The pep2pro database was built to support effective high-throughput proteome data analysis. Its database schema allows the coherent integration of search results from different database-dependent search algorithms and filtering of the data including control for unambiguous assignment of peptides to proteins. The capacity of the pep2pro database has been exploited in data analysis of various Arabidopsis proteome datasets. The diversity of the datasets and the associated scientific questions required thorough querying of the data. This was supported by the relational format structure of the data that links all information on the sample, spectrum, search database, and algorithm to peptide and protein identifications and their post-translational modifications. After publication of datasets they are made available on the pep2pro website at www.pep2pro.ethz.ch. Further, the pep2pro data analysis pipeline also handles data export to the PRIDE database (http://www.ebi.ac.uk/pride) and data retrieval by the MASCP Gator (http://gator.masc-proteomics.org/). The utility of pep2pro will continue to be used for analysis of additional datasets and as a data warehouse. The capacity of the pep2pro database for proteome data analysis has now also been made publicly available through the release of pep2pro4all, which consists of a database schema and a script that will populate the database with mass spectrometry data provided in mzIdentML format.

Keywords: database, mzIdentML, pep2pro, plant proteomics, standard format

THE pep2pro PUBLIC WEB INTERFACE

The public web interface of the pep2pro database is available at www.pep2pro.ethz.ch. Here, we provide all information contained in the published Arabidopsis datasets that have been analyzed using the pep2pro data analysis pipeline (Baerenfaller et al., 2008, 2011; Grobel et al., 2009, 2011; Bosch et al., 2011; Gfeller et al., 2011). This includes information on all identified proteins, their identified peptides, and whole genome hits, as well as their proteogenomic mapping. The information also includes the number of spectra per peptide together with the scores of peptide spectrum assignments, and the display of the spectra in a Spectrum Viewer. A normalized version of the Spectrum Summary displaying for each protein the number of spectra per organ has been added recently to account for the bias that is introduced based on the different total number of spectra available for the different organs. The data in pep2pro are exported to the PRIDE database (Vucinco et al., 2010) and the information on spectral counts in the different organs is also provided to MASCP Gator2 (Joshi et al., 2011). In MASCP Gator the information is linked to the pep2pro database. Following this link, all the datasets containing spectra for a given AGI are listed to allow the selection of the dataset that is of interest to the user. Additional functions are now available on the pep2pro website. First, peptides that are detectable in a complex protein sample and that unambiguously identify a protein are displayed. Second, it is now possible to search the database for specific peptide amino acid sequences. Third, all unambiguous theoretical tryptic peptides with minimum length of six amino acids and a molecular mass between 400 and 6000 Da are now provided for a protein of interest. We added this functionality to the database in response to valuable user input. The new functions are useful in the set-up of selected reaction monitoring (SRM) experiments (Lange et al., 2008), in which it is important to select peptides that unambiguously identify a protein. Information on unambiguous theoretical peptide sequences will also help users to assess if reported protein expression evidence for a protein of interest was based on spectrum assignments to unambiguous peptide sequences or whether expression evidence was solely based on a peptide sequence that could also belong to other proteins.

THE pep2pro DATABASE

While the different functionalities of the pep2pro website provide useful information in the pep2pro database to the plant community, the pep2pro database system is most powerful for efficient and effective proteome data analysis.

ANALYSIS OF HIGH-THROUGHPUT MASS SPECTROMETRY DATA USING pep2pro

The state of data of a typical mass spectrometry-based proteomics experiment is illustrated in Figure 1. When analyzing and interpreting the results of such an experiment, the data will be queried in different ways depending on the scientific questions of the experiment. For example, in the organ-specific proteome maps...
FIGURE 1 | State of data in a typical mass spectrometry experiment. In a mass spectrometry-based experiment different samples are analyzed, usually involving different extracts of each sample and different fractions of each extract. When injected several times into the mass spectrometer, this results in a significant multiplication of mass spectrometry runs per experiment. The mass spectrometry data are then subjected to one or several database-dependent peptide assignment algorithms. In the search results, one to several spectra are assigned to a peptide sequence and one to several peptides are assigned to a protein sequence. PTM, post-translational modification.

