PRIDE Converter 2 GUI User Manual

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1. Introduction

The PRIDE Converter 2 tool suite is a composed of 4 independent applications:

- The PRIDE Converter 2 application will convert MS search result files containing identification and spectra into PRIDE XML.
- The **PRIDE mzTab Generator** will produce skeleton mzTab files from MS search results files. These skeleton files require either manual or scripted editing to add quantitation and/or gel information.
- The PRIDE XML Filter will remove identifications or spectra from PRIDE XML files based on a series of configurable filters.
- The PRIDE XML Merger will combine several PRIDE XML files into a single one.

All tools have both a Graphical User Interface (GUI) and a Command-Line Interface (CLI). The GUIs have been designed to provide a rich, user-friendly interface while the CLIs have been developed mainly for tool and pipeline developers to be able to integrate the PRIDE Converter 2 tools in their own software to provide an efficient way to generate PRIDE XML from their own resources.

2. PRIDE Converter 2 Requirements

2.1. System Requirements

• Java: JRE 1.5 +

• CPU: 1 gigahertz (GHz) or faster 32-bit or 64-bit processor.

• Memory: 1 gigabyte (GB) RAM.

• Hard Disk: 55 MB available for installation, more if doing file conversions.

• Platform: Tested in Mac OS X, Linux, and Windows (XP, Vista, 7).

2.2. Additional Requirements

PRIDE Converter 2 requires Internet access for connection to the Ontology Lookup Service (OLS) web service and automatic access to PubMed records already published.

3. PRIDE Converter 2 Tool Suite GUI User Guide

To start the GUI, users must either double-click on the *PRIDE Converter 2* jar file or invoke the command-line without any arguments. The first screen that the user sees will be the tool selection screen. From there, the user can select which tool from the tool suite to launch (Figure 2).



Figure 2 – PRIDE Converter Tool Suite tool selection form

Each tool shares a common design framework to ensure a common and consistent look and feel across all applications (see Figure 3).

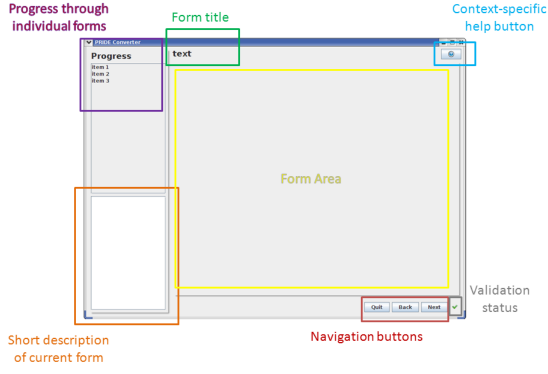


Figure 3. The Common GUI Navigation Panel.

Tools are made up of a series of forms, that can be navigated using the Next/Back/Quit buttons at the bottom right corner of the nagivation frame. Each form will provide information that is used to update the GUI: a form title, a short description of its purpose and required fields, a context-specific help page and a mechanism to validate the form content to ensure that all required fields are filled in and contain correctly-formatted values.

The result of this validation are shown next to the navigation buttons (\checkmark if all fields are OK, \triangle on warnings, \checkmark on errors). If errors are detected, the users will be shown an error dialog with more information on the validation rule that generated the error. Users will not be able to proceed with the conversion process until all errors are corrected.

PRIDE Converter 2 forms maintain a standard functionality scheme. Any field marked with a red asterisk (*) is mandatory and the user cannot proceed to the next form until all mandatory fields are filled. The button can be used to add entries. When entries are selected from a table, the button can be used to edit those entries and the button can be used to delete them.

3.1. PRIDE Converter 2

When launched in GUI mode, *PRIDE Converter 2* guides the user through a 12-step process to convert their search engine output files into well-annotated PRIDE XML files.

3.1.1. Format Selection

Input format selection is the first choice that the user must make. When selecting a format, the user will be given information on the exact file type(s) required, as well as links to the vendor and/or format specification documentation (Figure 4). Please note that Mascot conversion is not available on the Mac OS platform, due to limitations imposed by the Mascot API itself. Please refer to Table 1 for the supported formats in *PRIDE Converter 2*.

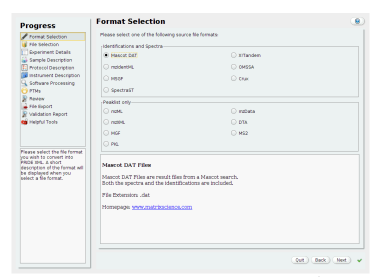


Figure 4 – PRIDE Converter 2 Format Selection form

Table 1. Supported Formats in *PRIDE Converter 2*.

Format Name	File Type	Data Content
Mascot	.dat	Spectra and Identifications
mzldentML	.xml	Spectra and Identifications
X!Tandem	.xml	Spectra and Identifications
OMSSA	.CSV	Spectra and Identifications
SpectraST	.txt	Spectra and Identifications
CRUX	.txt	Spectra and Identifications
MSGF	.txt	Spectra and Identifications
Proteome Discoverer	.msf	Spectra and Identifications
mzML	.xml	Spectra Only
DTA	.dta	Spectra Only
MGF	.mgf	Spectra Only
mzData	.xml	Spectra Only
mzXML	.xml	Spectra Only
PKL	.pkl	Spectra Only

3.1.2. File Selection

The next form is for file selection and setting *DAO* options, if applicable (Figure 5). Each format-specific *DAO* can have one or several custom options that can be set through the GUI (for example, the Mascot *DAO*, as shown in Figure 5). Sensible default options are always provided and only the basic required options are shown by default. Power users have the option to show all available options (if applicable) and those options will be stored in the report file such that the user can always review how the conversion process was configured. Tool tips will provide a description of the options and sensible/default values. Currently, only the peaklist-only *DAO*s do not have any options.

Some MS search result file formats link to spectra that are kept in external files (X!Tandem, mzIdentML and CRUX for example). For those *DAOs*, the user has the possibility to select a file containing the spectra to include in the conversion process. For the *DAOs* where externally-linked spectra are not applicable, the option to select spectra files is disabled.

