument, HCD stands for "higher-energy C-trap dissociation". However, a more generic term is oft-used: "higher energy collisional dissociation".

² Boersema et al. 2009. Multiplex peptide stable isotope dimethyl labeling for quantitative proteomics. Nature Protocols.

5.2 LOADING THE PROTEIN IDENTIFICATION RESULTS

The loading of the protein identification results for a phospho-proteomics project occurs in an identical fashion as for a non phospho-proteomics project, as described in Section 3.4, "Loading the Protein Identification Results". The main program window shows the same *Identifications* tab as described before (see Figure 5.4, "The MAIN WINDOW'S IDENTIFICATIONS TAB").

2/ ×	new_project - X!Tand	emPipeline 0.4	1.43 🗸	$\sim \otimes$
F <u>i</u> le <u>E</u> dit 😯				
_				
Identifications	Filter parameters	FDR	Mass precision	
	View protein list	View MS	identification list	
	Number of samples	6		
	Number of groups	2738		
	Number of subgroups	3200		
Numb	er of grouped proteins	: 5809		
Number of gro	ouped distinct peptides	27537		
View PTM isl	ands			
	~			
computing PTM isla	ands finished			

The *Identifications* tab inside of the main program window after loading a PTMs-based project's identification results files. The *View protein list* and *View MS identification list* buttons perform exactly as described earlier. In order to trigger the PTMs-based data exploration session, click the *View PTM islands* button.

FIGURE 5.4: THE MAIN WINDOW'S IDENTIFICATIONS TAB

When *i2MassChroQ* has done loading all the results, the *Protein list* window opens up as it usually does. This window, however, is not devoted to the exploration of phosho-proteomics data. In order to start the exploration of phospho-proteomics data, it is necessary to display a window that lists all the post-translational modification islands ("PTM islands"). That window shows up when the user clicks onto the *View PTM islands* button shown in **FIGURE 5.4, "The MAIN WINDOW'S IDENTIFICATIONS TAB**".

5.3 EXPLORING PTM ISLANDS IDENTIFICATION DATA

The workflow involved in the scrutiny of phospho-proteomics data is comparable to that for a conventional proteomis project, as described in SECTION 4.1, "THE PROTEIN LIST WINDOW". There are differences, however, both in terminology and in the kind of data presented to the user that require a detailed review, performed in

the following sections. The first step is to show the PTM islands, which is the step analogous to showing the identified proteins. The second step is to look into a given PTM island by scrutinizing the PTM peptides that make it; this step is analogous to showing the peptide list for a given protein.

5.3.1 THE PTM ISLANDS LIST WINDOW

The *PTM island list* window in FIGURE 5.5, "PTM ISLANDS LIST WINDOW" displays, in a table view, all the PTM islands that were identified (see Section 2.2.6.2, "PROTEIN IDENTIFICATION IN PHOSPHO-PROTEOMICS PROJECTS").

hecked	 PTM island ID 	accession	: description	ptm positions	spectrum	E PTM spectrum	sequence	i multi PTM	i start	: length
	ptmc314a1a1	AT2G23120.1	Late embryogenesis abundant protein%2C group 6	67	6	1	1	0	45	37
	ptma1a1a1	AT1G30230.1	translation elongation factor EF1B/ribosomal protein S6 family	91	3	3	1	0	66	33
	ptma1a2a1	AT1G30230.1	translation elongation factor EF1B/ribosomal protein S6 family	117	207	3	3	0	99	29
	ptma1a1a2	AT1G30230.2	translation elongation factor EF1B/ribosomal protein S6 family	91	3	3	1	0	66	33
	ptma1a2a2	AT1G30230.2	translation elongation factor EF1B/ribosomal protein S6 family	117	207	3	3	0	99	29
	ptmb19a1a1	AT5G24490.1	30S ribosomal protein Chr5:8365690-8367178 FORWARD	215	37	2	3	0	197	36
	ptmb67a1a1	AT1G06410.1	trehalose-phosphatase/synthase 7 Chr1:1955413-1958153	57	16	16	1	0	5	18
	ptmb67a1a2	AT1G06410.2	trehalose-phosphatase/synthase 7 Chr1:1955413-1958153	57	16	16	1	0	5	18
	ptmb67a1a3	AT1G06410.3	trehalose-phosphatase/synthase 7 Chr1:1955413-1958153	57	16	16	1	0	5	18
	ptmc245a1a1	AT5G56010.1	heat shock protein 81-3 Chr5:22681410-22683911 FORWARD	213 219	8	8	2	0	213	21
	ptmc245a3a2	AT5G56010.1	heat shock protein 81-3 Chr5:22681410-22683911 FORWARD	286 288	5	1	1	0	286	13
	ptmc245a1a2	AT5G56030.1	heat shock protein 81-2 Chr5:22686923-22689433 FORWARD	213 219	8	8	2	0	213	21
	ptmc245a3a3	AT5G56030.1	heat shock protein 81-2 Chr5:22686923-22689433 FORWARD	286 288	5	1	1	0	286	13
	ptmc800a1a1	AT3G10730.1	SAD1/UNC-84 domain protein 2 Chr3:3358556-3360522 REVERSE	63 65	2	2	2	0	56	18
	ptmc800a2a1	AT3G10730.1	SAD1/UNC-84 domain protein 2 Chr3:3358556-3360522 REVERSE	77	1	1	1	0	75	15
	ptmc527a1a1	AT5G11040.1	TRS120 Chr5:3495332-3500610 FORWARD LENGTH=1186	971	4	4	1	0	966	11
	ptmb95a1a1	AT1G48920.1	nucleolin like 1 Chr1:18098186-18101422 FORWARD LENGTH=55	96 97 98	10	2	3	0	84	23
	ptmb95a2a1	AT1G48920.1	nucleolin like 1 Chr1:18098186-18101422 FORWARD LENGTH=55	125 126 130 131	13	3	2	0	119	21
	ptmb95a3a1	AT1G48920.1	nucleolin like 1 Chr1:18098186-18101422 FORWARD LENGTH=55	186 187 191 195	3	1	1	0	183	21

The *PTM island list* table view has many columns that characterize each PTM island, as described in detail in the text below.

FIGURE 5.5: PTM ISLANDS LIST WINDOW

In the table view of all the PTM islands, each row corresponds to an island. It must be noted that multiple rows may appear as identical islands. This is not exactly true because, while the PTM islands appear to be exactly the same, these have been identified in proteins that are listed in the proteins database under different accession numbers (see the *accession* column of the table view).

The columns in the table view hold post-translational modification-specific data described below.

- *Checked*: if checked, the corresponding PTM island will be taken into account;
- *PTM island ID*: unambiguous identifier for the PTM island. The nomenclature is important and follows a precise syntax according to this scheme:
 - ptm<letter1><number><letter2><number><letter3><number>, with the following meaning:

³ The data presented in the examples below come from an experiment published in T. Y. Delormel *et al.* 2022. In vivo identification of putative CPK5 substrates in *Arabidopsis thaliana*. *Plant Science*. DOI: <u>HTTPS://DOI.ORG/10.1016/J.PLANTSCI.2021.11121</u> .

- <letter1><number>: identifies the group of proteins that share PTM islands;
- <letter2><number>: identifies the PTM island in the above group;
- <letter3><number>: identifies the accession number of the protein in the group above.
- *accession*: the accession number of the protein in the protein database file;
- *description*: the description of the protein in the protein database file;
- *ptm positions*: the number of positions in the PTM island that bear a post-translational modification;
- *spectrum*: number of distinct spectra that allowed defining this PTM island;
- *PTM spectrum*: number of distinct spectra that allowed the identification of more than one searched post-translational modifications;
- *sequence*: number of unique peptide sequences found in this PTM island;
- *multi PTM*: number of spectra that allowed the identification of more than one post-translational modification site;
- *start*: the position of the PTM island on the protein sequence (the first residue of the PTM island);
- *length*: the number of residues that encompass the PTM island.

In the same way as for the *Protein list* window's table view of the identified proteins, the cells of the table are "active": clicking onto any cell triggers the opening of a window that provides details about the corresponding PTM island.

5.3.2 Delving Inside the PTM Island Identification Data

The protein identifications list table view, as pictured in FIGURE 5.5, "PTM ISLANDS LIST WINDOW" is actually an active matrix where the user can easily trigger the exposition of the data that yielded any PTM island identification element of the table. This is simply done by clicking onto any cell of the table at the row matching the PTM island for which scrutiny of the data is requested.

Depending on the column at which the mouse click happens, there might be two different windows showing up:

• Clicking onto a cell in either the *accession* or *description* column opens the *Protein details* window shown in FIGURE 5.6, "PROTEIN DETAILS WINDOW".

2019_Boudsocq_146_PHOS - Protein details	~ ~ × `
AT3G56150.1 AT3G56150 eukaryotic translation initiation factor 3C Chr3:20833790-20836820 REVERSE	valid contaminant decoy
LENGTH=900 201606 MTSRFFTQVGSESEDESDYEVEVNEVQNDDVNNR <mark>YLQSGSEDDDDTDTKR</mark> VVKPAKDKRFEEMTYTVD ENFDKVNKQLEKVMRITEAVKPPTLYIKTLVMLEDFLNEALANKEAKKKMSTSNSKALNSMKQKLKK EVEEEKQPEDDDDDDDDDDEVEDDDDSSIDGPTVDPGSDVDEPTDNLTWEKMLSKKDKLLEKLMNKDF VAARGKKGTARFELVDQLTHLTKIAKTPAQKLEILFSVISAQFDVNPCLSGMMPINVWKKCVLMMLTI VEPDENETSKPTDYDGKIRVWGNLVAFLERVDTEFFKSLQCIDPHTREVVERLRDEPMFLALAQNIQU LRRVEAIYYKPQEVYDAMRKLAELVEEEETEEAKEESGPTSFIVVPEVVPRKFTFPESSRAMMDII AMLCDINHHALMDNFVTARDLLLMSHLQDNIQHMDISTQILFNRTMAQLGLCAFRAGMITESHSCLSE SQSRYHEKTPEQERMERRRQMPYHMHLNLELLEAVHLICAMLLEVPNMAANSHDAKRRVISKNFRRLI RDHVMAATRALTKGDFQKAFEVLNSLEVWRLLKNRDSILDMVKDRKEEALRTYLFTYSSYESLSLI IVSKMMINEELHASWDQPTRCIVFHEVQHSRLQSLAFQLTEKLSILAESNERAMESRTGGGGLDLSSF	NIKLYEDDINKYREAP KEITNDWVNKKFKEI ILDILVKYSNIVVDDT JYFERMGDFKAAAKVA LVSLIYRNGDERTKAR ELYSGQRVRELLAQGV LEISERQAFTAPPENV JQLAKMFDVSEPQVHS RRRDNNQDYAGAASGG
coverage 15.8889 % MW 102.884 kDa E-value 1.13437e-95 (log10: -94.9452)	get PDF

This window provides details about the protein identified as bearing one or more post-translation modification(s). The peptides that were matched as PSMs are highlighted in yellow. The $AT_3G_56_{150}$ provides a link to an external resource⁴. The other informational data bits are self-explanatory.

FIGURE 5.6: PROTEIN DETAILS WINDOW

• Clicking onto a cell in the *PTM island ID* column opens the *PTM peptide list* window for that specific island. That window is displayed in both figures FIGURE 5.7, "PTM PEPTIDES LIST WINDOW (FIRST COLUMNS)" and FIGURE 5.8, "PTM PEPTIDES LIST WINDOW (LAST COLUMNS)" in the following section (Section 5.4, "The PTM PEPTIDES LIST WINDOW").

5.4 The PTM Peptides List Window

Each PTM island is essentially defined by a set of related peptides that sport one or more post-translational modifications. The exploration of the PTM peptides thus involves looking into the different peptides of a PTM island. A number of characteristics of these peptides are described in the following text.

⁴ Here, the page https://www.arabidopsis.org/servlets/TairObject?type=locus&name=AT3G56150 🗗 opens in the browser.

peptide ID	sample	: scan	: RT	: charge	sequence
epb87b25	20160107_Boudsocq_146_Tx1F02	5004	1237.14	3	YLQSGSEDDDDTDTKR
epb87b25	20160107_Boudsocq_146_Tx1F03	4566	1141.31	3	YLQSGSEDDDDTDTKR
epb87b25	20160107_Boudsocq_146_Tx1F03	4809	1185.39	3	YLQSGSEDDDDTDTKR
epb87b25	20160107_Boudsocq_146_Tx1F03	4915	1205.11	2	YLQSGSEDDDDTDTKR
pepb87b25	20160107_Boudsocq_146_Tx1F03	5032	1226.03	3	YLQSGSEDDDDTDTKR
pepb87b25	20160107_Boudsocq_146_Tx1F03	5149	1246.95	2	YLQSGSEDDDDTDTKR
pepb87b25	20160107_Boudsocq_146_Tx1F03	5281	1269.35	3	YLQSGSEDDDDTDTKR
pepb87b25	20160107_Boudsocq_146_Tx1F04	4346	1198.67	3	YLQSGSEDDDDTDTKR
pepb87b25	20160107_Boudsocq_146_Tx1F05	4670	1237.95	3	YLQSGSEDDDDTDTKR
pepb87b24	20160107_Boudsocq_146_Tx1F02	4133	1106.29	3	YLQ <mark>S</mark> GSEDDDDTDTKR
pepb87b24	20160107_Boudsocq_146_Tx1F02	4303	1130.7	2	YLQSGSEDDDDTDTKR
pepb87b24	20160107_Boudsocq_146_Tx1F02	4443	1151.22	3	YLQ <mark>S</mark> GSEDDDDTDTKR
epb87b23	20160107 Poudcoca 146 Tv1E02	4472	1155.06	ο.	YLQ <mark>S</mark> GSEDDDDTDTKR
pepb87b23	MOD:00552)LQS(MOD:00696)GSEDE				YLQ <mark>S</mark> GSEDDDDTDTKR
vepb87b23	MOD:00552)LQSGS(MOD:00696)EDD	DDTDTK(N	10D:00552)R	(6) 4	YLQ <mark>S</mark> GSEDDDDTDTKR
pepb87b22	20160107_Boudsocq_146_Tx1F02	4174	1112.4	3	YLQ <mark>S</mark> GSEDDDDTDTKR
pepb87b22	20160107_Boudsocq_146_Tx1F02	4174	1112.4	3	YLQSGSEDDDDTDTKR
pepb87b22	20160107_Boudsocq_146_Tx1F02	4306	1131.24	2	YLQ <mark>S</mark> GSEDDDDTDTKR
pepb87b22	20160107_Boudsocq_146_Tx1F02	4306	1131.24	2	YLQSG <mark>S</mark> EDDDDTDTKR
pepb87b22	20160107_Boudsocq_146_Tx1F02	4454	1153	3	YLQ <mark>S</mark> GSEDDDDTDTKR

Every PTM island has, associated to it, a list of post-translationally modified peptides. This figure illustrates the first columns of the table view that lists all the peptides making a PTM island. The contextual menu visible near peptide ID *pepb87b23* will be detailed later (SECTION 5.4.1.2, "AMBIGUITIES ON THE POST-TRANSLATIONAL MODIFICATION SITES").