of the AtProteome and pep2pro datasets we assessed how many proteins were identified in each plant organ based on the number of peptide spectrum assignments and which proteins were identified only in one organ. We also determined which peptides were identified best in which organ and extract, as sample context was shown to determine peptide detectability (Baerenfaller et al., 2008, 2011). For the characterization of whole genome hits and post-translational modifications we used PepSplice (Roos et al., 2007), for which we determined the optimal search parameters by calculating local false discovery rates based on the number of peptide spectrum assignments against a decoy database (Elias and Gygi, 2007; Baerenfaller et al., 2008). In the Arabidopsis phosphoproteomics datasets, which were obtained following affinity chromatography on IMAC and TiO2, we searched for phosphoproteome differences between end-of-day and end-of-night by integrating the search results from two different search algorithms (Reiland et al., 2009). In wild type and stn8 kinase mutants we identified STN8 kinase substrates (Reiland et al., 2011). To investigate the extent to which jasmonates control wound-induced re-patterning, we compared the normalized spectral counts in aos mutants to those in wild type leaves after wounding (Gfeller et al., 2011). The identification of proteins that were changed in abundance required statistical tests that are best performed outside of the database in a program such as R (R Development Core Team, 2010) using lists of normalized spectral counts for each biological replicate of each sample that were created as an output of the pep2pro database. The same approach was used to investigate the proteome of Tic159 import receptor mutants, which revealed putative Tic159-dependent and Tic159-independent plastid precursor proteins (Bischof et al., 2011).

PROTEOME DATA-SPECIFIC CHALLENGES SOLVED IN pep2pro

The types of analyses described above required proteome data integration in a relational database. Considering the typical state of proteomics data illustrated in Figure 1, this posed a challenge that we solved in the pep2pro database (Baerenfaller et al., 2011). For example, one challenge was the integration of search results from different search algorithms. Results of the different search algorithms have different formats and quality scores and they can contradict each other. Such contradictions are possible when different search algorithms annotate the same spectrum with different peptide sequences, different post-translational modifications, or with post-translational modifications at different
While the data in pep2pro are public, until now the pep2pro information was not usable and detailed data analysis by querying pep2pro for different issues related to proteome data. The relational database schema integrated in the pep2pro data analysis pipeline addresses most issues related to proteome data (Baerenfaller et al., 2011). Then, it can be used for subsequent analyses and biological interpretations and the exclusion of the decoy hits from the final datasets that can cause high false discovery rates in the datasets after data upload and filtering so that they can be distinguished from the forward hits. Integration of the decoy database it was first checked, whether their sequences in the T AIR10 database (Lamesch et al., 2012) 15.2% of all theoretical trypptic peptide sequences and 5.1% of the theoretical trypptic peptides with a minimum length of six amino acids and a molecular mass between 400 and 6000 Da are contained in more than one protein, while 138 peptide sequences are assigned to several loci. We therefore developed a peptide ambiguity filter, which only integrates peptide sequences into pep2pro that are unambiguously assigned to one protein only. This does not apply to different splice variants of the same locus that most of the times cannot be distinguished by the identified peptides anyway or to different genes with exactly the same amino acid protein sequence. Another important aspect during construction of pep2pro was the integration of the decoy database hits. For this we first needed to create a look-up table containing those peptide sequences that both occur in the forward and the decoy database (553 peptide sequences in the TAIR10 database). Upon import of the decoy hits into the pep2pro database it was first checked, whether their sequences were contained in the look-up table, and if not, they were flagged so that they can be distinguished from the forward hits. Integration and flagging of decoy database hits allows the assessment of local false discovery rates in the datasets after data upload and filtering and the exclusion of the decoy hits from the final datasets that can then be used for subsequent analyses and biological interpretations (Baerenfaller et al., 2011).