By default, users will be shown a form that will only provide single-file selection dialogues but users who wish to do multiple file batch conversion can switch to an alternative view, which will allow them to select multiple files of each type (source, mzTab, spectra, etc).

When converting multiple files at once, a warning message will be displayed to the user indicating that some metadata will be copied across all files while some will not. The information that will be common across all files will include: sample annotations, post-translational modifications (PTM), protocol and instrumentation description and any other experiment-level annotations. Software processing, which can include file-specific software settings, score thresholds and false discovery rates will be taken from the individual report files associated with each input file to be converted.

Once the files have been selected, the *PRIDE Converter 2* runs in *prescan* mode to generate the report files, where the annotations that the user will input in the coming steps will be stored prior to actual conversion. Report file generation can take anywhere between seconds and several minutes and is directly proportional to the size and complexity of the input files. If several files are being converted at once, a report file needs to be generated for each input file, which will compound the time required.

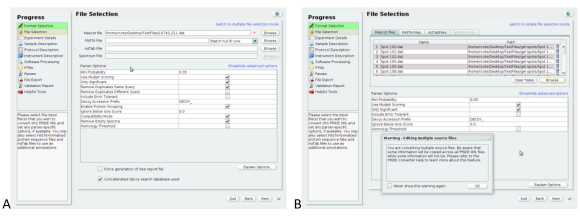


Figure 5 – *PRIDE Converter 2* File Selection form. Panel A shows the single-file selection mode while panel B shows the multi-file selection mode. Note that panel A shows the full set of options for the Mascot DAO while panel B only shows the default options.

Users have the possibility to link their search database protein sequences to the PRIDE XML files that they generate. This is particularly helpful in the case where custom-made protein sequence databases are used. The protein sequence database must be exported in FASTA format. From experience, we have observed that the protein accession that is assigned to an identification can differ significantly from the FASTA header, depending on if and how the search engine processes the protein sequence database. To solve this issue, PRIDE Converter 2 has several options on how the FASTA file should be processed during conversion.

3.1.2.1. FASTA - First word

In this scenario, the search engine assigns the first word of the FASTA ID line as the identification accession. Given a FASTA ID line that looks like the following:

```
>P30443 Full=HLA class I histocompatibility antigen, A-1 alpha chain;
```

The report file identifications, after being enriched with the FASTA information, will look like this:

The FastaSequenceReference points to a Sequence element, as illustrated below.

```
<Sequence id="8" accession="P30443">
MAVMAPRTLLLLSGALALTQTWAGSHSMRYFFTSVSRPGRGEPRFIAVGYVDDTQFVRFDSDAASQKMEPRAPWIEQEGPEYWDQETRNMKA
HSQTDRANLGTLRGYYNQSEDGSHTIQIMYGCDVGPDGRFLRGYRQDAYDGKDYIALNEDLRSWTAADMAAQITKRKWEAVHAAEQRRVYLEG
RCVDGLRRYLENGKETLQRTDPPKTHMTHHPISDHEATLRCWALGFYPAEITLTWQRDGEDQTQDTELVETRPAGDGTFQKWAAVVVPSGEEQ
RYTCHVQHEGLPKPLTLRWELSSQPTIPIVGIIAGLVLLGAVITGAVVAAVMWRRKSSDRKGGSYTQAASSDSAQGSDVSLTACKV
</Sequence>
```

3.1.2.2. FASTA - Full ID line

In this scenario, the search engine assigns the complete ID line FASTA ID line as the identification accession. This occurs when the search engine does not process or index the source FASTA database, and is especially useful when custom databases or custom identifiers are used.

The report file identifications, after being enriched with the FASTA information, will look like this:

3.1.2.3. FASTA - Match UniProt AC

If the FASTA file used contains ID lines that are generated according to the UniProtKB specification, they should look something like the following:

```
>sw|P31946|1433B_HUMAN RecName: Full=14-3-3 protein beta/alpha;AltName: Full=Protein 1054;AltName: Full=Protein kinase C inhibitor protein 1;Short=KCIP-1;Contains:RecName: Full=14-3-3 protein beta/alpha, N-terminally processed;
```

Some search engines will process these FASTA ID lines and only assign the UniProtKB protein accession. The report file identifications, after being enriched with the FASTA information, will look like this:

3.1.2.3. FASTA - Match UniProt ID

Using the same FASTA ID line as in the preceding section, some search engines will assign the UniProtKB ID. The report file identifications, after being enriched with the FASTA information, will look like this:

Note, in this last case, the presence of a *CuratedAccession* element where the correct UniProtKB accession has been linked to the UniProtKB identifier. Users are strongly urged to use UniProt accessions instead of identifiers, as entry identifiers are notoriously changeable, making data comparisons and downstream analysis more difficult and cumbersome.

3.1.3. Experimental Details

The first annotation form deals with experimental and project details (Figure 6). Users must provide a project name, an experiment title and a short label to describe the file being converted. In the case where multiple files are being converted at once, the user needs to provide a project name in this form but will have the opportunity to provide individual experiment titles and short labels in a special form which is only displayed for this purpose. Users need to provide contact information for at least one person – usually the data submitter or the primary investigator. A full name, valid e-mail address and institute contact details are required for each contact. Contacts can be saved as templates to avoid the necessity to re-enter the information at each run of the application. Users also have the option to provide a reference that supports the data being submitted to PRIDE. References can either be provided as free text or use an automatic PubMed ID lookup to retrieve the full citation, if indexed and available online. An Internet connection is required for this.

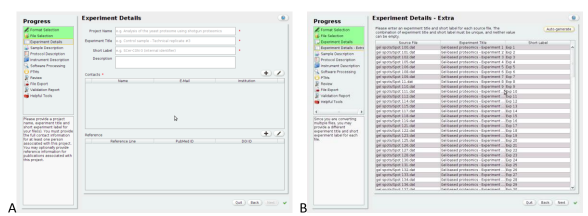


Figure 6 – *PRIDE Converter 2* Experiment Details form. Panel A shows the single-file conversion mode while panel B shows the additional step that is required to provide experiment titles and short labels for each source file to be converted.