FIGURE 5.7: PTM PEPTIDES LIST WINDOW (FIRST COLUMNS)

modifs	: start	i length	top Evalue	theoretical MH+	delta MH+	top hyperscore	top PTM positions	observed PTM positions
Y:28.03 45:79.97 65:79.97 15K:28.03	35	16	2.7e-09	2060.77	0.000265847	36.5	46	4 6
Y:28.03 45:79.97 65:79.97 15K:28.03	35	16	0.00036	2060.77	0.000404847	21.6	46	4 6
Y:28.03 45:79.97 65:79.97 15K:28.03	35	16	1.4e-10	2060.77	0.00327185	30	4 6	4 6
Y:28.03 45:79.97 65:79.97 15K:28.03	35	16	0.024	2060.77	0.00140985	17.6	4 6	4 6
Y:28.03 45:79.97 65:79.97 15K:28.03	35	16	5.5e-11	2060.77	0.00157785	39.6	46	4 6
Y:28.03 4S:79.97 6S:79.97 15K:28.03	35	16	0.00035	2060.77	0.00179385	24.4	4 6	4 6
IY:28.03 45:79.97 65:79.97 15K:28.03	35	16	1.1e-10	2060.77	7.38466e-05	37.5	4 6	4 6
Y:28.03 4S:79.97 6S:79.97 15K:28.03	35	16	0.0018	2060.77	0.00150185	17.9	4 6	4 6
Y:28.03 4S:79.97 6S:79.97 15K:28.03	35	16	4e-09	2060.77	0.00209085	36.3	4 6	4 6
Y:36.08 4S:79.97 15K:36.08	35	16	6.6e-07	1996.9	0.000649549	32.4	4	4
Y:36.08 6S:79.97 15K:36.08	35	16	1.5e-05	1996.9	0.00314155	25.6	6	6
Y:36.08 4S:79.97 15K:36.08	35	16	1.8e-07	1996.9	-0.00163045	36.6	4	4
IY:32.06 4S:79.97 15K:32.06	35	16	3.5e-05	1988.86	0.00194488	32.9	4	4 6
IY:32.06 4S:79.97 15K:32.06	35	16	2.2e-12	1988.86	0.00276688	47.2	4	4
1Y:32.06 4S:79.97 15K:32.06	35	16	3.9e-05	1988.86	0.00415188	24.3	4	4 6
Y:28.03 4S:79.97 15K:28.03	35	16	4.2e-05	1980.81	0.000237847	26.8	4	4
1Y:28.03 6S:79.97 15K:28.03	35	16	4.2e-05	1980.81	0.000237847	26.8	6	6
IY:28.03 4S:79.97 15K:28.03	35	16	1.5e-08	1980.81	0.00133285	36	4	4
Y:28.03 6S:79.97 15K:28.03	35	16	1.5e-08	1980.81	0.00133285	36	6	6
IY:28.03 45:79.97 15K:28.03	35	16	3.8e-07	1980.81	0.000709847	36.3	4	4

Every PTM island has, associated to it, a list of post-translationally modified peptides. This figure illustrates the last columns of the table view that lists all the peptides making a PTM island.

FIGURE 5.8: PTM PEPTIDES LIST WINDOW (LAST COLUMNS)

The *PTM peptide list* window contains a large set of peptide characteristics organized in a number of columns, as described below:

- *peptide ID*: the unambiguous identity of the peptide;
- *sample*: the file name of the sample in which the peptide was found and sequenced;
- scan: the scan number of the MS/MS spectrum in which the precursor peptidic ion was fragmented;
- *RT*: the retention time at which the peptide eluted;
- *charge*: the charge of the peptidic ion;
- *sequence*: the sequence of the peptide. Note that the residues that do (or might) bear a post-translational modification are printed in red color;
- *modifs*: a semi-column-separated list of modified positions. The modification is identified by the net mass change that occurs upon the chemical modification. The *1Y:32.06 4S:79.97 15K:32.06* example indicates that the tyrosine at position 1 is dimethylated, the serine at position 4 is phosphorylated and that the lysine at position 15 is methylated (the dimethyl modification was commented above).

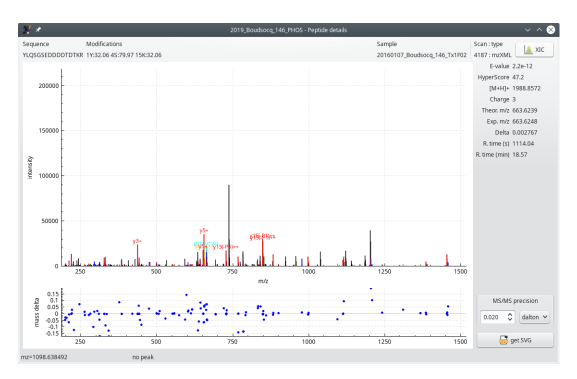
- *start*: the position of the peptide's first residue in the protein sequence;
- *length*: the number of residues in the peptide;
- *top Evalue*: best Evalue from those calculated for different peptide spectrum matches that occurred in a single fragmentation scan (see Section 5.4.1.2, "AMBIGUITIES ON THE POST-TRANSLATIONAL MODIFICA-TION SITES" below for a thorough description of this situation);
- *theoretical MH*+: the calculated mass of the [M+H]⁺ peptidic ion;
- *delta MH*+: the difference between the measured and the calculated [M+H]⁺ masses;
- *top hyperscore*: best Hyperscore from those calculated for different peptide spectrum matches that occurred in a single fragmentation scan (see Section 5.4.1.2, "AMBIGUITIES ON THE POST-TRANSLATIONAL MODIFICATION SITES" below for a thorough description of this situation);
- *top PTM positions*: positions of modified residues for the peptide having the best Evalue (see above) for the current scan;
- observed PTM positions: space-separated list of all the modified positions found for the current scan.

5.4.1 Delving Inside the PTM Peptide Identification Data

The *PTM peptide list* table view (FIGURE 5.7, "PTM PEPTIDES LIST WINDOW (FIRST COLUMNS)") is actually an active matrix where the user can easily trigger the exposition o fthe data that yielded any PTM peptide identification element of the table. This is simply done by clicking onto any cell of the table at the row matching the peptide for which scrutiny of the data is requested.

5.4.1.1 The Peptide Details Window

Clicking a cell in the *peptide ID* column opens the *Peptide details* window shown in FIGURE 5.9, "PEPTIDE DETAILS WINDOW". Notice how this window is similar to the one described for conventional non PTM-based projects at SECTION 4.2.3.1, "THE PEPTIDE DETAILS WINDOW".



This window displays a large amount of informational data bits that characterize the MS/MS spectrum *vs* peptide match (PSM) for the peptide ID that was clicked in the *PTM peptide list* table view window. Almost all the information data bits shown in this figure are self-explanatory.

FIGURE 5.9: PEPTIDE DETAILS WINDOW



Tip

The nomenclature of the product ions in the MS/MS spectrum shown in the figure above is simple: when the ion under a given MS/MS spectrum peak is the result of a neutral loss of phosphoric acid, the ion is labelled "yx(-Pz)", with x being the index of the y ion, and (-Pz) indicating the loss of z phosphoric acid neutral molecule(s).

When ions actually bear any number of post-translational modifications, these are not listed along with the ion series (b or y) and index text because that would quickly become unwieldy, from a graphical point of view.

5.4.1.2 Ambiguities on the post-translational modification sites

The *PTM peptide list* table view in FIGURE 5.7, "PTM PEPTIDES LIST WINDOW (FIRST COLUMNS)" lists the *pep-b87b23* peptide ID thrice because that peptide was identified in three different MS/MS scans (see the *scan* column values for the three rows). What is interesting with these three rows, is that the peptide sequence showing the modification position(s) varies from a row to the other. In one row, the sequence only shows a single red-colored serine residue. That serine was positively identified as a phosphorylated residue. In the other two rows,

one serine is red-colored and the other is orange-colored. For these two scans, the position of the phosphorylated residue is not ascertained: the MS/MS data only tell that it is possible that one or the other serine residue be phosphorylated (but not both).

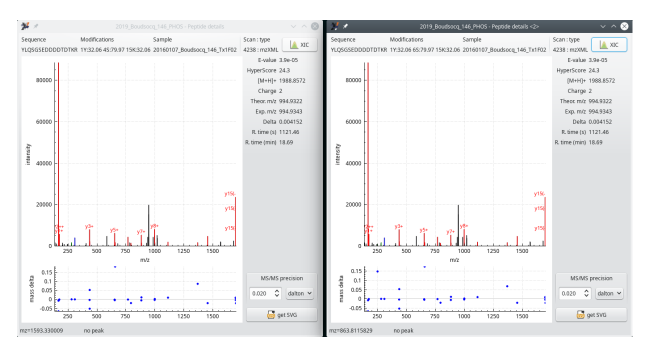


Note

The coloring of these two serine residues is arbitrary in this case: since the PSMs that yielded this same sequence are absolutely identical from a score point of view (when opening the peptide details window, one can check that the Evalues are identical for the two PSMs), the software labels in red the first modification position and in orange all the remaining ones.

When clicking any cell of any one of the two rows where there is an ambiguity over the location of the phosphorylation event, *i2MassChroQ* shows a contextual menu that displays the possible phosphorylation positions. The user can thus select one or the other of the menu items to display the details of the corresponding peptide.

The ambiguity about the phosphorylation site above, on *Peptide ID pepb87b23*, is intereresting. Indeed, when the user selects the first item of the contextual menu and then (by keeping the Ctrl key pressed) selects the second item of that menu, the windows that open up show exactly the same informational data bits about the MS/MS scan: the two PSMs were calculated by the search engine on the basis of the very same MS/MS scan (FIGURE 5.10, "Two MASS SPECTRUM VS PEPTIDE MATCHES (PSM) IN A SINGLE MS/MS scan"). This is one reason why both positions could not be ascertained: the engine says that one position is plausible (with a pretty low Evalue) and that the other position is equally plausible (with the exact same low Evalue).



This figure shows the two *Peptide details* windows that are opened when the user clicks onto the first and the the second menu items of the contextual menu shown in the *PTM peptide list* window shown in FIGURE 5.7, "PTM **PEPTIDES LIST WINDOW (FIRST COLUMNS)**". In both windows, the scan number is the same (4238), demonstrating that both PSMs were computed from the same MS/MS spectrum acquisition. Further, the same pretty low E-value should hint at a low reliability of the phosphorylation site identification.

FIGURE 5.10: TWO MASS SPECTRUM VS PEPTIDE MATCHES (PSM) IN A SINGLE MS/MS SCAN

5.4.1.3 The XIC Viewer Window for the PTM Peptide Details

One interesting feature of the *Peptide details* window, is the *XIC* button (top right) that triggers the calculation of an extracted ion current chromatogram, as pictured in FIGURE 5.11, "THE EXTRACTED ION CURRENT (XIC) CHROMATOGRAM VIEWER WINDOW". Although very similar to the window described at Section 4.2.3.2, "THE XIC VIEWER WINDOW FOR THE PEPTIDE DETAILS", this phospho-proteomics-specific version of the *XIC viewer* has specific informational data bits described below.



The extracted ion current (XIC) chromatogram viewer shows the peptide sequence interspersed with post-translational modifications data. The modifications are listed as PSIMOD OBO accession number (MOD:xxxxx) text elements⁵.

FIGURE 5.11: THE EXTRACTED ION CURRENT (XIC) CHROMATOGRAM VIEWER WINDOW

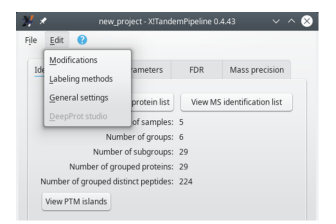
The phospho-proteomics-specific informational data bits provided in the *XIC viewer* window (see Figure 5.11, "The EXTRACTED ION CURRENT (XIC) CHROMATOGRAM VIEWER WINDOW") are located right below the top border of the plot frame.

⁵ Montecchi-Palazzi et al. 2008. PSI-MOD community standard for representation of protein modification data. Nature biotechnol.

6 Advanced Proteomics Configurations

This chapter describes in detail all the various advanced configurations that determine the workings of *i2Mass-ChroQ* in various contexts.

Some of the advanced proteomics configurations can only be carried over when protein identification results have been loaded in *i2MassChroQ*. Indeed, only at that point, does the menu of the main program window show the submenus of interest, as shown in FIGURE 6.1, "THE EDIT MENU SHOWING VARIOUS CONFIGURATION TASKS".



This menu becomes available in the main program window only when identification data have been loaded in *i2MassChroQ*

FIGURE 6.1: THE EDIT MENU SHOWING VARIOUS CONFIGURATION TASKS

6.1 Configuring Modifications

It is often required to specify the chemistry of a given chemical modification that might be performed (or spontaneously observed) on the proteins of a sample. For example, proteins in a sample often undergo a chemical modification of the cysteine residues in their reduced form to block them from reoxidizing into a sulphide bond. The reagent used is iodoacetamide and the reaction name is *carbamidomethylation*.

In order to be able to access the modifications configuration, select *Modifications* from the *Menu* menu. The window that opens up is shown in FIGURE 6.2, "MODIFICATIONS EDITOR WINDOW (FOLDED)". In that *Modifica-tions editor* window, the currently defined modifications are listed with no details about them.

¥ *	Modifications editor	~ ^ 😣
	Modification list	
[MOD:00397] iodoacetamide deriva	tized residue	
[MOD:01160] deaminated residue		
[MOD:00408] mono N-acetylated re	sidue	
[MOD:00719] L-methionine sulfoxid	e	

When the *Modifications editor* window is displayed, all the modifications data are folded.

FIGURE 6.2: MODIFICATIONS EDITOR WINDOW (FOLDED)

To start editing an existing modification, click the correponding item from the list, which opens up a *Selected modification* group box widget, as shown in FIGURE 6.3, "MODIFICATIONS EDITOR WINDOW (UNFOLDED)".