Together, the upload and filtering algorithms implemented and integrated in the pep2pro data analysis pipeline address most issues related to proteome data. The relational database schema in which the proteome data reside allows users to perform flexible and detailed data analysis by querying pep2pro for different information.

**pep2pro4all: MAKING THE pep2pro DATABASE SYSTEM BROADLY AVAILABLE.**

While the data in pep2pro are public, until now the pep2pro database system was available only to users who provided their mass spectrometry raw files, which we then analyzed using our data analysis pipeline. Alternatively, users could re-build the pep2pro analysis pipeline in their own environment based on the published database schema and the detailed explanations of the different building blocks (Baerenfaller et al., 2011). To make the analysis pipeline more broadly available to users we now built the pep2pro4all system.

**THE pep2pro4all DATABASE SCHEMA.**

The pep2pro4all database schema enables users to locally build a light version of the pep2pro data analysis pipeline and integrate their mass spectrometry search results in mzdentML format into the pep2pro4all database schema (Figure 2). For this, users can download all necessary files and data from www.pep2pro4all.ethz.ch, install the empty pep2pro4all database schema and use the p2p4a script for populating the database. To deal with unambiguous peptides and forward peptide sequences in the decoy database, information on loci, splice variants, and protein sequences from the TAIR10 Arabidopsis search database is provided and used by the p2p4a script for data processing. This assures that only unambiguous peptides are uploaded for analysis and that only peptide sequences that exclusively occur in the decoy database are flagged as decoy hits. Further, upon population of the pep2pro4all database, the sequence-specific protein sequences and protein and locus information are stored in the corresponding database tables. In the pep2pro4all version provided here the sequence-specific information is based on the TAIR10 release of the Arabidopsis protein search database. However, this can be extended to other search databases, plant species, and organisms upon request.

**THE mzdentML DATA STANDARD.**

The second version (v1.1) of the mzdentML file format was released recently and represents the stable and recommended exchange format for peptide and protein identification data (Jones et al., 2012). It was developed by the Proteomics Standards Initiative (PSI) in collaboration with instrument and software vendors, and the developers of major open-source proteomics projects. Software implementations have been developed to enable conversion into mzdentML format from various search algorithm output formats and the major public proteomics data repositories will soon support the format. Additionally, the various methods used by mzdentML to reference external spectra are supported by a common interface implemented by parsers of the jmzReader library that parse the most commonly used mass spectrometry data formats (Griss et al., 2012). Therefore we decided to develop the p2p4a script based on the mzdentML standard file format to facilitate the exchange of MS proteomics data irrespective of search engine and analysis software.

FROM SEARCH RESULTS TO AN INTEGRATED DATASET IN RELATIONAL FORMAT.**

The p2p4a script using the mzdentML format was developed based on the examples provided with the release of mzdentML v1.1 (Jones et al., 2012) and on a mzdentML file converted with the script.
Mass spectrometry files are first converted to standard formats.  
(1) They are then searched with different peptide spectrum assignment algorithms, which create search results files in different formats.  
(2) For integration of the mass spectrometry result files into pep2pro, a separate parser reads out the different formats and applies different import settings. Before integration into the relational database, the data are filtered with a peptide ambiguity filter and inconsistencies in the results are flagged.  
(3) After data integration, the data are ready for data analysis, are displayed on the pep2pro website and can be exported to the PRIDE database.  
(4) For import of mass spectrometry result files into pep2pro4all, they are first converted into mzIdentML standard format using a mzIdentML converter.  
(5) The p2p4a script then parses the data, performs a peptide ambiguity check using the provided sequence-specific information from the search protein database, assembles the dataset, and finally writes the data and all necessary SQL commands into a .sql file. For creating the pep2pro4all database populated with mass spectrometry result data, first the empty pep2pro4all database schema is imported into a local MySQL server, and afterward the .sql output file containing all the data is imported into the database.  
(6) The mass spectrometry result files now reside in relational format in the pep2pro4all database and can be analyzed by data querying.

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