If multiple files are being converted, the project name will be identical across all files, but the combination of experiment title and short label must be unique to each file. Users can auto-generate placeholder data, based on the project name, but are encouraged to fill in these fields with meaningful information.

3.1.4. Sample Description

The sample description form (Figure 7) allows the user to provide taxonomy, tissue type and cell type annotations using pull-down menus that contain the most commonly used values already in PRIDE. A sample name and species are required. Users still have the possibility to use the comprehensive Ontology Lookup Service (OLS) to look up alternative terms that are not already provided. If multiple files are being converted, the information in the "master report" will be automatically copied over to each file. Users will have the possibility to add file-specific controlled vocabulary (CV) terms by clicking on the "add custom sample information" button and selecting the proper source file. Files with custom annotations will be shown in bold. Any annotation coming from the master report will not be editable or deletable from the individual files' panel, but users will be able to add new terms. Any terms added or deleted from the master report view will always be propagated to all files.

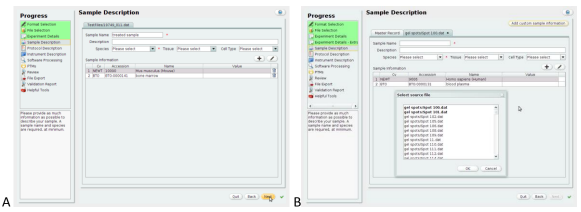


Figure 7 – *PRIDE Converter 2* Sample Description form. Panel A shows the single-file conversion mode while panel B shows the how annotation of multiple files. Note that, in panel B, the entries for the selected individual file cannot be deleted, as they were set in the 'Master Record' tab.

3.1.5. Protocol Description

In the protocol description form (Figure 8), *PRIDE Converter 2* requires a protocol name to be provided, at minimum. Protocols can have multiple steps, and each step can be described with one or several CV terms. Please look at the templates provided with the tool for examples. The "Load" button can be used to load existing templates. A popup window will be shown, where users can select the desired template from a list. To save a template once all of the information is correctly entered, users simply need to enter a name (in this case, a protocol name) and click on the "Save" button. A file will be saved to the templates folder in the installation directory of *PRIDE Converter 2*. Should there already be a file with the same name, the user will be prompted to either rename the template or allow the existing one to be overwritten.

PRIDE Converter 2 provides the possibility to save commonly-used annotations such as sample, instrumentation and protocols as templates which can be reused in subsequent conversions. A set of basic templates is provided with the tool suite. Users can load and update templates to better suit their own requirements.

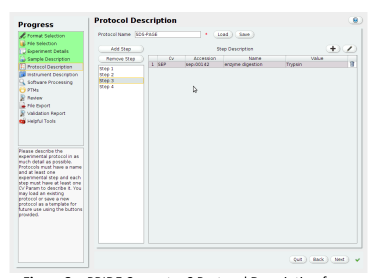


Figure 8 – *PRIDE Converter 2* Protocol Description form.

3.1.6. Instrument Description

The instrument description form (Figure 9) is another form that can use templates, as in most cases, the instrumentation available to a user will not change dramatically over time. The instrument name in mandatory. "Source", "Analyzer" and "Detector" elements need to be annotated, preferably with CV terms from the Proteomics Standards Initiative (PSI) Mass Spectrometry (MS) controlled vocabulary (http://psidev.cvs.sourceforge.net/viewvc/psidev/psi/psi-ms/mzML/controlledVocabulary/psi-ms.obo).

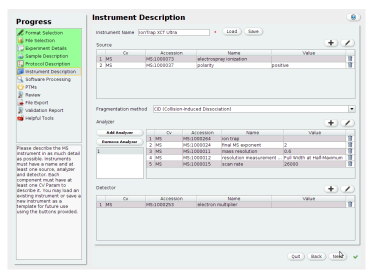


Figure 9 – PRIDE Converter 2 Instrument Description form.

3.1.7. Software Processing

The software processing form (Figure 10) report information on the software settings and the search database used. The "software" and "processing method" information will more often than not be reported directly by the DAO and should not be modified. **Users should be aware that If converting multiple files**, these two sections will be taken directly from the source file and any additions made to the GUI will not be carried across all converted files.

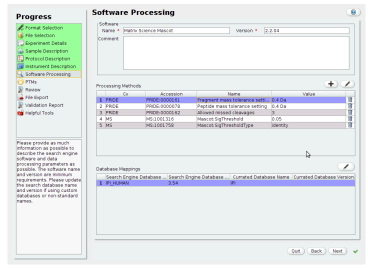


Figure 10 – *PRIDE Converter 2* Software Processing form.

The "search database" mappings will help standardize the names of the sources of the protein sequences that have been used to generate the search databases. The most common sources of protein sequences have been included (UniProtKB/Swiss-Prot, UniProtKB/TrEMBL, International Protein Index (IPI), Ensembl, RefSeq, NCBI nr database) and the users are encouraged to provide either a release number or the date the search database was built. Users can also manually enter a custom database name, if the supplied options are not appropriate. The "curated database name" is mandatory, but the "curated database version", while desired, is not. A warning will be generated if the database version is omitted however.

3.1.8. PTMs

The next step is the review and/or correction of PTM assignments (Figure 11). Please refer to the section entitled "PRIDE Converter Automatic PTM Assignment" for more information on how PTMs are assigned and how they can be manually corrected or annotated.

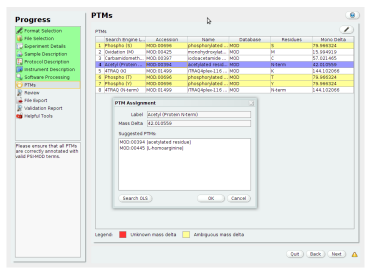


Figure 11 – PRIDE Converter 2 PTM form.