2/ *	Modifications editor	~ ^ 😣
	Modification list	
[MOD:00397] iodoacetamide derivati	zed residue	
[MOD:01160] deaminated residue		
[MOD:00408] mono N-acetylated res	idue	
[MOD:00719] L-methionine sulfoxide		
	Selected modification	
accession MOD:00397		
description iodoacetamide derivatize	d residue	
mass delta 57.0214637211		
	Replace by	
accession		browse
description		
mass delta		
replace		

This view shows the details of the modification item currently selected.

FIGURE 6.3: MODIFICATIONS EDITOR WINDOW (UNFOLDED)

The *Modifications editor* window displays the details of the modification currently selected in the *Modifications list*. The main informational data bits are the following:

- *accession*: the PSI-MOD¹ identificator. This value matches the id field of the corresponding PSI-MOD ontology record (see an example below);
- *description*: the description of the PSI-MOD entity. This value corresponds to the name field of the corresponding PSI-MOD ontology record;
- *mass delta*: the net mass change that results from a protein modification using this modification entity.



Note: A typical term definition from the PSI-MOD ontology

[Term]

```
id: MOD:00397
name: iodoacetamide derivatized residue
def: "A protein modification that is produced by reaction
with iodoacetamide, usually replacement of a reactive hydrogen
with a methylcarboxamido group."
[PubMed:11327326, PubMed:11510821, PubMed:12422359, Unimod:4]
subset: PSI-MOD-slim
synonym: "Carbamidomethyl" RELATED PSI-MS-label []
synonym: "Iodoacetamide derivative" RELATED Unimod-description []
xref: DiffAvg: "57.05"
xref: DiffFormula: "C 2 H 3 N 1 0 1"
xref: DiffMono: "57.021464"
xref: Formula: "none"
xref: MassAvg: "none"
xref: MassMono: "none"
xref: Origin: "X"
xref: Source: "artifact"
xref: TermSpec: "none"
xref: Unimod: "Unimod:4"
is_a: MOD:00848 ! reagent derivatized residue
```

If a change, that is, a replacement, is required in the modification, the first step is to click onto the *browse* button located in the *Replace by* group box, which opens the *PSI MOD selection* window shown in FIGURE 6.4, "PSI MOD SELECTION WINDOW".

¹ The PSI-MOD ontology is perusable at https://raw.githubusercontent.com/HUPO-PSI/psi-мod-CV/мaster/PSI-MOD.ово 🗖

PSI MC	DD selection ? ~ ^ (
	MOD:00397 iodoacetamide derivatized residue "A protein modification that is produced by reaction with iodoacetamide, usually replacement of a reactive hydrogen with a methylcarboxamido group."
MOD:01214 57.0215 iodoacetamide - site C MOD:01215 57.0215 iodoacetamide derivatized aspartic acid MOD:01216 57.0215 iodoacetamide derivatized glutamic acid	PubMed:11327326 PubMed:11510821 PubMed:12422359 Unimod:4 C 2 H 3 N 1 O 1 57.0215 X X Image: Cancel image

This window allows one to browse the PSI-MOD ontology on the basis of the modification net mass value.

FIGURE 6.4: PSI MOD SELECTION WINDOW

The window shows the various PSI-MOD-defined ontology terms that match the net mass value of the modification that was selected when the *browse* button was clicked.

It is possible to search for other PSI-MOD terms that match a different net mass change by modifying the mass value in the spin box widget that is found under the control of the *Mass filter* check box. The other spin box widget next to the first one allows one to enter the tolerance of the net mass change match. Available "units" for that tolerance are Dalton, ppm and res.

If the *Mass filter* check box is unchecked, the list beneath it displays *all* the PSI-MOD ontology terms.

Once the desired term has been found and selected, click the *OK* button. In order to effectively replace the old term with the new one, do not forget to click the *replace* button.

TIP: WHY BOTHERING WITH PSI-MOD ONTOLOGY MODIFICATION TERMS ?

It is beneficial to define chemical modifications based on the PSI-MOD ontology terms because the defined modification is then used as its code (for example *MOD:00397*) in the *X!Tandem* presets chemical modifications sections (see SECTION 3.3, "SETTING THE X!TANDEM RUN PRESETS"). Indeed, instead of only entering the net mass, defining a modification preset with the PSI-MOD ontology term code allows *i2MassChroQ* to compute a number of informative supplementary data. For example, when an isotopic cluster is simulated for a peptide that is modified with a PSI-MOD-documented modification, the formula of the modification recorded in the term is taken into account in the isotopic cluster calculation.

6.2 Configuring Labeling Methods

i2MassChroQ can handle stable isotope labeling methods. In order to list the available methods, select the *Labeling methods* menu item of the *Edit* menu in the main program window. The *Labeling methods editor* window that opens up is shown in FIGURE 6.5, "LABELING METHODS EDITOR WINDOW".

^ 😣
Ж
C

This window lists all the available labeling methods.

FIGURE 6.5: LABELING METHODS EDITOR WINDOW

Configuring a labeling method is as easy as selecting one item from the list shown in FIGURE 6.5, "LABELING METHODS EDITOR WINDOW" and clicking *OK*. If unsure about the selection, just click *Cancel*.



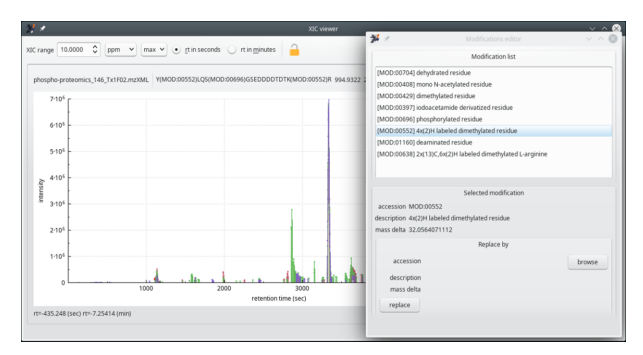
Once the user has clicked the *OK* button, it is no more possible to change the labeling method without recomputing the whole data set.

Using a labeling method involves a crucial step during the *X*!*Tandem* configuration, as described in SECTION 3.3, "SETTING THE X!TANDEM RUN PRESETS" and in CAUTION: PHOSPHO-PROTEOMICS PROJECTS OFTEN INVOLVE LA-BEL-BASED QUANTIFICATION. Whenever an experiment involves chemical labeling of any protein residue, the scientist is expected to configure the labeling modifications in the *X*!*Tandem* presets before running it to search the database, so that the engine models correctly-modified peptides.

When the *X*!*Tandem* database search engine has finished identifying peptides and proteins, the modified peptides are only tagged using the modifications' net masses. This is visible in the peptide details window shown in the upper left part of FIGURE 5.9, "PEPTIDE DETAILS WINDOW". There, one can see that the peptide is modified on its lysine 15 residue with one of the labels defined in the *Fixed modifications 1* field of the presets configuration window shown in FIGURE 5.3, "CONFIGURING THE PHOSPHORYLATED RESIDUES".

Selecting a labeling method, as described above, will allow *i2MassChroQ* to report residue modifications in a much more detailed and informative way. For example, the name of the chemical modification will be used instead of the net mass change due to that modification. For this to happen and the effects to be visible, *i2Mass-*

ChroQ goes back through all the identification process and makes sure that the modified peptides are tagged according to the modifications specified in the labeling method. The effects are visible in the *XIC viewer* window as shown in FIGURE 6.6, "CHEMICAL MODIFICATIONS ARE TAGGED WITH THEIR PSI-MOD CODE".



When the user selects a labeling method from the list of available chemical modifications, the chemically modified residue gets an associated tag with the PSI-MOD code. This is visible in this figure, at the top line of the *XIC viewer* window, where the last peptide's arginine residue is modified with MOD:00552, that corresponds to the 4x(2)H labeled dimethylated residue modification referenced in the *Modification editor* window on the right hand side of this figure.

FIGURE 6.6: CHEMICAL MODIFICATIONS ARE TAGGED WITH THEIR PSI-MOD CODE

6.3 THE *i2MassChroQ* GENERAL SETTINGS

There are general settings that might be configured for the whole *i2MassChroQ* program. These settings are available upon selecting the *General settings* menu of the *Edit* menu of the main program window. The window that opens up is shown in FIGURE 6.7, "GENERAL SETTINGS WINDOW".

27 ×	General settings	? ~ ^ 😣
М	S run XIC reader settings	
🔘 di <u>r</u> ect	• <u>b</u> uffered	
	R settings	
R binary path		
Select the R.exe par	th	
Rscript binary path		
Select the Rscript.e	xe path	
Temporary directory	3ruker timsTOF settings	
Select the tempora		
Origin of the MS/MS	data	
	=	
	✓ <u>О</u> К	⊘ <u>C</u> ancel

General settings might be configured in this window, as described below.

FIGURE 6.7: GENERAL SETTINGS WINDOW

The settings that can be configured are detailed below.

- *MS run XIC reader settings*: if *direct* is selected, the XIC chromatogram is computed by read the MS data right from the file; if *buffered* is selected, the XIC chromatogram is computed by using a cache mechanism that buffers data in the computer's memory.
- R settings:
 - *R binary path*: full path to the R program;
 - *Rscript binary path*: full path to the Rscript program;
- *Bruker timsTOF settings*: the fields below need filling-in if the data in the proteomics project originated in the Bruker's timsTOF line of instruments.
 - *Temporary directory*: optionally set the full path to the directory where the program stores temporary data;
 - Origin of the MS/MS data: if MGF is selected, the data are read from the Bruker-generated MGF file; if *TDF* is selected, the data are read from the Bruker native TDF file.



WARNING

If the user selects the MGF option above (because this is what they have), *i2MassChroQ* will not be able to extract XIC chromatograms, because the MGF file format does not contain MS data, it contains only MS/MS data. If having the native TDF files at hand (in the .d directory), the user is advised to select TDF.

7 *i2MassChroQ* and Quantitative Proteomics

This chapter describes in detail the way to prepare the work that will be carried over by the MassChroQ module.

7.1 INTERFACE TO THE *MassChroQ* QUANTITATIVE PROTEOMICS MODULE

While it is certainly possible to perform pretty thorough analyses by exploring data by way of peptide identification—protein inference scrutiny strategies, it is necessary to expand the boundaries of these strategies if quantitative proteomics projects are being developed. We have now integrated *MassChroQ* in *i2MassChroQ*, which makes it straightforward to perform quantitative proteomics work right after the identification—protein inference process.

The way the *MassChroQ* program is harnessed in *i2MassChroQ* is according to the following outline:

- Open an *i2MassChroQ* project or load protein identification results files;
- Configure all the aspects of the *MassChroQ* run in a specific *MassChroQ* configuration window;
- Use *i2MassChroQ* to run the external *MassChroQ* software or have *i2MassChroQ* only write the file that *MassChroQ* uses to perform its quantitative proteomics task at a later stage and outside of *i2MassChroQ*.

7.1.1 Preparing sample associations for MassChroQ

Performing quantitative proteomics experiments most likely involves comparing samples between them. That means that most often multiple samples need to be associated into meaningful groups. Before going on with the *MassChroQ* configuration, it is thus necessary to first define the sample associations. In fact, since a given sample is actually a given LC-MS run, and that each MS run's data are then used to perform protein identifications, these associations are performed between MS runs.

To perform a quantitative proteomics experiment, the very first step is to load either the protein identification results (see Section 3.4, "Loading the Protein Identification Results" or an <u>xpip</u> project file (see Section 3.5, "Loading 12MassChroQ Projects").

Once the protein identification results have been loaded (or the *i2MassChroQ* project file), the sample associations (that is, between MS run files) need to be performed by first clicking onto the *View MS identification list* button of the main program window (see SECTION 3.4.2, "DISPLAYING THE MS IDENTIFICATIONS LIST"). The MS runs are displayed in a table and sample associations can be performed by right-clicking onto the cells of the *Alignment group* column label, as shown in FIGURE 7.1, "DEFINING SAMPLE ASSOCIATIONS FOR XIC ALIGNMENTS".



Note

The sample (MS run) associations are critical not only because one wants to compare quantitative data about somehow related samples, but also because of the way *MassChroQ* performs quantification of proteomics data. Indeed, *MassChroQ* uses not *spectral count*-based strategies but an *area under the curve* strategy where the area of mass peaks is determined by looking at XIC chromatograms for these mass peaks. The associations will thus allow the software to perform the alignment of the XIC chromatograms that will be essential for the quantification analysis. Indeed, even LC-MS runs of an identical sample will not provide identical (m/z,retention time) pairs. But, to be able to quantify proteomics data on the basis of the area under the curve of XIC chromatogram peaks, it is necessary that all the XIC chromatograms for all the associated samples be properly aligned.

The associations between samples can be performed in any arbitrary way, according to the user's experimental scheme. Any number of groups can be defined that may contain any number of samples. The process is described in Figure 7.1, "Defining sample associations for XIC alignments" and Figure 7.2, "Sample associations are done by grouping samples into groups".

Alignments G	roups A Identificatio	n Run ID I Identification result file	Ms Run File	Identification engine Fasta file
Select Group	Group >	+ New Group 06_proteomics_extract_1_A01_urnb-1.xml	20120906_proteomics_extract_1_A01_urnb-1.mzML	XITandem 2017.2.1.4 contaminants
Select Group		20120906_proteomics_extract_1_A05_urnb-2.xml	20120906_proteomics_extract_1_A05_urnb-2.mzML	XITandem 2017.2.1.4 contaminants
Select Group	→ 글 As msRun reference	20120906_proteomics_extract_1_A09_urnb-3.xml	20120906_proteomics_extract_1_A09_urnb-3.mzML	🗹 X:Tandem 2017.2.1.4 contaminants
Select Group	☐ <u>R</u> emove reference	20120906_proteomics_extract_1_B01_urnb-4.xml	20120906_proteomics_extract_1_B01_urnb-4.mzML	🖄 X:Tandem 2017.2.1.4 contaminants
Select Group	Ungroup	20120906_proteomics_extract_1_B05_urnb-5.xml	20120906_proteomics_extract_1_B05_urnb-5.mzML	XITandem 2017.2.1.4 contaminants

By right-clicking into the cells of the *Alignment groups* column, groups can be defined and samples can be associated to the groups.

FIGURE 7.1: DEFINING SAMPLE ASSOCIATIONS FOR XIC ALIGNMENTS

roup-1	identa0	20120906_proteomics_extract_1_A01_urnb-1.xml	20120906_proteomics_extract_1_A01_urnb-1.mzML	🗹 X!Tandem 2017.2.1.4
roup-1	identa1	20120906_proteomics_extract_1_A05_urnb-2.xml	20120906_proteomics_extract_1_A05_urnb-2.mzML	🗹 X!Tandem 2017.2.1.4
roup-2	identa2	20120906_proteomics_extract_1_A09_urnb-3.xml	20120906_proteomics_extract_1_A09_urnb-3.mzML	🖄 X!Tandem 2017.2.1.4
roup-2	identa3	20120906_proteomics_extract_1_B01_urnb-4.xml	20120906_proteomics_extract_1_B01_urnb-4.mzML	🖄 X!Tandem 2017.2.1.4
roup-3	identa4	20120906_proteomics_extract_1_B05_urnb-5.xml	20120906_proteomics_extract_1_B05_urnb-5.mzML	🗹 X!Tandem 2017.2.1.4

Three groups have been defined, two groups having two samples each and one group having only one sample.