3.1.9. Review

Once all of the required annotations are provided, the GUI displays a quick review screen before proceeding to copy the metadata across all report files (Figure 12). Any warnings generated by the automatic form validation that occurs at each step will be displayed here. If multiple files are being converted at once, the total number of input files will also be shown. Users can add additional experiment-level CV terms that would reflect the experiment as a whole but are not appropriate in any of the previous sections. The DAOs and *PRIDE Converter 2* itself will automatically add params for 'XML generation software' and 'Original MS data file format'.

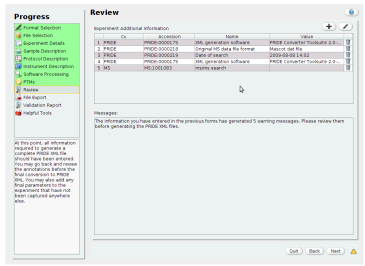


Figure 12 – PRIDE Converter 2 Review form.

Updating the report file (if performing a single conversion) or copying the data across all report files (if converting multiple files) can take anywhere between seconds and several minutes and is directly proportional to the number of files that need to be updated.

3.1.10. File Export

The next step is the generation of the PRIDE XML files (Figure 13). Alternatively, the graphical conversion process can be halted at this point, as all of the report files are now complete and validated and the conversion process can be scripted and batched using the Command Line Interface (CLI). This is generally only practical for users who need to convert a significantly large number of files and who have access to a cluster of computers where the conversions can be parallelized. In most cases, a single desktop machine with sufficient memory and disk space will be more than sufficient.

PRIDE Converter 2 can generate compressed files using the gzip algorithm to save on disk space. This is enabled by default, as is the option to remove any temporary work files, such as report files, once the application exits. PRIDE Converter 2 will always include all spectra in the generated PRIDE XML file unless the "include only identified" spectra option is selected. If the experiment contains interaction data that would benefit from being submitted to the IntAct interaction database (http://www.ebi.ac.uk/intact) at the EBI, users are encouraged to select the relevant option in the export dialog. This will add a CV term in the generated XML file that will be recognized by the submission process and the person whose contact details have been captured will be contacted by an IntAct curator for further discussion.

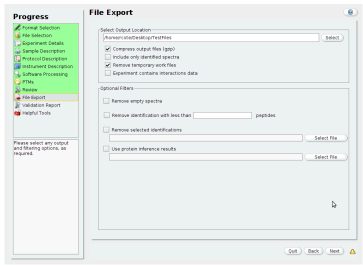


Figure 13 – *PRIDE Converter 2* File Export form.

This panel also sets the various filtering options, which are discussed in greater detail in the section entitled "PRIDE XML Filter".

Generating the PRIDE XML files can take anywhere between minutes and hours. This is proportional to the size, number and complexity of the source files. Users should ensure that they have sufficient disk space. If filtering is enabled, the full PRIDE XML file is first written to disk, then written again in its filtered form after it has been processed. The report file and the intermediate PRIDE XML file are only deleted when *PRIDE Converter 2* exits if the "remove temporary files" option is selected.

3.1.11. Validation Report

Once the PRIDE XML files are written, a status report is shown, indicating how many PRIDE XML files have been generated (Figure 14).

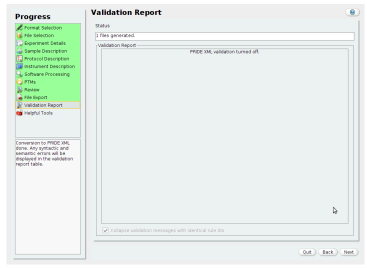


Figure 14 – *PRIDE Converter 2* Validation Report form.

3.1.12. Helpful Tools

The final screen (Figure 15) of the GUI invites users to review the generated PRIDE XML files using the PRIDE Inspector tool (http://code.google.com/p/pride-toolsuite/wiki/PRIDEInspector) and to submit their data *via* the ProteomeXchange (PX) consortium (http://www.proteomexchange.org).

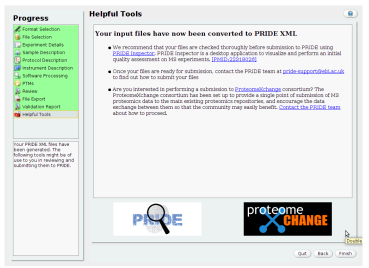


Figure 15 – *PRIDE Converter 2* Helpful Tools form.

3.2. PRIDE mzTab Generator

3.2.1. Format Selection

The input format selection form for the PRIDE mzTab Generator is identical to the one used in PRIDE Converter 2 (Figure 4), with the exception that peaklist-only formats are disabled and cannot be selected.

3.2.2. File Selection

The look and usability of the file selection form for the PRIDE mzTab Generator is almost identical to that described in the "File Selection" section of the PRIDE Converter 2 GUI user guide, with the addition of 4 mzTab-specific configuration options (Figure 16).

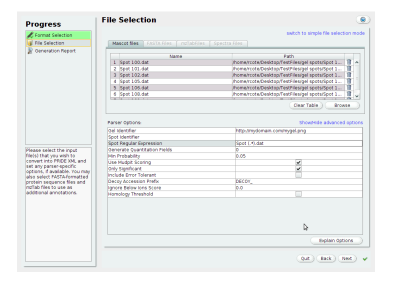


Figure 16 – PRIDE mzTab Generator File Selection form.

The DAO options, if any, will be identical to those present in the PRIDE Converter 2 and will be stored in the mzTab file. In order to avoid unpredictable DAO behaviour, users **must** use the same settings when generating the mzTab files and when using the PRIDE Converter 2. As the options are stored in both the report file and the mzTab file, validity checks will be performed to ensure that both sets of options are identical. Any differences will cause an exception and the Conversion process will be blocked until all options match.

3.2.2.1. Adding Gel-based information

The "gel identifier" option is used to set the gel identifier to be used for identifications in the generated mzTab file. This identifier should correspond to the gel's "name" in the publication (f.e. "Gel A", "Gel B" etc.). It should enable reviewers and readers to easily link the submitted data to the data presented in the corresponding manuscript.