FIGURE 7.2: SAMPLE ASSOCIATIONS ARE DONE BY GROUPING SAMPLES INTO GROUPS

TIP: SAMPLE ASSOCIATIONS WITH SPECIFIC SAMPLE SETS

Sample associations play a critical role when samples (that is, MS runs) have conceptual relationships. For example, let's assume that a project used polyacrylamide gel electrophoresis as a protein separation method. Five related samples (be them biologically-relevant variants or technical replicates, for example) have been loaded onto five different lanes of the gel. The migration pattern between the five lanes is very similar and one could observe reproducible bands (albeit with different intensities) from one lane to the other, say, in sample 1 a band A, below a band B and so on. Sample 2 would also have that pattern, with a band A and a band B, and the same for the remaining samples (that is, lanes). Bands would be excised and subjected to trypsin digestion, the peptides would be extracted and analysed by mass spectrometry. The sample associations, here, would typically involve the definition of groups that associate related "horizontal" bands on the gel. For example, group A would associate all the bands A from the five samples, group B would associate all the bands B from the samples and so on. The sample associations would thus allow the quantification and comparison of kin proteins from the various samples.

The alignement of XIC chromatograms computed for samples from a given association group is performed by having one reference sample in that group. Each group must have a reference sample. The definition of the reference sample can be performed by the user at this stage (or at a later stage, described later) by using the context menu shown in Figure 7.3, "Setting the reference sample for the alignment".

port <u>C</u> ol	umns <u>M</u> assChroQ	
Alignme	nts Groups 🗸 🗄 Identification Run	ID I Identification result file
group-1	identa0	20120906_proteomics_extract_1_A01_urnb-1.xml
group-1	♦ <u>G</u> roup	20120906_proteomics_extract_1_A05_urnb-2.xml
group-2	<u> </u>	20120906_proteomics_extract_1_A09_urnb-3.xml
group-2	<u> </u>	20120906_proteomics_extract_1_B01_urnb-4.xm
	😵 Ungroup	20120906 proteomics extract 1 B05 urnb-5.xml

Use the context menu by right-clicking on the cells in the *Alignment groups* column to set the alignment reference in each group.

FIGURE 7.3: SETTING THE REFERENCE SAMPLE FOR THE ALIGNMENT



Note

Selecting the proper alignment reference is not something to do without thinking because the reference sample will serve as the basis for the alignment of all the samples in the group. The best sample to be chosen as alignment reference is the sample that shares the most precursor ions' m/z values with all the other samples. It is possible to delegate to *i2MassChroQ* the choice of the alignment reference sample, as described later.

Now that the sample associations have been performed, the next step is to configure *MassChroQ* from within *i2MassChroQ*. This is described in the next sections.

7.1.2 CONFIGURATION OF *MassChroQ*

i2MassChroQ provides an interface to *MassChroQ*, the software that performs XIC extractions for a list of precursor ions' m/z values. That interface is shown by selecting the *MassChroQ* menu item of the main *File* menu. The window that opens up is shown in FIGURE 7.4, "THE MASSCHROQ INTERFACE WINDOW (SAMPLE ASSOCIA-TIONS)", and is described below.

2. *		MassChroQ parameter	s	?	~ ^	8
Sample associations	Alignment	Peak quantification	MassChroQ			
All samples		Samples (MS runs) in the selected group				
		eference sample:				
Check MS run data	Check MS run data files					
ODS V Results format ON Compare samples						
		Cancel	💾 Save File	S Run M	assChro	Q

This window offers an interface to the *MassChroQ* program. The *Sample associations* tab allows one to define groups of samples that will be processed together. All the configurations in the tabs are described in the sections below.

FIGURE 7.4: THE MASSCHROQ INTERFACE WINDOW (SAMPLE ASSOCIATIONS)

7.1.2.1 The Sample associations TAB

This tab allows one to configure the sample associations. The window state shown in FIGURE 7.4, "THE MASS-CHROQ INTERFACE WINDOW (SAMPLE ASSOCIATIONS)" corresponds to a situation in which the user did not define sample associations according to the way described in SECTION 7.1.1, "PREPARING SAMPLE ASSOCIATIONS FOR MASSCHROQ". In this case, it is assumed that the user wants to treat all the samples as a single group, (the *All_samples* group). To reveal all the samples (that is, MS runs) that are being handled, check the *All_samples* check button, which will associate all the samples in that single group and display them in the right hand side list widget, as shown in FIGURE 7.5, "THE MASSCHROQ INTERFACE WINDOW (SAMPLE ASSOCIATIONS) - ALL SAMPLES LISTES".

2! *		MassChroQ parameter	s	? ~ ^ 😣	
Sample associations Alignmen		Peak quantification	MassChroQ		
All samples		Samples (MS runs) in group "All samples"			
		20120906_proteomics_e 20120906_proteomics_e 20120906_proteomics_e 20120906_proteomics_e 20120906_proteomics_e	xtract_1_A05_urnb xtract_1_A09_urnb xtract_1_B01_urnb	p-2 p-3 p-4	
Check MS run data		eference sample:	 Find the best 	t reference sample	
ODS V Results forma	t ON O	Compare samples	P Save File	🗞 Run MassChroQ	

By checking the *All_samples* check box on the left hand side list widget, all the samples in the project are associated in a single *All_samples* group and displayed in the list widget on the right hand side of the window.

FIGURE 7.5: THE MASSCHROQ INTERFACE WINDOW (SAMPLE ASSOCIATIONS) - ALL SAMPLES LISTES

If the user has crafted groups of associated samples, as described in Section 7.1.1, "Preparing sample associations for MassChroQ", the window displays different settings at start (see Figure 7.6, "The MassChroQ INTERFACE WINDOW (Sample associations) - pre-defined sample associations ").

! *		MassChroQ parameter	s	? ~ ^ 😣	
Sample associations	Alignment	Peak quantification	MassChroQ		
All samples		Samples (MS runs) in group "group-2"			
group-1		20120906_proteomics_extract_1_A09_urnb-3			
group-2	2	20120906_proteomics_extract_1_B01_urnb-4			
group-3					
	¥ ×			n	
	E <u>x</u> port <u>C</u> olumns <u>M</u> assChroQ				
	Alignme	ent groups 🗸 🖓	Identification Ru	n ID 🕴	
	group-1	identa	0	20120906_pi	
	group-1	identa	1	20120906_pr	
	group-2	identa	2	20120906_pr	
Check MS run da	group-2	identa	3	20120906_pr	
	group-3	identa	4	20120906_pr	
ODS 🗸 Results forr				·	

When the sample associations were defined before opening the *MassChroQ* interface window (the inserted window corresponds to Section 7.1.1, "Preparing sample associations for MassChroQ"), the groups of associated samples are displayed in the list widget on the left hand side of the window. Selecting group names in that list allows one to display the samples associated in a given group. To include a group in the *MassChroQ* computations, check the corresponding check box widget.

FIGURE 7.6: THE MASSCHROQ INTERFACE WINDOW (SAMPLE ASSOCIATIONS) - PRE-DEFINED SAMPLE ASSOCIATIONS

To verify which samples are being associated in a given group, select that group in the list widget on the left hand side of the window.

To make sure a given group is going to be accounted for by *i2MassChroQ* during the preparation of the file that lists all the precursor ions' peaks for which the XIC extractions needs to be performed at a later stage by *MassChroQ*, check the corresponding check box.

The *Check MS run data files* button allows the user to make sure that all the samples associated in the various groups can be found as mass spectrometry data files (\underline{mzML} or \underline{mzXML} files). This is a hard requirement because *MassChroQ* does the quantification of peptide mass spectrometric signals by extracting ion current for the peptide's precursor ion (XIC extraction). For this to be possible, the software needs to access the mass spectrometry data files.

The *Reference sample* drop-down list widget allows one to select the alignment reference sample for the currently selected sample association group in the left hand side list. The alignment reference sample must be chosen with care, as explained in Figure 7.3, "Setting the reference sample for the alignment".



Tip

If the selection of an alignment reference sample is not possible, the user might ask *i2MassChroQ* to search for it by clicking the *Find the best reference sample* button. *i2MassChroQ* will look into all the sample files associated in the current group and search for the sample that shares the maximum number of precursor ions with all the other samples. The discovered MS run file is then set to the drop-down list widget.

The *Results format* drop-down list widget allows the user to select the kind of format that the quantification results should be written in. The *ODS* format is the standard format for the *LibreOffice* software suite. The *TSV* format is a "tab-separated values" text format.

The *Compare samples* switch indicates if the results output file should display a low-details version of the data but arranged in a manner that allows the user to easily compare the quantification data about the various samples.

7.1.2.2 THE Alignment TAB

This tab allows one to configure the way the XIC chromatograms obtained for the different associated samples are aligned (see Section 7.1.1, "Preparing sample associations for MassChroQ") as shown in Figure 7.7, "The MassChroQ interface window (Alignment)".

2! *		MassChroQ parameter	s	? ~ ^ 😣			
Sample associations	Alignment	Peak quantification	MassChroQ				
	✓ Stor	e time corrections in thi	s director <u>y</u> :				
times.d							
	N	IS2 alignment paramete	ers				
MS2 tendency 10	\$						
MS2 smoothing 15	٥						
MS1 smoothing 0	\$						
ODS V Results forma	ODS V Results format ON Compare samples						
CD3 + Results forma			Deverte				
		Cancel	P Save File	Cun MassChroQ			

This tab configures the way *i2MassChroQ* performs the XIC chromatograms alignment between associated samples in the various groups. If the user is interested in the results of the alignment, the XIC retention time corrections can be stored in the directory specified at *Store time corrections in this directory* for later scrutiny.

FIGURE 7.7: THE MASSCHROQ INTERFACE WINDOW (ALIGNMENT)

The *MS2 alignment parameters* group box widget gathers parameters that are critical to the XIC chromatogram alignment algorithm for all the samples associated in a given group, as described below.

- *MS2 tendency*: half size of the window used to apply a moving median on the MS/MS retention time deviation curve. Used to create the tendency deviation curve. Of course the appropriate value for this window depends on the number of identified peptides that the two runs (reference run and run being aligned) have in common. Usually a good value is 10. While aligning, *MassChroQ* outputs the number of peptides in common which can be used to readjust this parameter if necessary.
- *MS smoothing*: half size of the window used to apply a moving average on the MS/MS retention time deviation curve. Smooths the deviation curve. Same as the above parameter, usually a good value is 10.
- *MS1 smoothing*: half size of the window used to apply a moving median on the MS retention time corrections curve. This smoothing parameter is optional, and it is not necessary most of the time. It could be used in place of the MS2 smoothing parameter in cases of a small number of shared identified peptides (< 100), in which case a good value is 20.

7.1.2.3 THE Peak quantification TAB

This tab allows one to configure the way the peaks in the XIC chromatograms are evaluated from a quantification stand point. These parameters need some testing as they might depend on the instrument whence the data originated.

2! *	MassChroQ	parameters	? ~ ^ 😣				
Sample associations	Alignment Peak quar	ntification MassChroQ					
	XIC extraction parameters						
XIC range		10.0000 🗘 ppm	~				
Inside the range, take	the	max	~				
	Peak detectio	on parameters					
smoothing	1 🗘						
minmax half window	2 🗘						
maxmin half window	з 🗘						
minmax threshold	5000 🗘						
maxmin threshold	3000 🗘						
ODS V Results format ON Compare samples							
	E	Cancel Save File	Run MassChroQ				

This tab configures the way *i2MassChroQ* performs the evaluation of the peaks in the XIC chromatograms from a quantification stand point. The settings in this dialog window might need some tweaking as they might depend on the instrument whence the data originated.

FIGURE 7.8: THE MASSCHROQ INTERFACE WINDOW (PEAK QUANTIFICATION)

- *XIC extraction parameters*: these parameters govern the way the program searches for m/z values in the mass spectral data.
 - *XIC range*: the m/z width (mass tolerance) for searching m/z values in the mass data during the XIC extraction. Units can be part-per-million (*ppm*), resolution (*res*) or Dalton (*dalton*). The wider the window, the rougher the XIC extraction. This value typically depends on the resolving power of the instrument that acquired the data.
 - *Inside the range, take the*: once the m/z window has been located in the mass spectral data, it will contain a nubmer of points. This settings determines what kind of signal intensity to compute for the m/z window (that is, what to do with the m/z points contained in the m/z window). If *max*is

selected, only the max-intensity point in the m/z window is used as the signal intensity corresponding to the m/z window. If *sum* is selected, the sum of the intensities of all the m/z points in the window is used.

- *Peak detection parameters*: these parameters govers the way the program detects peaks.
 - *smoothing*: number of points around the point being considered in the XIC chromatogram. If set to one, the rolling window will contain three points: one before the considered point, one after it and the considered point itself. This setting thus determines the width of the rolling window that is used to iterate in the XIC chromatogram in search for peaks. This window, whatever the setting, will shift by one point at each iteration in the XIC chromatogram.
 - *minmax half window*: the half window size used to apply the close (min/max) transform on the XIC intensities. This window determines the number of scan points over which two peaks will be considered separately, otherwise they would have been merged. A good half window value is usually 3 (which makes a window of 7).
 - *maxmin half window*: same as above but for the close (max/min) transform. This window determines the minimum peak width (in scan points number) below which the peak would not be detected. A good half window value is usually 2 (which makes a window of 5).
 - *minmax threshold*: threshold on the close signal: a minimum intensity value below which peaks are not detected on the closing signal. This threshold is usually two or three times the background noise intensity level, which depends on your mass spectrometer.
 - *maxmin threshold*: threshold on the open signal: a minimum intensity value below which peaks are not detected. It corresponds to the opening signal upper limit and it represents the background signal upper level. A good value would thus be slightly bigger than your background noise intensity level.

7.1.2.4 ТНЕ *MassChroQ* ТАВ

This tab allows one to configure the way *MassChroQ* actually performs the quantification (if using the *Run MassChroQ*) or the way *i2MassChroQ* writes the <u>masschroqml</u> file to be fed to the *MassChroQ* program.

2/ ×		MassChroQ paramete	rs	? ~ ^ 😣			
<u>Sample associations</u>	Alignment	Peak quantification	MassChroQ				
	C Edit MassChroQ execution						
Select the MassChro	Select the MassChroQ binary path						
	Ru	n MassC <u>h</u> roQ through	HTCondor				
MassChroQ memor	y requirement (MB) 5000		\$			
		MassChroQ paramete					
		MassChroQ paramete	15				
Number of CPUs	Number of CPUs 1 🗘						
Temporary directory	Temporary directory Select the temporary dir (default: /tmp/)						
Use the tempora	Use the temporary directory to store detected peaks						
Perform peptide	Perform peptide <u>t</u> ext file parsing only						
ODS V Results forma	ODS V Results format ON Compare samples						
		🗵 Cancel	Save File	S Run MassChroQ			

This tab configures the way either *MassChroQ* actually performs the quantification or *i2MassChroQ* writes the <u>masschroqml</u> file that *MassChroQ* will be fed with to perform the task.

FIGURE 7.9: THE MASSCHROQ INTERFACE WINDOW (MASSCHROQ)

- *Edit MassChroQ execution*: activate the check button to use the directory icon to locate the *MassChroQ* program on disk. The full path to the program will be printed in the line edit widget next to the icon.
- *Run MassChroQ through HTCondor*: activate the check button to set the memory requirements for HT-Condor.
- *MassChroQ parameters*: these settings govern the actual *MassChroQ* quantification process:
 - *Number of CPUs*: set the number of central processing units that *MassChroQ* is allowed to use (these are actually called "threads").
 - *Temporary directory*: use the directory icon to select a specific temporary directory where *Mass-ChroQ* will write processing-related data. By default the directory is <u>/tmp/</u>. The temporary files are eliminated when no more used.
 - *Use the temporary directory to store detected peaks*: if checked, the detected peaks might be stored in files in the temporary directory described above. This can be construed as a swap area where to store peaks data if the available memory is insufficient.

SAVING THE FILE AND OPTIONALLY RUNNING MassChroQ 7.1.2.5

Once all the configuration has been done, the user can either only save the masschrogml file by clicking on the Save File button or immediately start MassChroQ by clicking on the Run MassChroQ button.



Νοτε

Even if the user decides to go down the direct Run MassChroQ route, the program will ask to save the masschrogml file. This is because that file is read by MassChroQ when iz MassChroQ internally calls it to run the quantification process.

The masschroqml file describes the proteins and peptides that were retained during the protein identification results analysis session. The contents of the file are shown in FIGURE 7.10, "CONTENTS OF THE MASSCHROQML FILE".

```
path="1 A01 urnb-1.mzML'
                                                                                                                                                                                       type="centroid"/>
                                                                                                               path="1_A01_urnb-1.mzML" type= centroid />
path="1_A05_urnb-2.mzML" type="centroid"/>
path="1_A09_urnb-3.mzML" type="centroid"/>
path="1_B01_urnb-4.mzML" type="centroid"/>
path="1_B05_urnb-5.mzML" type="centroid"/>
    <data_file id="msruna2" format="mzml"
<data_file id="msruna3" format="mzml"
<data_file id="msruna4" format="mzml"</pre>
     <data_file id="msruna5" format="mzml"
      <group id="All_samples" data_ids="msruna1 msruna2 msruna3 msruna4 msruna5"/>
   sprotein_ctst>
<protein id="a1.a1.a1" desc="P04711 Phosphoenolpyruvate carboxylase"/>
<protein id="b88.a1.a1" desc="NP_001152746 ascorbate peroxidase"/>
<protein id="b88.a2.a1" desc="NP_001150192 APx1 - Cytosolic Ascorbate Peroxidase"/>
<protein id="c388.a1.a1" desc="B6SIF9 Histone H2B seq=translation"/>
     cycpeptide id="pepa1a4" mods="[C 74 H 121 0 26 N 18 S 1] " prot_ids="a1.a2.a1" mh="1709.8414637049" seq="ALLDEMAVVATEEYR">

    cobserved_in data="msrunal" scan="14760" i= 372
    //peptide>
    /opeptide>
    cobserved_in data="msruna1" scan="15553" z="2"/>
         cobserved_in data="msruna1" scan="15687" z="2"/>
         cobserved_in data="msruna2" scan="15687" z="2"/>
         cobserved_in data="msruna3" scan="15715" z="3"/
          cobserved_in
</peptide>
</peptide_list>
                                           <!--write_time_values_output_dir="directory" to write retention time corrections--->
<ms2_tendency_halfwindow>10</ms2_tendency_halfwindow>
<ms2_smoothing_halfwindow>15</ms2_smoothing_halfwindow>
<ms1_smoothing_halfwindow>
                                           <ms1_smoothing_halfwindow>0</ms1_smoothing_halfwindow>
         </msi_smootheng_nderwendows
</msi_smootheng_nderwendows
</alignment_methods
<align group_id="All_samples" method_id="my_ms2" reference_data_id="msruna1"/>
```

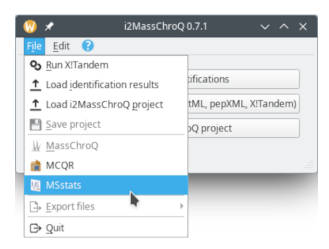
The <u>masschroqml</u> file contains all the required data and configuration bits to perform the XIC extractions for all the peptidic precursor ions that allowed identifying proteins. This file is read by the *MassChroQ* program. (In this screen dump, the file contents were obviously redacted for brevity.)

FIGURE 7.10: CONTENTS OF THE MASSCHROQML FILE

7.2 INTERFACE TO THE *MSstats* STATISTICS MODULE

The statistical analysis of the quantified peptide data by *MassChroQ* is assigned to the *MSstats* software authored by M. Choi and colleagues (2014) *MSstats: An R package for statistical analysis of quantitative mass spectrometry-based proteomic experiments* in *Bioinformatics*. As a prerequisite, peptide quantification must thus have been performed by *MassChroQ* as described in earlier sections.

This section describes in a step-by-step fashion the interface that *i2MassChroQ* provides to the *MSstats* software. To start the process, select the the *MSstats* menu item of the main *File* menu, as shown in FIGURE 7.11, "MSSTATS MENU ITEM IN THE MAIN 12MASSCHROQ MENU".



Menu that loads the interface to the *MSstats* statistics software that processes the *MassChroQ*-quantified peptide data to provide protein quantifications.

FIGURE 7.11: MSSTATS MENU ITEM IN THE MAIN I2MASSCHROQ MENU

When the *MSstats* interface is loaded it looks like that shown in FIGURE 7.12, "MAIN MSSTATS INTERFACE WIN-DOW". The actions to be carried over are shown on the left hand side region of the window in the form of a series of actions that are materialized by gradient-filled buttons. Each one of these actions are described in the following sections.

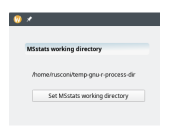
*	MSstats	~ ^
MSstats working directory	Log book	
not selected		
Set MSstats working directory		
	h	
	Status	
	Program output	
	You can change or check the number of thread(s) used by MCQR at anytime with the functions 'mcq.set.max.threads' and 'mcq.get.max.threads'.	Î
Load	See ?mcq.set.max.threads or ?mcq.get.max.threads for more information.	
MCQR dubious data filter	MCQRInfo: MCQR version : 0.6.11	
MSstats data processing		
MSstats quantification by sa	Attaching package: 'MSstats'	
MSstats quantification by gro	The following object is masked from 'package:grDevices':	
Rscript	required libraries are ready to use	٣
RSCript		

The main *MSstats* interface window has two main regions: the left hand side part of the window contains all the workflow steps that are to be carried out from the topmost item to the bottommost item; the right hand side part of the window contains three elements: the *Log book* view at the top, the *Program output* at the bottom, and the central *Status* widget.

FIGURE 7.12: MAIN MSSTATS INTERFACE WINDOW

7.2.1 Setting the temporary *MSstats* working directory

The first choice to be made by the user is to define where the *MSstats* package will write the data it needs to fulfill its statistical analysis tasks (FIGURE 7.13, "SETTING THE MSSTATS WORKING DIRECTORY"). The directory needs to be created if it does not exist already.

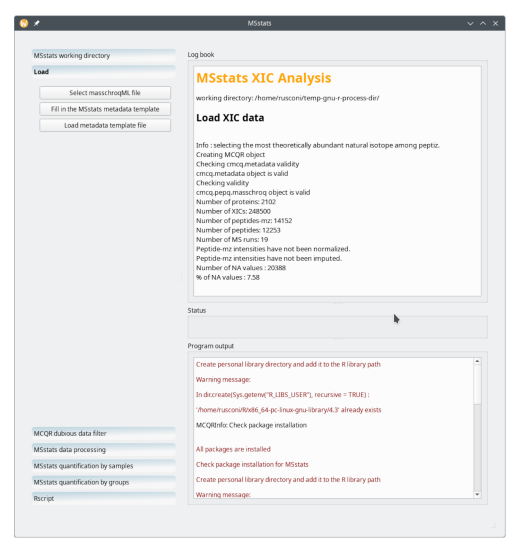


The directory is used by *MSstats* to store all the files and directories it creates during its fulfilling of its tasks. That directory can be located anywhere on the disk and needs to be created if it does not exist already.

FIGURE 7.13: SETTING THE MSSTATS WORKING DIRECTORY

7.2.2 Loading the peptide quantification data file by MassChroQ

The *MassChroQ*-based peptide quantification process must have been performed already and must have produced an XML file with the <u>masschroqml</u> extension. That file can be loaded by clicking the *Select masschroqML file* button. Upon completing the data file loading process, the right hand side pane of the windows shows a summary of the data just loaded in the *Log book* widget (FIGURE 7.14, "LOADING MASSCHROQ-GENERATED DA-TAFILE").



The peptide quantification process by *MassChroQ* produces a file that the user must load by clicking onto the *Select masschroqML file* button. As shown in the *Log book* pane on the right hand side of the window, the data have been loaded and a summary is provided.

FIGURE 7.14: LOADING MASSCHROQ-GENERATED DATA FILE

Once the data have been loaded, *i2MassChroQ* crafts a brand new spreadsheet data set in memory that needs to be stored on disk. For this, the user clicks the *Fill in the MSstats metadata template* button, which will permit saving the file to disk (with the OpenDocument format, ods extension, typically in the working directory created

earlier). *i2MassChroQ* will try to automatically open that file right after having written it to disk. If the file cannot be opened, then user needs to open it manually and start filling in the metadata for *MSstats* to performs its work as intended. A typical unmodified metadata template file looks like shown in FIGURE 7.15, "METADATA TEMPLATE FILE FOR USE BY MSSTATS" where only the MS run file names are listed.

D	R		
<u>F</u> ile	<u>E</u> dit <u>V</u> iew <u>I</u> nsert	F <u>o</u> rmat St <u>y</u> les <u>S</u> heet <u>D</u> ata <u>T</u> ools <u>W</u> indow <u>H</u> elp	
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_			
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F22	~	f _× ∑ × =	
	A	B C D	
1	msrun	msrunfile BioReplicate Condition	
2	msruna1	20120906 balliau extract 1 A01 urnb-1.mzML	
3	msruna2	20120906 balliau extract 1 A02 urzb-1.mzML	
4	msruna3	20120906 balliau extract 1 A04 teal-1.mzML	
5	msruna4	20120906 balliau_extract 1_A05_urnb-2.mzML	
6	msruna5	20120906 balliau extract 1 A06 urzb-2.mzML	
7	msruna6	20120906 balliau extract 1 A07 tca1-2.mzML	
8	msruna7	20120906 balliau extract 1 A08 teal-2.mzML	
9	msruna8	20120906_balliau_extract_1_A09_urnb-3.mzML	
10	msruna9	20120906_balliau_extract_1_A10_urzb-3.mzML	
11	msrunb10	20120906_balliau_extract_1_A11_tca1-3.mzML	
12	msrunb11	20120906_balliau_extract_1_A12_teal-3.mzML	
13	msrunb12	20120906_balliau_extract_1_B01_urnb-4.mzML	
14	msrunb13	20120906_balliau_extract_1_B02_urzb-4.mzML	
15	msrunb14	20120906_balliau_extract_1_B03_tca1-4.mzML	
16	msrunb15	20120906_balliau_extract_1_B04_teal-4.mzML	
17	msrunb16	20120906_balliau_extract_1_B05_urnb-5.mzML	
18	msrunb17	20120906_balliau_extract_1_B06_urzb-5.mzML	
19	msrunb18	20120906_balliau_extract_1_B07_tca1-5.mzML	
20	msrunb19	20120906_balliau_extract_1_B08_teal-5.mzML	
21			_

A typical *MSstats* metadata template file as created by *i2MassChroQ*. That file needs to be modified according the user requirements in terms of grouping the samples according to the experiment plan. When created, the file only contains the list of MS runs from which peptide quantifications were performed.

FIGURE 7.15: METADATA TEMPLATE FILE FOR USE BY MSSTATS

Once modified by the user to annotate the MS run file names and to group the samples according to the experiment plant, the file looks like shown in FIGURE 7.16, "ANNOTATED METADATA FOR USE BY MSSTATS".

	*			m
<u>F</u> ile	<u>E</u> dit <u>V</u> iew	r <u>I</u> nsert F <u>o</u> rmat Styles <u>S</u> heet <u>D</u> ata <u>T</u> ools <u>W</u> indo	w <u>H</u> elp	
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A21		✓ f _* ∑ - =		
	А	В	С	D
1	msrun	msrunfile	Condition	BioReplicate
2	msruna1	20120906 balliau extract 1 A01 urnb-1.mzXML	urnb	1
3	msruna2	20120906 balliau extract 1 A02 urzb-1.mzXML	urzb	2
4	msruna3	20120906 balliau extract 1 A04 teal-1.mzXML	teal	3
5	msruna4	20120906 balliau extract 1 A05 urnb-2.mzXML	urnb	4
6	msruna5	20120906 balliau extract 1 A06 urzb-2.mzXML	urzb	5
7	msruna6	20120906 balliau extract 1 A07 tca1-2.mzXML	tca	6
8	msruna7	20120906 balliau extract 1 A08 teal-2.mzXML	teal	7
9	msruna8	20120906_balliau_extract_1_A09_urnb-3.mzXML	urnb	8
10	msruna9	20120906 balliau extract 1 A10 urzb-3.mzXML	urzb	9
11	msrunb10	20120906_balliau_extract_1_A11_tca1-3.mzXML	tca	10
12	msrunb11	20120906_balliau_extract_1_A12_teal-3.mzXML	teal	11
13	msrunb12	20120906_balliau_extract_1_B01_urnb-4.mzXML	urnb	12
14	msrunb13	20120906_balliau_extract_1_B02_urzb-4.mzXML	urzb	13
15	msrunb14	20120906_balliau_extract_1_B03_tca1-4.mzXML	tca	14
16	msrunb15	20120906_balliau_extract_1_B04_teal-4.mzXML	teal	15
17	msrunb16	20120906_balliau_extract_1_B05_urnb-5.mzXML	urnb	16
18	msrunb17	20120906_balliau_extract_1_B06_urzb-5.mzXML	urzb	17
19	msrunb18	20120906_balliau_extract_1_B07_tca1-5.mzXML	tca	18
20	msrunb19	20120906_balliau_extract_1_B08_teal-5.mzXML	teal	19

Once the template metadata have been completed to inform *MSstats* about the sample grouping and the analysis logic, it might look like shown in this figure.