The "gel spot identifier" option sets the gel spot identifier to be used for identifications in the generated mzTab file. This represents the gel spot's description used in the manuscript – similar to the "gel identifier" for the gel itself. Expected values would be "Spot 1", "Spot 2", etc. Together with the "gel identifier" it is therefore possible for the reviewer and reader to unambiguously link findings presented in the manuscript to specific identifications reported in the PRIDE XML file. This option is ignored if the "gel spot regex" option is set. The "gel spot regex" option is used to extract the gel spot identifier based on the source file's name. The value of this option must correspond to a valid Java regular expression. The first matching group in the pattern will be used as a spot identifier.

Both of these fields can also be added manually to an existing mzTab file: the "gel spot identifier" is mapped to the "opt_gel_identifier" column and the "gel spot identifier" to the "opt_gel_spotidentifier" column in the mzTab file. Additional gel specific information can be supplied using the following optional columns in the mzTab file:

opt_gel_url The gel's URL to use as a link.

opt_ycoord_pixelopt_xcoord_pixelopt_mwY coordinates in pixel of the spot in the gelX coordinates in pixel of the spot in the gelobserved) molecular weight.

opt_pi The protein's (observed) pl.

Once these columns are present in the supplied mzTab file, PRIDE Converter will automatically add the corresponding values to the PRIDE XML file. For more information on mzTab and on optional columns in mzTab files please refer to the mzTab format specification at http://mztab.googlecode.com.

3.2.2.2. Adding quantitative data

The "generate quant fields" option will add empty placeholder quantitative fields to the generated mzTab file for the number of specified labels. These fields and columns will need to be manually or programmatically filled in with the proper values before the mzTab file is ready to be used by the PRIDE Converter 2. If, for example, a 4-plex iTRAQ approach was used in which 4 different samples were labelled, the "generate quant fields" should be set to "4".

As a result, the mzTab generator will automatically add the required meta-data fields to the generated mzTab file:

- A placeholder for the used quantification method ("[UNIT ID]-quantification method")
- A placeholder for the used protein quantification unit ("[UNIT_ID]-protein-quantification_unit")

A placeholder for the used peptide quantification unit ("[UNIT_ID]-peptide-quantification_unit")

Additionally, so-called "subsamples" will be added to the mzTab file for every label. These subsamples will contain place holders to describe the used label as well as the characteristics of the labelled (biological) sample:

- A placeholder for the used quantification reagent
- A placeholder for the human readable description of the labelled sample (f.e. "healthy control")

The way these data is represented in the mzTab file is documented in detail in the mzTab format specification (http://mztab.googlecode.com).

At last, the mzTab generator will also add columns to the mzTab file's protein and peptide section to hold the actual measured quantitative values for each protein and peptide identified in the MS run. The user must then manually or preferably automatically insert the actual quantification values into this mzTab file. Once PRIDE Converter is supplied with an mzTab file containing such quantitative information, these data will automatically be reported in the generated PRIDE XML file.

A step-by-step description of how to generate quantitative mzTab file using the PRIDE mzTab Generator can be found in the PRIDE Converter 2 wiki: https://code.google.com/p/pride-converter-2/wiki/QuantitativeMzTabFiles.

3.2.3. Generation Report

The final screen of the PRIDE mzTab Generator (Figure 17) will show the path of the newly generated mzTab file corresponding to each input file. If a gel identifier and/or a spot identifier or spot regex have been provided, the relevant information is also included in the report. This is particularly useful when a gel spot regular expression has been used to confirm that the correct spot identifier has been extracted from the input file name.

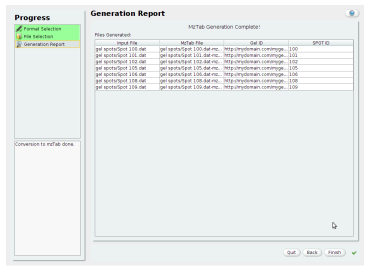


Figure 17 – PRIDE mzTab Generator Generation Report form.

3.3. PRIDE XML Filter

3.3.1. Filter Information

The first screen of the PRIDE XML Filter provides an overview of the role of the tool and informs the user of several design assumptions and restrictions. All the PRIDE XML files must be valid according to the PRIDE XML schema. Any number of filters can be selected in one go and all the filters will be applied to each selected input file. Some of the supported use cases are:

- Remove contaminants from the list of reported proteins (e.g. keratins).
- Remove decoy identifications if a concatenated target-decoy database was used.
- Provide a way to match more closely the final list of reported proteins to what is actually reported in the manuscript taking into account the protein inference problem.
- Remove protein identifications that contain less than a specified number of peptide IDs
 (useful to remove "one-hit wonders", for example). Please note that If a reported
 identification does not contain any supporting peptides after peptide filtering, it will be
 omitted from the final PRIDE XML file.
- Remove empty spectra or include only the identified spectra in the submission.

3.3.2. File Selection

The look and usability of the file selection form for the PRIDE XML Filter is almost identical to that described in the "File Selection" section of the PRIDE Converter 2 GUI user guide, with the exception that all other file type selectors have been disabled and there are no configuration options to set. The PRIDE XML Filter can either take compressed (gzipped) or uncompressed PRIDE XML files as input, but all files need to be in the same state.

3.3.3. File Export

The look and usability of the file export form for the PRIDE XML Filter is almost identical to that described in the "File Export" section of the *PRIDE Converter 2* GUI user guide, with the exception that some options have been disabled (Figure 18).

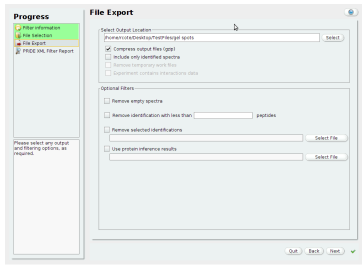


Figure 18 – PRIDE XML Filter File Export form.

The "include only identified spectra" filter is the only filter that does not need to generate an intermediate PRIDE XML file, as this can be done at the *DAO* level. All other filters will first write a complete PRIDE XML file, as normal, then post-process this file to generate the fully-filtered PRIDE XML file. This will have consequences on the time and disk space required to fully produce the final file.

The "remove empty spectra" filter will remove any spectra that do not contain any peaks.

The "remove identification with less than X peptides" filter will scan and remove any protein identification from the PRIDE XML file that contains less peptides than the number given as parameter.

The "remove selected identifications" filter requires a list of contaminant protein identifications in the form of a text file with one protein accession per line. It will use this blacklist to remove corresponding identifications from the XML file where the submitted protein accession identically matches one of the blacklisted protein identifiers.

The "use protein inference results" filter also requires a list of protein identifications in the same manner as the previous filter but works in the complete opposite fashion. Only identifications found in the whitelist are retained in the final PRIDE XML file. This is important taking into account how PRIDE Converter 2 reports peptide to protein assignments (for more details see section 7).

- PRIDE Converter 2 reports all possible peptide-to-protein assignments.
- *PeptideItems* are duplicated in different proteins in the PRIDE XML files but reference the same spectrum in the mzData section of the file.
- PeptideItems are duplicated to report ambiguous modification positions (again referencing the same spectrum).

So, using the "use protein inference results" filter, users can provide a list of the proteins that are reported in the manuscript.

3.3.4. Filter Report

The final screen of the PRIDE XML Filter (Figure 19) will show the path of the newly generated filtered PRIDE XML file corresponding to each input file.

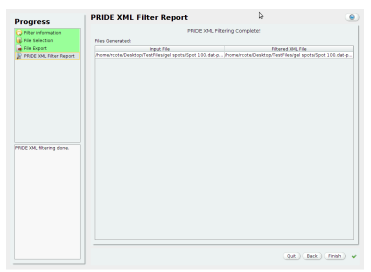


Figure 19 – PRIDE XML Filter Report form.

3.4. PRIDE XML Merger

3.4.1. Merger Information

The first screen of the PRIDE XML Merger provides an overview of the role of the tool and informs the user of several design assumptions and restrictions. All the PRIDE XML files must be valid according to the PRIDE XML schema. A master file will be selected, which will serve as the basis for the final PRIDE XML file in terms of metadata. The identifications and spectra from the other files will be merged into the final output file. Users should note that the spectrum IDs from the source files will not necessarily be kept, but the internal references between peptide-spectrum_refs will be maintained.

3.4.2. File Selection

The look and usability of the file selection form for the PRIDE XML Merger is almost identical to that described in the "File Selection" section of the PRIDE Converter 2 GUI user guide, with the exception that all other file type selectors have been disabled and the only DAO option available will be the option to compress the final output file (Figure 20). The PRIDE XML Merger can either take compressed (gzipped) or uncompressed PRIDE XML files as input, but all files need to be in the same state.

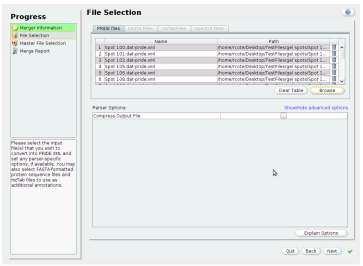


Figure 20 – PRIDE XML Merger File Selection form.

3.4.3. Master File Selection

A master file must be selected. All metadata annotations will be taken from this file to as a basis to generate the merged PRIDE XML file (Figure 21).

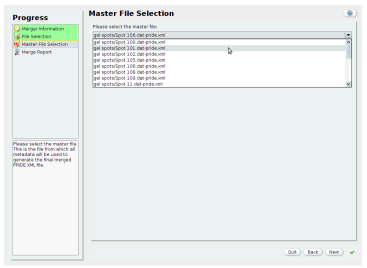


Figure 21 – PRIDE XML Merger Master File Selection form.

3.4.4. Merger Report

Once all PRIDE XML files have been merged, the user is shown a report form showing where the output file was written to and all the source PRIDE XML files which were used for the merge (Figure 22).

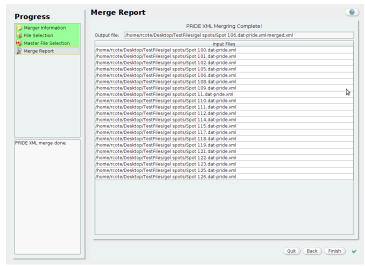


Figure 22 – PRIDE XML Merger Report form.

4. Troubleshooting

4.1. The Application Selector window appears but no tools start when a button is clicked

When *PRIDE Converter 2* starts in the GUI mode, the first screen that is displayed is the application selector (Figure 3). In rare occasions, it is possible that nothing happens when the users click on a button to start a specific application (*Converter, Filter, Merger, mzTab generator*). This is generally a problem caused with improper Java configuration. Please ensure that JAVA_HOME is correctly set in your environment and that the java runtime executable file is in your path. Please refer to the Java documentation on how to properly install and configure the Java Runtime Environment (http://java.com/en/download/help/download options.xml).

When a button is clicked from the application selector, a new process is launched based on the settings found in the *converter.properties* file that is located in the same directory as the *PRIDE Converter 2* jar file.

If JAVA_HOME isn't properly set, the new process might not properly start. One way to circumvent this problem is to update the *java.home* setting in the properties file. You need to include the full path to the java executable, including the final slash, but do not include the java application itself.

```
java.home=/home/rcote/dev/jdk1.6.0_13/bin/ - correct for linux and Mac OS X java.home=c:\Program Files\jdk1.6.0_13\bin\ - correct for Windows
```

If this still does not work correctly, try starting *PRIDE Converter 2* from the command line, as useful debugging information is logged to the console window.

```
rcote@bobble: target$ java -jar pride-converter-2.0-SNAPSHOT.jar

Reading properties file: ./converter.properties
Bootstrap command: /home/rcote/dev/jdk1.7.0_03/jre/bin/java -Xms128M -Xmx4000M -cp pride-
converter-2.0-SNAPSHOT.jar
uk.ac.ebi.pride.tools.converter.gui.ConverterApplicationSelector$ConverterLauncher

Name of the OS: Linux
Version of the OS: 2.6.32-41-generic
Architecture of The OS: amd64
Validator Loaded
```

This information can be communicated with the PRIDE helpdesk to help fix the problem.