FIGURE 7.16: ANNOTATED METADATA FOR USE BY MSSTATS

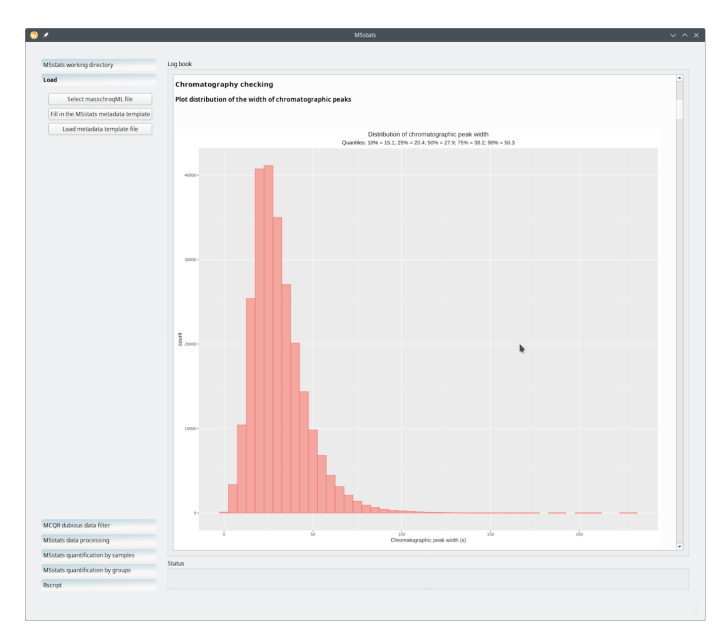
The template file, as modified by the user should next be loaded by clicking on the *Load metadata template file* button. The *Log book* widget now shows informational data like shown in FIGURE 7.17, "PRELIMINARY PROCESS-ING PERFORMED BY MCQR UPON LOADING OF THE METADATA TEMPLATE FILE". The output data are comprehensive and illustrated with graphs like the one described below.

*	MSstats	\sim
MSstats working directory	Log book	
.oad		
	Ponding metadata file	
Select masschroqML file	Reading metadata file	
Fill in the MSstats metadata template	Columns "Condition" and "BioReplicate" specific to MSstats	
Load metadata template file		
	Merging metadata	
	Warning : Please carefully check your data and metadata because there are differences between object and metadata. Your 'msrunfile' have been renamed. Number of proteins: 2102	
	Number of XICs: 248500	
	Number of peptides-mz: 14152 Number of peptides: 12253	
	Number of MS runs: 19 Peptide-mz intensities have not been normalized.	
	Peptide-mz intensities have not been imputed. Number of NA values : 20388	
	% of NA values : 7.58	
	Chromatography checking	
	Plot distribution of the width of chromatographic peaks	
	Distribution of chromatographic peak width Quantiles: 10% = 15.1; 25% = 20.4; 50% = 27.9; 75% = 38.2; 90% = 50.3	
	4000-	
	•	
	3000-	
	Status	
	Program output	
	In dir.create(Sys.geterw("R_LIBS_USER"), recursive = TRUE) :	
	'/home/rusconi/R/x86_64-pc-linux-gnu-library/4.3' already exists	
CQR dubious data filter	MCQRInfo: Check package installation	
ISstats data processing	All packages are installed	
Sstats quantification by samples	Check package installation for MSstats	
ISstats quantification by groups	Create personal library directory and add it to the R library path	
tscript		

The template metadate file loading step triggers preliminary computing tasks by MCQR (a *GNUR* script developed in our facility) and the output is provided in the *Log book* widget.

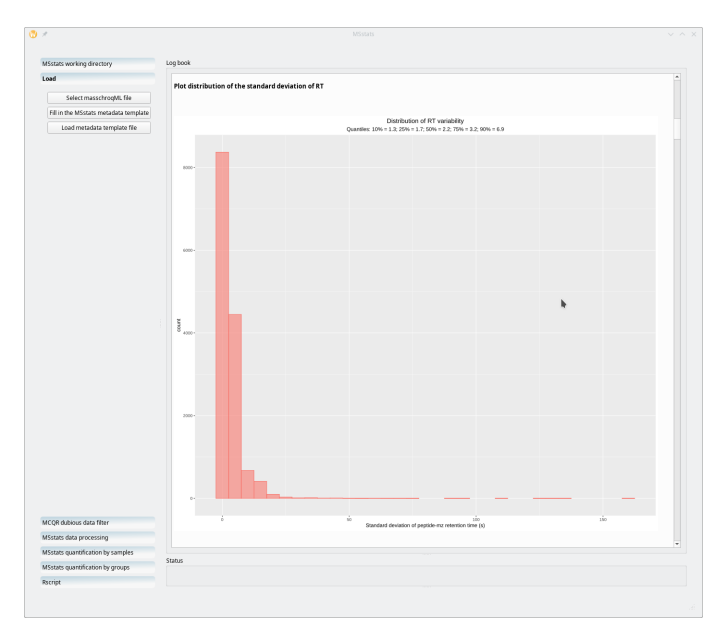
FIGURE 7.17: PRELIMINARY PROCESSING PERFORMED BY MCQR UPON LOADING OF THE METADATA TEMPLATE FILE

One particular informational bit that is of use in a later step of the processing workflow is the distribution of the chromatographic peak width over all the samples and over the whole retention time (FIGURE 7.18, "CHROMATOG-RAPHY DATA CHECKS: THE DISTRIBUTION OF THE CHROMATOGRAPHY WIDTHS"). Equally useful is the distribution of retention time variability for all the (m/z,z) pairs that were extracted from the whole set of MS run acquisitionsFIGURE 7.19, "CHROMATOGRAPHY DATA CHECKS: THE DISTRIBUTION OF RETENTION OF RETENTION TIME VARIATIONS".



MSstats prints out data used by *i2MassChroQ* to plot a number of graphics like this histogram showing the distribution of the peak widths (in seconds). One can assume that a given ion might be reasonably contained in a retention time range (0—150) seconds.

FIGURE 7.18: CHROMATOGRAPHY DATA CHECKS: THE DISTRIBUTION OF THE CHROMATOGRAPHY WIDTHS



The histogram plot here shows the variability of the retention time values for all the (m/z,z) pairs extracted from the whole set of MS run acquisitions. One might consider that the maximum standard peak width variation acceptable is 30.

FIGURE 7.19: CHROMATOGRAPHY DATA CHECKS: THE DISTRIBUTION OF RETENTION TIME VARIATIONS

The two information bits described in the two figures above are of use in the next step of the processing workflow, as described in the next section.

7.2.3 FILTERING DUBIOUS DATA BY RUNNING MCQR

This step is optional. It is performed by MCQR. The idea is that the user should be able to scrap some dubious data from the data set if these data are outside of "reasonable ranges". For example, one should be able to filter out (m/z,z) pairs if they match retention times of too large a range (that is, for example, an ion being detected in the MS run acquisition over too long a retention time range, which is suspect).

A		
MSstats working directory	Log book	
5 ,		
Load		
MCQR dubious data filter	Before MCQR dubious data filter	
Remove peptides with large chromatographics peaks	Number of proteins: 2102	
Peak width cutoff 150 🗘	Number of XICs: 248500	
	Number of peptides-mz: 14152 Number of peptides: 12253	
Remove peptides with Retention time variation	Number of MS runs: 19	
	Peptide-mz intensities have not been normalized.	
Retention time variation cutoff 30 🗘	Peptide-mz intensities have not been imputed. Number of NA values : 20388	
S Filter dubious data	% of NA values : 7.58	
S Filter dubious data	10011474010037730	
	After MCQR dubious data filter	
	Number of proteins: 2102 Number of XICs: 247514	
	Number of xICs: 24/514 Number of peptides-mz: 14090	
	Number of peptides: 12214	
	Number of MS runs: 19	
	Peptide-mz intensities have not been normalized.	
	Peptide-mz intensities have not been imputed. Number of NA values : 20196	
	% of NA values : 7.54	
//Sstats data processing		-
	4	► F
MSstats quantification by samples	Status	
MSstats quantification by groups		
Rscript		

Dubious data might be filtered out on the basis of the two criteria shown. The numerical values set in this example are based on the output of *MSstats* as shown in the previous histograms.

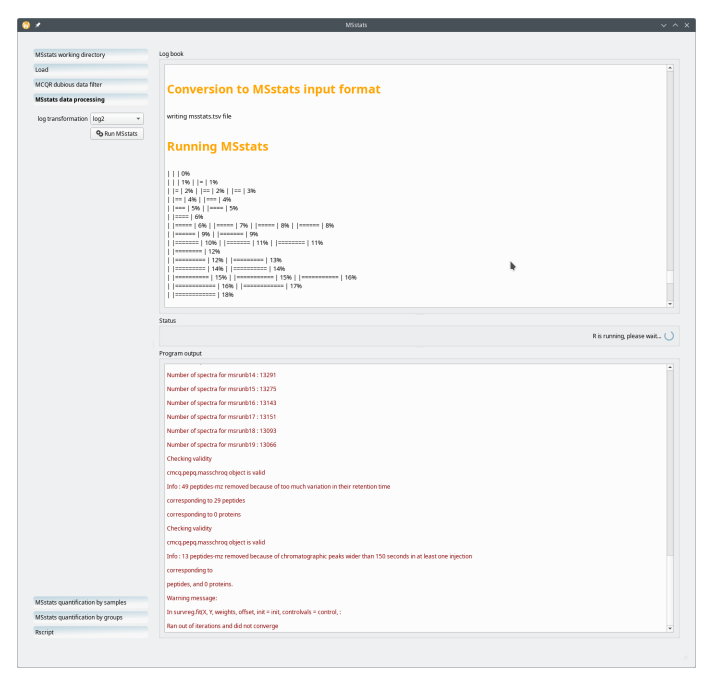
FIGURE 7.20: FILTERING DUBIOUS DATA USING MCQR

The dubious data filtering is performed on the basis of two criteria: the retention time peak width (*Peak width cutoff*) and the retention time variability (*Retention time variation cutoff*). *i2MassChroQ* documents the filtering step in the *Log book* widget as shown in FIGURE 7.20, "FILTERING DUBIOUS DATA USING MCQR".

As visible in FIGURE 7.20, "FILTERING DUBIOUS DATA USING MCQR", in the ouput printed in the *Log book* widget, the data set is pretty good, since applying the filters did only remove 39 peptides over more than twelve thousands and not a single protein.

7.2.4 RUNNING MSstats ON THE CONFIGURED DATA SET

The next step in the wokflow is to actually run *MSstats*. However, one last bit of configuration is required: the user is requested to select the log transformation (either log2 or log10) because that is a prerequisite for *MSstats* to run. Once that configuration bit has been set, the user might click on the *Run MSstats* button.



After having chosen the log tranformation (log2 or log10) that is *required* by *MSstats*, the user clicks on the *Run MSstats* button. The output shows the advancement of the computations.

FIGURE 7.21: RUNNING MSSTATS ON THE CONFIGURED DATA SET

At the end of the computation, as shown in Figure 7.21, "Running MSstats on the configured data set", the starts the next workflow step.

7.2.5 RUNNING THE *MSstats* QUANTIFICATION BY SAMPLES

One of the manners in which the *MSstats*-based quantification process can be run is the "quantification by samples" mode. This is triggered by clicking on the *MSstats quantification by samples* workflow item, as shown in **FIGURE 7.22**, "MSstats QUANTIFICATION BY SAMPLES MODE".

,	MSstats	~
ISstats working directory	Log book	
pad	90%	91%
ICQR dubious data filter	92%	
Sstats data processing	93%	
Sstats quantification by samples	94%	
Stats sample quantification	95%	
Save protein abundance results	=====================================	
	97% 	
	98%	
	99% 99%	
	100%	
	MSstats quantification process on samples	
	missible quantification process on sumples	
	done	

In the quantification by samples mode, the samples are taken as individual samples depending on their *BioReplicate* number in the metadata template file. See text for details.

FIGURE 7.22: MSSTATS QUANTIFICATION BY SAMPLES MODE

The quantification process depicted in FIGURE 7.22, "MSSTATS QUANTIFICATION BY SAMPLES MODE" is very quick. The "processing by samples" mode will quantify protein on the basis of the "*BioReplicate*" variable in the metadata template file (FIGURE 7.16, "ANNOTATED METADATA FOR USE BY MSSTATS"). Because, in the example, each MS run acquisition (that is, each row of the spreadsheet page) is marked as a different *BioReplicate*, the quantification by samples mode will quantify proteins found in each individual sample separately.

The user will want to save the protein quantification results by saving them to a spreadsheet file. This step is achieved by clicking the *Save protein abundance results* button. The saved spreadsheet file is shown in FIGURE 7.23, "Spreadsheet view of the QUANTIFICATION BY SAMPLES RESULTS".