4.2. A progress window appears but the application seems to hang

If you are converting a large number of files or files that are very big or contain a large number of spectra, the conversion process can take anywhere from minutes to hours. This is normal and the time required will be in direct proportion to the number, size and complexity of the source files. The GUI will always show message dialogs if exceptions occur. If the application still seems to hang but no error message is shown, users should look in the log file that is in the *log* directory where the *PRIDE Converter 2* was installed.

4.3. The application runs out of memory

The *PRIDE Converter 2* tool suite was written to be as memory-efficient as possible. However, if converting a large number of big files, it is always possible to run out of memory. If this should happen, please update the *jvm.args* setting in the *converter.properties* file.

4.4. The application cannot use the OLS

The *PRIDE Converter 2* tool suite requires a working internet connection to connect to the OLS web service to perform CV term related queries. If you are on a computer that requires a proxy configuration to access external network resources, you will need to uncomment and update the five *http.proxy* settings in the *converter.properties* file.

5. PRIDE Converter 2 Automatic PTM Assignment

PRIDE Converter 2 attempts to assign the correct PTM annotation based on a curated list of the most commonly observed PTMs and the mass delta as reported by the search engine. Please refer to Table 2 below for the full list of automatically assigned PTMs. If a unique PTM can be assigned to a mass delta within a 0.1 Da mass tolerance, the annotation is automatically shown to the user (Figure 16).

In cases where multiple PTMs can be assigned to a mass delta with a precision of 0.1 Da, *PRIDE Converter 2* will try to locate a unique PTM assignment to within 0.01 Da. If a unique match is found at the higher precision threshold, it will be assigned but the GUI will report the fact that multiple PTMs have been observed. In the case that multiple PTMs are still found at the higher precision threshold, no mapping is done. The GUI will report the fact that multiple PTMs have been observed within the mass tolerance window by highlighting the conflict in yellow. PTMs that have not been automatically assigned will be highlighted in red (Figure 16A).

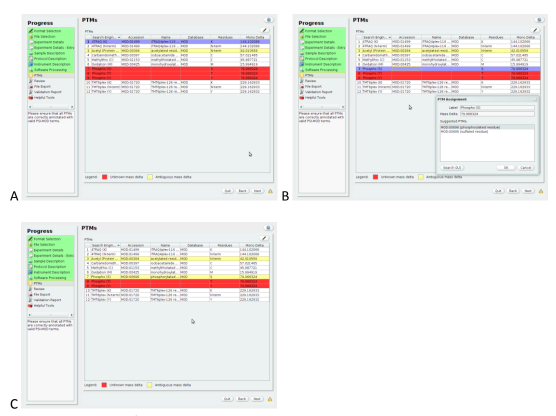


Figure 16. Display of automatically assigned PTMs in the *PRIDE Converter* 2 GUI. Rows with a white background have been unambiguously assigned. Rows in yellow have been assigned, but there is ambiguity in a 0.1 Da window around the reported mass delta. Rows in red have could not be assigned.

By double-clicking on a row highlighted in red, the user will be presented with a window showing the preferred PTMs (those that map to a mass delta with a precision of 0.1 Da), if any. The user has the choice to either select a preferred PTM, if applicable, or to search the PSI-MOD ontology via the OLS (Figure 16B). See Table 2 to see how the different mass deltas are mapped to the PSI-MOD ontology.

Table 2. List of PTMs that *PRIDE Converter 2* will attempt to automatically assign, based on the mass delta reported in the PSI-MOD ontology.

Accession MOD:00394 MOD:00674 MOD:01885 MOD:00696 MOD:00397 MOD:00398	1 2 3 21 4 5	acetylated residue amidated residue biotinylated residue phosphorylated residue	Mass Delta (Da) 42.01057 -0.98402 226.07760
MOD:00674 MOD:01885 MOD:00696 MOD:00397	2 3 21 4 5	amidated residue biotinylated residue phosphorylated residue	-0.98402 226.07760
MOD:01885 MOD:00696 MOD:00397	3 21 4 5	biotinylated residue phosphorylated residue	226.07760
MOD:00696 MOD:00397	21 4 5	phosphorylated residue	
MOD:00397	4 5		70 0000
	5		79.96633
MUD:00308		iodoacetamide derivatized residue	57.02146
10100.00336		carbamoylated residue	43.00581
MOD:00399	6	iodoacetic acid derivatized residue	58.00548
MOD:00400	7	deamidated residue	0.98402
MOD:00403	10	homoserine	-29.99281
MOD:00404	11	homoserine lactone	-48.00337
MOD:00410	17	S-(N-isopropylcarboxamidomethyl)-L-cysteine	99.06841
MOD:00704	23	dehydrated residue	-18.01057
MOD:00417	24	S-carboxamidoethyl-L-cysteine	71.03711
MOD:00419	26	(R)-5-oxo-1,4-tetrahydrothiazine-3-carboxylic acid	39.99492
MOD:00425	35	monohydroxylated residue	15.99492
MOD:01160	28	deaminated residue	-17.02655
MOD:01813	29	morpholine-2-acetylated residue	127.06333
MOD:00423	30	monosodium salt	21.98194
MOD:00424	31	S-pyridylethyl-L-cysteine	105.05785
MOD:00599	34	monomethylated residue	14.01565
MOD:01153	39	methylthiolated residue	45.98772
MOD:00695	40	sulfated residue	79.95682
MOD:00127	42	N6-lipoyl-L-lysine	188.03296
MOD:00437	44	farnesylated residue	204.18780
MOD:00438	45	myristoylated residue	210.19837
MOD:00128	46	N6-pyridoxal phosphate-L-lysine	229.01401
MOD:00440	47	palmitoylated residue	238.22967
MOD:00441	48	geranylgeranylated residue	272.25040
MOD:00159	49	O-phosphopantetheine-L-serine	340.08579
MOD:00697	50	flavin modified residue	783.14149
MOD:00445	52	L-homoarginine	42.02180
MOD:00493	122	formylated residue	27.99492
MOD:00405	12	Applied Biosystems original ICAT(TM) d8 modified cysteine	450.27521
MOD:00406	13	Applied Biosystems original ICAT(TM) d0 modified cysteine	442.22499
MOD:00480	105	Applied Biosystems cleavable ICAT(TM) light	227.12699
MOD:00481	106	Applied Biosystems cleavable ICAT(TM) heavy	236.15719
MOD:00544	188	6x(13)C labeled residue	6.02013
MOD:00546	193	(18)O label at both C-terminal oxygens	4.00849
MOD:01152	299	carboxylated residue	43.98983