0 🗶		quant	fication-by-sample-results - LibreOffice Calc			
<u>File Edit View Insert Format Styles</u>	<u>Sheet Data Tools Window H</u> elp					
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		••••		-		
A1	ein					
A B C	D E F	G H I	J K L	M N O	P Q R	S T
1 Protein tca 6 tca 10	tca 14 tca 18 teal 3	teal 7 teal 11 teal 15	teal 19 urnb 1 urnb 4 u	rnb 8 urnb 12 urnb 16	urzb 2 urzb 5 urzb 9 u	rzb 13 urzb 1
2 GRMZM2G0 24.99149 25.2030613	3 24.9085943 24.7369918 25.2471146	25.2366565 25.0799455 25.2712639	25.0308132 23.7139587 24.5164874	24.2058997 24.5228861 24.3433468	24.6023815 24.7734412 24.0609365	24.5270675 24.503
3 GRMZM2G@ 24.2142014 23.8940373	3 24.2144983 24.0074594 23.0379256	23.1742483 23.1277379 22.9903432	23.504089 23.4311045 23.9066337	23.9430848 23.8345213 24.1349405	24.0056634 23.9027203 24.0583954	23.9086968 23.913
4 GRMZM2G) 24.2828722 24.2491433	3 24.2412872 24.0985407 23.7564139	24.1713002 24.1171434 24.0092049	24.163269 24.1604197 24.1626585	24.2622599 24.1209541 24.2531187	24.1548299 24.3816418 24.0814061	24.1446607 24.255
5 GRMZM2G? 24.5804333 24.5494043	3 24.5581736 24.4120157 22.1210576	23.7081262 22.5824257 23.0073712	22.9118607 24.3863739 24.1002344	24.1139798 24.0976093 24.1829865	24.1175504 23.7264604 23.9623617	24.1325538 23.975
6 GRMZM2G3 22.1800067 20.638628	22.5087298 22.4728324 25.7342082	25.5362149 23.5516447 23.1664744	25.5586894 23.4838089 22.7373527	23.0139565 23.0494026 23.0403918	22.8782096 22.9149991 22.6217807	22.9979969 21.87
7 GRMZM2G) 24.1706033 24.1334423	3 24.1634204 23.7570512 23.9051287	24.1288791 24.03655 24.0609592	23.9215876 24.3736719 24.3470932	24.0317881 24.3470706 24.3278334	24.3067646 24.2682762 24.2567991	24.2506848 24.209
8 GRMZM2G0 23.76814 23.6572042	2 23.5963359 23.5542755 23.0119632	23.5734328 22.1458353 22.6883096	22.7935342 23.7941676 23.8154495	23.7804384 23.6710182 23.7476813	23.9118348 23.4467422 23.1984283	23.649604 23.780
9 GRMZM2G0 25.5944947 26.0473367	26.0655645 25.9582716 25.7083421	25.753472 25.6985986 25.785011	25.6445028 25.882529 25.46919	25.6331737 25.6871481 25.4528868	25.3022065 25.5674751 25.3673691	25.6298127 25.667
10 AC234157.1 26.4371452 26.1754312	2 26.3257097 26.4736952 26.6577783	26.4432699 27.3101185 26.7225562	26.7804765 25.6548028 25.9548268	26.1449254 26.0529609 26.0767347	26.0151289 25.9190147 25.9396926	25.9564025 25.961
11 GRMZM2G# 25.5132374 25.1282174	25.2316531 25.211796 25.5473325	25.2708734 25.5393336 25.7516532	25.4501353 26.0588964 25.8137166	25.4186889 25.6624547 25.8392889	25.703141 25.7878555 25.7354365	25.8212799 25.561
12 GRMZM2G0 23.6059748 23.3567495	5 23.4016451 23.4958117 24.1257384	23.741926 24.253306 24.2124685	24.2139682 22.5721485 23.5436861	23.2287041 22.5127526 2.1004834	23.341311 23.3427301 23.3000225	22.899138 22.94
13 GRMZM2G> 23.5775546 23.8941526	23.570007 23.3027719 22.9454379	23.5301193 23.1783634 23.4464302	23.6911901 23.2767406 23.3796094	23.8611058 23.1886836 23.2253297	26.2194721 26.3289109 22.9509712	23.5254699 23.677
14 GRMZM2G? 24.7551618 24.763397	24.6962298 24.9362583 24.6263543	24.7584957 24.7500337 24.9382337	24.8642908 25.0398982 24.8019072	25.1172658 24.9826424 24.8104419	24.8290495 24.8278396 24.6826869	24.8820369 24.877
15 GRMZM2G? 26.0173647 25.9301647	25.8779207 26.1776739 25.6914874	25.9559881 25.9010924 25.8000601	25.9726493 26.1488921 25.9220936	26.1257349 26.0556673 26.0376858	25.8394741 26.0309442 25.8857576	26.0688823 25.937
16 GRMZM2G) 23.0431394 22.9714926	22.7664769 23.2425906 23.0586359	23.1617575 22.7082082 23.1581848	23.0882948 23.2352567 23.0393947	23.1769869 23.0212603 22.9351532	23.0906809 23.06101 23.0278385	23.0526321 22.937
17 GRMZM2G@ 26.1184947 26.0751918	3 25.9570704 26.0154477 25.9328394	26.2831897 26.2003816 25.9981566	26.2459892 25.9880591 26.0237671	25.7705258 25.697581 25.9939536	25.6291979 25.5212082 25.5506413	25.4360126 25.658
18 GRMZM2G3 24.3018529 24.261854	1 24.4498349 24.5895886 23.6564672	23.8464348 24.1840714 23.9895215	24.1845867 24.258298 24.2967222	24.3009282 24.2245127 24.2688698	24.1781539 24.1718807 24.2926755	24.1427893 24.114
sampleQuantification						
S Find	V Find All Formatted Display	Match Case 🔍				
Sheet 1 of 1		mp1	English (USA)	I. 🗈	Average: ; Sum: 0	

In the metadata template file, each MS run acquisition was listed as a different *BioReplicate* identity. This means that proteins were quantified independently in each MS run. The spreadsheet view in this figure shows quantification data for each protein (each row) found in each each sample (the columns).

FIGURE 7.23: SPREADSHEET VIEW OF THE QUANTIFICATION BY SAMPLES RESULTS

7.2.6 RUNNING THE *MSstats* QUANTIFICATION BY GROUPS

The other manner in which the *MSstats*-based quantification process can be run is the "quantification by groups" mode. This is triggered by clicking on the *MSstats quantification by groups* workflow item, as shown in FIG-URE 7.24, "MSSTATS QUANTIFICATION BY GROUPS MODE".

stats working directory	Log book	
ad	======================================	
CQR dubious data filter		
Sstats data processing		
ISstats quantification by samples		
Sstats quantification by groups	MSstats quantification process on samples	
Stats group quantification		
Save protein abundance results		
	done	
	MSstats quantification process on groups (for each condition described in	
	metadata)	
	includita)	
	done	
	4	
	Status	
	Status Program output	
	Status Program output Number of spectra for msrunb17 : 13151	
	Status Program output Number of spectra for msrunb17 : 13151 Number of spectra for msrunb18 : 13093	
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	Status Program output Number of spectra for msrunb17: 13151 Number of spectra for msrunb18: 13093 Number of spectra for msrunb18: 13093 Number of spectra for msrunb19: 13066 Checking validity cmcq.pepq.masschroq object is valid Info: 49 peptides-mz removed because of too much variation in their retention time corresponding to 29 peptides corresponding to 0 proteins Checking validity rmcq.pepq.masschroq object is valid Info: 13 peptides-mz removed because of chromatographic peaks wider than 150 seconds in at least one injection corresponding to peptides.mz removed because of chromatographic peaks wider than 150 seconds in at least one injection corresponding to memory and proteins. Warning message:	
	Status Program output Number of spectra for msrunb17 : 13151 Number of spectra for msrunb18 : 13093 Number of spectra for msrunb19 : 13066 Checking validity cmcq.pepq.masschroq object is valid Info : 49 peptides-mz removed because of too much variation in their retention time corresponding to 29 peptides corresponding to 0 proteins Checking validity cmcq.pepq.masschroq object is valid Info : 13 peptides-mz removed because of chromatographic peaks wider than 150 seconds in at least one injection corresponding to peptides, and 0 proteins. Warning message: In survreg.fit(X, Y, weights, offset, init = init, controlvals = control, : Ran out of iterations and did not converge	
	Status Program output Number of spectra for msrunb17: 13151 Number of spectra for msrunb18: 13093 Number of spectra for msrunb18: 13093 Number of spectra for msrunb19: 13066 Checking validity cmcq.pepq.masschroq object is valid Info: 49 peptides-mz removed because of too much variation in their retention time corresponding to 29 peptides corresponding to 0 proteins Checking validity rmcq.pepq.masschroq object is valid Info: 13 peptides-mz removed because of chromatographic peaks wider than 150 seconds in at least one injection corresponding to peptides.mz removed because of chromatographic peaks wider than 150 seconds in at least one injection corresponding to peptides.and 0 proteins. Warning message: In survreg.ft(X, Y, weights, offset, init = init, controlvals = control, :	

In the quantification by groups mode, the samples are first grouped into groups defined in the metadata template file. In that file, the column that specifies the required grouping has the *Condition* header. See text for details.

FIGURE 7.24: MSSTATS QUANTIFICATION BY GROUPS MODE

The quantification process depicted in FIGURE 7.24, "MSSTATS QUANTIFICATION BY GROUPS MODE" is very quick. The "processing by groups" mode will quantify proteins on the basis of the "*Condition*" variable in the metadata template file (FIGURE 7.16, "ANNOTATED METADATA FOR USE BY MSSTATS"). Because, in the example, there are four different *Condition* values, there will be four groups of proteins (different protein solubilization methods).

The user will want to save the protein quantification results by saving them to a spreadsheet file. This step is achieved by clicking the *Save protein abundance results* button. The saved spreadsheet file is shown in FIGURE 7.25, "Spreadsheet view of the QUANTIFICATION BY GROUPS RESULTS".

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	А	В	с	D	E ^	E
1	Protein	tca	teal	urnb	urzb	
2	GRMZM2G035268 P0	25.5733642001247	25,56094786201	25.67674667236	25.650668000879	L I
3	GRMZM2G015384 P0	23.5947503096769	23.541259181921	23.59237042983	23.558290389996	
4	GRMZM2G029186 P0	23.2272801317162	22.656693558875	23.20828389314	23.386550238628	<u>م</u> ر
5	GRMZM2G072909 P0	22.4160674243762	23.087016696402	22.51190586447	22.405613253861	
б	GRMZM2G036609 P0	24.7436269193898	3 24.519785781034	24.77447143046	24.731584626287	Ø
7	GRMZM2G076950 P0	22.7199419246104	22.830281219265	22.81619338008	22.971124217844	
8	GRMZM2G096153 P0	25.8631149447885	5 25.780612615971	26.04172314717	25.97264073261	f
9	GRMZM2G030072 P0	23.4126109258736	3 23.586061181892	23.53460648158	23.814259759169	
0	GRMZM2G436710 P0	26.63947005.0985	5 26.987070205132	26.6817893881	26.730416888372	
1	GRMZM2G054300 P0	25.7625453220619	24.665920528769	23.48761934929	26.052059230537	
2	GRMZM2G027640 P0	24.0448689271279	24.526046791103	24.35159999402	24.200810282812	
3	GRMZM2G081037 P0	21.4258564208647	21.321004583551	21.52525443746	21.530049443805	
4	GRMZM2G041381 P0	25.0993658026596	23.391370704689	24.69780528655	24.664119761677	
5	GRMZM2G139441 P0	23.2519668716281	23.132990364069	23.31100118274	23.516154221023	1
6	GRMZM2G157007 P0	26.0878234853339	25.958090187309	26.00666926119	25.906068917439	
7	GRMZM2G442129 P0	23.93691816735	5 23.998916852552	23.72573124362	23.53360844792	Γ
8	GRMZM2G135186 P0	26.8107094262947	27.248199633594	26.4042039045	26.425070510229	
9	GRMZM2G098039 P0	22.7497018824786	22.885220088261	22.31975985513	22.568085404565	
0	GRMZM2G157443 P0	25.4480690217536	24.673167816682	25.83390978048	25.745066715829	
1	GRMZM2G120579 P0	24.2267829137487	24.266909066281	24.47738574131	24.333355923869	
2	GRMZM2G097900 P0	23.5425497517129	23.632210742833	23,49074044701	23.258789061468	
3	GRMZM2G081883 P0	22.1357714896681	21.572644507788	21.6906191481	22.144071826553	
4	GRMZM2G070422 P0				26.394876538092	
15	GRMZM2G127361 P0				24.200322734689	
6	GRMZM2G178415 P0		3 25.011039050724		25.110140218141	
7	GRMZM2G473001 P0		24.641219694859			
8	GRMZM2G003765 P0		22.332993196383		24.088512888843	
9	GRMZM2G175134 P0				25.173408387027 ~	
	0111120170104 PV	23.213133032012	24.732031303112	20.21110200024	<	
•	♦ ► ► + groupQ	uantification				
8	Find	v ~ ~	Find All 📄 Forma	atted Display 📃 Ma	atch Case 🔍	
	et1of1 mp1	English (USA)	I_ 🖺	Average: ; Sum: 0		1009

In the metadata template file, MS run acquisitions were grouped using the value of the *Condition* variable. The grouping of the MS runs involve thus four groups (different protein solubilization methods). This means that the proteins will be quantified in each group of MS run acquisitions.

FIGURE 7.25: SPREADSHEET VIEW OF THE QUANTIFICATION BY GROUPS RESULTS

7.2.7 RUNNING THE MSstats GNU R AND RMARKDOWN SCRIPTS

The workflow as has developed since the beginning to this *MSstats* work session has been recorded both in the form of a pure *GNU R* script and as a RMarkdown script. By clicking onto the *Rscript* workflow item (FIG-URE 7.26, "LOAD THE GNU R AND RMARKDOWN SCRIPTS"), the user is presented with two options: load the *GNU R* script and/or the RMarkdown script. The RMarkdown script might be run in the *RStudio* (*RStudio* has changed its name to become *Posit*) environment.

0	*
	MSstats working directory
	Load
	MCQR dubious data filter
	MSstats data processing
	MSstats quantification by samples
	MSstats quantification by groups
	Rscript
	Download R script
	Download Rmd script

Use the button of interest to download the *GNUR* or the RMarkdown script corresponding to all the workflow steps that were run up to this one.

FIGURE 7.26: LOAD THE GNU R AND RMARKDOWN SCRIPTS

Upon saving the RMarkdown version of the script, if available on the system, *i2MassChroQ* will try to load it automatically in *RStudio*.

8 Specific procedures for the timsTOF line of instruments

This chapter describes the very few specific procedures to carry out when the proteomics data at hand are from the Bruker timsTOF line of instruments.

8.1 GENERAL CONSIDERATIONS

The mass spectrometers from the timsTOF line of instruments by Bruker implement ion mobility mass spectrometry by trapping ions and subjecting them to a gas flow that moves them in the trap according to their collisional cross section. Interestingly, the outcome of that operation is the reverse of conventional drift tube-based ion mobility: large ions are released first and smaller ions are released last.

Apart from the observation above, the result is nonetheless that ions entering the instrument are separated according to a new dimension that is orthogonal to the other two retention time and m/z ratio dimensions: the ion mobility dimension. This new dimension inevitably introduces more complexity and greater volume to the mass data.

In a historic move, Bruker has decided to publish the technical details of their data format. During the acquisition, data are stored in two separate files located in their data directory that has the <u>..</u> extension:

- <u>analysis.tdf</u>: this file is a *SQLite3* relational database that contains all the metadata about the acquisition. The generic *metadata* term defines data that describe the actual data. So this file contains data that explain how the data are organized in the actual data file below.
- <u>analysis.tdf_bin</u>: this file is a binary-format file that holds the data in the form of a succession of numbers packed according to a specific scheme that Bruker has decided to make public.