MOD:00428	425	dihydroxylated residue	31.98983
MOD:01499	214	iTRAQ4plex-116 reporter+balance reagent acylated residue	144.10206
MOD:01549	730	iTRAQ8plex-116 reporter+balance reagent acylated residue	304.20536
MOD:01720	737	TMT6plex-126 reporter+balance reagent acylated residue	229.16293

6. PRIDE Converter 2 Protein Inference Implementation

Protein inference as well as modification ambiguity cannot be modelled fully in PRIDE XML files. PRIDE Converter 2 still tries to preserve this vital information of a proteomics result set following the here explained convention:

- PRIDE Converter 2 reports all possible peptide-to-protein assignments.
- *PeptideItems* are duplicated in different proteins in the PRIDE XML files but reference the same spectrum in the mzData section of the file.
- PeptideItems are duplicated to report ambiguous modification positions (again referencing the same spectrum).

6.1. Multiple peptide assignments

One problem in bottom-up proteomics approaches is the assignment of peptide identifications to proteins. Often, because of sequence similarity, a given peptide sequence cannot be unambiguously assigned to a single protein but only to multiple proteins.

In PRIDE XML file every peptide item references a given spectrum in the mzData section of the PRIDE XML file:

This feature is used by *PRIDE Converter 2* to report peptides that can be assigned to multiple proteins.

Given the following example where the peptide ABC can be assigned to two given proteins: P12345 and P54321. To represent this ambiguous assignment *PRIDE Converter 2* creates two *PeptideItems* (one for each protein) which both reference the same spectrum:

As a default setting, *PRIDE Converter 2* reports every possible peptide-to-protein assignment and does not use any specific protein inference algorithm.

6.2. Ambiguous peptide modifications

Some search engines are able to report ambiguous modification positions in peptides. The PRIDE XML format was not originally designed to support this use case. In order to provide this information *PRIDE Converter 2* uses two approaches:

- Same approach as with ambiguous protein assignments to report ambiguous peptide modification positions.

For example, assume that a peptide is identified with a sequence of MELAYM and is reported to have an oxidized methionine - either the 1st or the last. In this case *PRIDE Converter 2* again will generate two *PeptideItems* both referencing the same spectrum in the mzData section of the PRIDE XML file but with a methionine oxidation once on the first and once on the last methionine.

```
<?xml version="1.0" encoding="UTF-8" standalone="yes"?>
<GelFreeIdentification>
    <Accession>P12345</Accession>
    <PeptideItem>
       <Sequence>MELAYM</Sequence>
<SpectrumReference>10</SpectrumReference>
       <ModificationItem>
         <ModLocation>1</ModLocation>
<ModAccession>MOD:00425</modAccession>
         <ModDatabase>MOD</ModDatabase>
       </ModificationItem>
    <PeptideItem>
       .
<Sequence>MELAYM</Sequence>
       <SpectrumReference>10</SpectrumReference>
       <ModificationItem>
         <ModLocation>5</ModLocation>
<ModAccession>MOD:00425</ModAccession>
         <ModDatabase>MOD</ModDatabase>
</GelFreeIdentification>
```

- Add the reliability of the modification position using additional CV params, provided that this information is made available by the DAO.

For example, the following snippet was generated using the ProteomeDiscoverer DAO:

```
<?xml version="1.0" encoding="UTF-8" standalone="yes"?>
<GelFreeIdentification>
    <Accession>P12345</Accession>
    <PentideTtem>
           <Sequence>VPQLEIVPNSAEER</Sequence>
           <Start>106</Start>
           <End>119</End>
            <SpectrumReference>1124</spectrumReference>
            <ModificationItem>
                 <ModLocation>10</ModLocation>
                 <ModAccession>MOD:00696</ModAccession>
<ModDatabase>MOD</ModDatabase>
                 <ModDatabaseVersion>1.010.7</modDatabaseVersion>
<ModDatabaseVersion>1.010.7</modMonoDelta>79.966331</modMonoDelta>
                 <ModAvgDelta>79.9799</ModAvgDelta>
                 <additional>
                      <<toolar:

<<toolar:

<<toolar:

<<toolar:</pre>
<<toolar:</pre>
<<toolar:</p>

                 </additional>
           </ModificationItem>
           <additional>
                 <cvParam cvLabel="MS" accession="MS:1001969"
                 name="ProteomeDiscoverer:phosphoRS score" value="152.07938"/>
<cvParam cvLabel="MS" accession="MS:1001970"
                          name="ProteomeDiscoverer:phosphoRS sequence probability" value="1.0"/>
           </additional>
      </PeptideItem>
```

</GelFreeIdentification>

6.3. Adding protein inference algorithms to PRIDE Converter 2

During the conversion process *PRIDE Converter 2* generates a report file for every input file. These report files contain every possible peptide-to-protein assignment. Any change made to these assignments in the report file (i.e. removing a peptide from a protein) will be reflected in the generated PRIDE XML file.

Therefore, it is possible to plug-in protein inference algorithms that simply process the generated report files.