The *SQLite3* <u>analysis.tdf</u> file contains a set of tables. Most often, records in one table make reference (*relate*) to other record(s) in other table(s). This is why this database file is said to be a *relational database*. A view of the tables making that relational database file is shown in FIGURE 8.1, "VIEW OF THE RELATIONAL DATABASE FILE".

Create Table	Execute SQL	Edit Database Cell 🔷
Create Table 🛛 😽 Create Index 📑 Modify Ta		
v	ble 🔢 Delete Table 🚔 Print	Mode: Text 🗸 🎲 📄 🗐 🎼 🧹
e : Ty	pe ^E Schema	1
Tables (17)		
CalibrationInfo	CREATE TABLE CalibrationInfo (KeyPolarity CHAR(1) CHECK (KeyPolarity IN ('+', '-')), KeyName	TEX
DiaFrameMsMsInfo	CREATE TABLE DiaFrameMsMsInfo (Frame INTEGER PRIMARY KEY, WindowGroup INTEGER N	OT N
DiaFrameMsMsWindowGroups	CREATE TABLE DiaFrameMsMsWindowGroups (Id INTEGER PRIMARY KEY)	
DiaFrameMsMsWindows	CREATE TABLE DiaFrameMsMsWindows (WindowGroup INTEGER NOT NULL, ScanNumBegin	INT
III ErrorLog	CREATE TABLE ErrorLog (Frame INTEGER NOT NULL, Scan INTEGER, Message TEXT NOT NUL	L)
FrameMsMsInfo	CREATE TABLE FrameMsMsInfo (Frame INTEGER PRIMARY KEY, Parent INTEGER, TriggerMass	REA
FrameProperties	CREATE TABLE FrameProperties (Frame INTEGER NOT NULL, Property INTEGER NOT NULL, V	/alue
Frames	CREATE TABLE Frames (Id INTEGER PRIMARY KEY, Time REAL NOT NULL, Polarity CHAR(1) CHI	ECK
GlobalMetadata	CREATE TABLE GlobalMetadata (Key TEXT PRIMARY KEY, Value TEXT)	Type of data currently in cell Apply
GroupProperties	CREATE TABLE GroupProperties (PropertyGroup INTEGER NOT NULL, Property INTEGER NOT	Size of data currently in table
MzCalibration	CREATE TABLE MzCalibration (Id INTEGER PRIMARY KEY, ModelType INTEGER NOT NULL, Digi	itizer DB Schema 🔷
PasefFrameMsMsInfo	CREATE TABLE PasefFrameMsMsInfo (Frame INTEGER NOT NULL, ScanNumBegin INTEGER N	IOT I Name
Precursors	CREATE TABLE Precursors (Id INTEGER PRIMARY KEY, LargestPeakMz REAL NOT NULL, Averag	ieMz
PropertyDefinitions	CREATE TABLE PropertyDefinitions (Id INTEGER PRIMARY KEY, PermanentName TEXT NOT NU	VILL,
PropertyGroups	CREATE TABLE PropertyGroups (Id INTEGER PRIMARY KEY) WITHOUT ROWID	CalibrationInfo
Segments	CREATE TABLE Segments (Id INTEGER PRIMARY KEY, FirstFrame INTEGER NOT NULL, LastFrar	> DiaFrameMsMsInfo
TimsCalibration	CREATE TABLE TimsCalibration (Id INTEGER PRIMARY KEY, ModelType INTEGER NOT NULL, CO	0, C1 DiaFrameMsMsWindowGroups
Indices (4)		> DiaFrameMsMsWindows
FramesTimeIndex	CREATE UNIQUE INDEX FramesTimeIndex ON Frames (Time)	> ErrorLog
PasefFrameMsMsInfoPrecursorIndex	CREATE INDEX PasefFrameMsMsInfoPrecursorIndex ON PasefFrameMsMsInfo (Precursor)	FrameMsMsInfo
PrecursorsParentIndex	CREATE INDEX PrecursorsParentIndex ON Precursors (Parent)	FrameProperties
PropertyDefinitionsIndex	CREATE UNIQUE INDEX PropertyDefinitionsIndex ON PropertyDefinitions (PermanentName))
Views (1)		GlobalMetadata
Properties	CREATE VIEW Properties AS SELECT s.Id Frame, pd.Id Property, COALESCE(fp.Value, gp.Value) \	Value
Triggers (0)		MzCalibration
		PasefFrameMsMsInfo Precursors

The mass spectrometers of the tims TOF line of instruments by Bruker produce mass data that are stored in two files. This figure shows the table structure of the relational database <u>analysis.tdf</u> file displayed in *Sqlite-Browser* (a Free Software application).

FIGURE 8.1: VIEW OF THE RELATIONAL DATABASE FILE

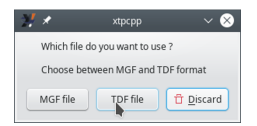
When dealing with proteomics projects that have their data originating in timsTOF instruments, some specific steps are to be taken so as to inform *i2MassChroQ* that specific handling is required. These will be reviewed below in the same succession as they need be implemented when running *i2MassChroQ*.

8.1.1 RUNNING X! Tandem IDENTIFICATIONS WITH BRUKER TIMS TOF DATA

It is possible to load Bruker tims TOF data right in the *i2MassChroQ* program's graphical user interface, as shown in the window pictured in FIGURE 3.2, "X!TANDEM-BASED IDENTIFICATION CONFIGURATION". The very first specific step to take in this case is to select the data files by clicking the *Add Bruker timsTOF folders*.

The *Add Bruker timsTOF folders* button lets the user choose the Bruker data directory (<u>...d</u> extension) and then asks if the data to be loaded are in the TDF or MGF format (see FIGURE 8.2, "FILE FORMAT SELECTION DIALOG FOR BRUKER TIMSTOF DATA").

Indeed, the MGF file generated by the Bruker software is automatically installed in the <u>.d</u> extension data directory.



When handling Bruker tims TOF data, two file formats are available: MGF and TDF. See text for details.

FIGURE 8.2: FILE FORMAT SELECTION DIALOG FOR BRUKER TIMSTOF DATA

The *DataAnalysis* software from Bruker allows one to export proteomics MS/MS data into MGF format files (Mascot generic format). Their native data format, though, is the TDF format. It is important to keep in mind that the MGF format only stores MS/MS spectral data, no MS data. By using this format, *i2MassChroQ* and *MassChroQ* won't be able to access MS data, which are required in a number of situations, in particular when extracting ion currents for given m/z ratios, for example, or for area-under-the-curve quantitative proteomics. It is thus always recommended to use the native TDF format whenever available.

When loading Bruker timsTOF data right into *i2MassChroQ* as described above, the software performs some under-the-hood operations that the user might want to be aware of. The hidden operations are unveiled in the following sections, as they involve command line programs shipped along with *i2MassChroQ* that might be of interest to the user.

8.1.2 Converting Bruker timsTOF data to mZXML with mZXMLCONVERTER

X!Tandem needs the mass spectrometric data that it uses for the database searches to be in the mzXML format. For this very reason, *i2MassChroQ* cannot work by harnessing the capabilities of *X!Tandem* starting from Bruker timsTOF data. These data need to be converted to an mzXML file before they can then be fed to *X!Tandem*.

In order to be able to store a mzXML file on disk, the user may convert timsTOF data files (TDF or MGF) to mzXML using the *mzxmlconverter* program that is shipped along with *izMassChroQ*. This program is a command line program that takes a data file in input and that writes a mzXML file in output. The command line syntax is easy, as picture in FIGURE 8.3, "CONVERTING MASS DATA FILES TO MZXML DATA FILES".

To obtain help about the program, run the following:

\$ ^I <path_to>/mzxmlconverter --help RETURN

¹ The prompt character might be % in some shells, like *zsh*.



Files of any format handled by *ProteoWizard* or files from the Bruker's timsTOF line of instruments can be converted to mzXML using *mzxmlconverter*. Conversion from timsTOF format to mzXML is performed entirely by our own software.

FIGURE 8.3: CONVERTING MASS DATA FILES TO MZXML DATA FILES

WARNING: THE MZXML FILE FORMAT DOES NOT CONTAIN MOBILITY DATA FROM THE TIMSTOF DATA FILES

It is important to grasp that the mzXML file that is generated by *mzxmlconverter* does not contain all the mass data that are contained in the Bruker's timsTOF data files. When loading the produced mzXML format files in *izMassChroQ*, the *X!Tandem* program will be able to perform peptide and protein identifications. Later, however, when the user will try to activate features in *izMassChroQ* that require the original data, the software will not be able to provide the expected results, like XIC reports or ion mobility values, because it won't have access to the original data.

The *mzxmlconverter* program is practical because it allows storing the mzXML file on disk and loading it in *izMassChroQ* for *X!Tandem* to consume it for the identifications. However, as stated in the warning above, that mzXML file has not a full copy of the data in the original mass spectrometry data file (be that a mzML, or MGF, or TDF file). *izMassChroQ* has a solution for this problem: using an integrated workflow to convert the original data file to mzXML, make *X!Tandem* use it, write out the *X!Tandem* results file and finally rewrite that file into a new version by adding a connection between the *X!Tandem* run results and the original mass data file. In this way, when the user will activate features in *izMassChroQ* that need accessing the original mass data file, the expected results will be effectively displayed. This process is described in detail in the next section.

8.1.3 DATA CONVERSION PROCESS WITH BRUKER TIMSTOF TDF DATA AND TANDEMWRAPPER

In order to seamlessly use Bruker tims TOF data in the context of performing *X*.*Tandem* identifications and later *MassChroQ*-based quantifications, the *tandemwrapper* program is made available to users who want to perform database searches using the command line interface. The *tandemwrapper* program performs an under-the-hood

file format conversion as described in the previous section before automatically feeding the generated mzXML file to *X!Tandem*. After *X!Tandem* has produced its results file, that file is rewritten by *tandemwrapper* in such a manner that the connection with the original mass spectrometry data files is reinstated for further use in the *i2MassChroQ* graphical interface.

There is, however, a way to convert Bruker timsTOF data files to mzXML using a standalone program, called *tandemwrapper*, that is shipped along with *i2MassChroQ*. The *tandemwrapper* program is a command line program that takes as input a XML configuration file. The XML file is most similar to the configuration file that *X!Tandem* uses.

To obtain help about the program, run the following:

\$ <path_to>/tandemwrapper --help RETURN

A typical *tandemwrapper* input configuration file is shown below:

The XML configuration file that is provided to *tandemwrapper* on the command line is a replicate of the file that *X*.*Tandem* itself expects. That file is shown in **FIGURE 8.4**, "**CONFIGURATION FILE FOR TANDEMWRAPPER**".

	xml version="1.0" encoding="UTF-8"?
2	<bioml label="example-tandemwrapper-mass-data-file.mzxml"></bioml>
3	<note type="heading">Paths</note>
- 4	<pre><note label="list path, default parameters" type="input">/full_path_to/xtandem-presets-file.xml</note></pre>
5	<pre><note label="list path, taxonomy information" type="input">/full_path_to/database.xml</note></pre>
6	<pre><note label="spectrum, path" type="input">full_path_to/mass-data-file.mzxml</note></pre>
7	<note type="heading">Protein general</note>
8	<note label="protein, taxon" type="input">usedefined</note>
9	
10	<pre><note label="output, path" type="input">/full_path_to/tandemwrapper-output.xml</note></pre>
11	

The *tandemwrapper* program takes as input a configuration file that is most similar to the configuration file that is fed to *X*!*Tandem*.

FIGURE 8.4: CONFIGURATION FILE FOR TANDEMWRAPPER

The following elements need an explanation:

- *default parameters*: In the example, the <u>/full_path_to/xtandem-presets-file.xml</u> file is the *X!Tandem* presets file, already discussed in Section 3.3, "Setting the X!Tandem Run Presets".
- *taxonomy information*: In the example, the <u>/full_path_to/database.xml</u> file is the file that configures the location of the FASTA protein database files that are searched by *X!Tandem*. This file is described below.
- *path*: In the example, the <u>/full_path_to/mass-data-file.mzxml</u> file is the mass spectrometry data file in the mzXML format.



Tip

The mass spectrometry data file might be of any format that can be handled by ProteoWizard (open data formats only, particularly mzML) and also the Bruker's timsTOF TDF format that is handled by our own code.

• *output, path*: In the example, the <u>/full_path_to/tandemwrapper-output.xml</u> file is the file in which the *X!Tandem* configuration file is written for immediate use by *X!Tandem*.

The configuration file that indicates where the FASTA protein database files are located, that is referenced in the *tandemwrapper* input configuration file is shown in FIGURE 8.5, "CONFIGURATION FILE POINTING AT THE FASTA PROTEIN DATABASES".



This file tells *tandemwrapper* the location of the FASTA protein databases required when *X*.*Tandem* will actually perform the searches.

FIGURE 8.5: CONFIGURATION FILE POINTING AT THE FASTA PROTEIN DATABASES

The *tandwrapper* program performs the following tasks in sequence:

- Convert the input mass data file to the mzXML file format that *X!Tandem* needs to perform the database searches. This step is only performed if the original mass spectrometry data file has not the mzXML format. *X!Tandem* produces an identification results file in an XML format;
- The mzXML format that is consumed by *X!Tandem* is a pretty simple format that was not designed to store a large variety of data/metadata, like ion mobility data, for example. For this very reason, *tandemwrapper* reads the identification results file produced by *X!Tandem* (that file is also in a specific

XML format) and rewrites it to a new analogous file that has all the necessary connections to the original mass data file. In this way, when the new version of the *X!Tandem* identification results file is loaded in *i2MassChroQ* all the original mass data can be accessed to provide the user with all the data, like XIC chromatograms, ion mobility data, for example. To load the *X!Tandem* identification results, see Section 3.4, "LOADING THE PROTEIN IDENTIFICATION RESULTS".

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Version 3, 29 June 2007

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Colophon

About the authors. Filippo Rusconi is a senior research scientist at the French national research council (*Centre national de la Recherche scientifique*, CNRS). Filippo has a background in biochemistry and organic chemistry and was trained during his Ph.D. as a bioanalytical chemist. He has extensive knowledge of analytical techniques involved in the study of biopolymers.

Filippo Rusconi is the author of a handbook about mass spectrometry for biochemists (French). The book was published by the French sci/tech publisher Lavoisier (HTTPS://WWW.Lavoisier.FR) **7**.

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The frog on the book title page is a frog from Papua. This frog is able to hover when performing downwards leaps. This picture is courtesy HTTP://www.PAPUAWEB.ORG

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