

Building high-quality assay libraries for targeted analysis of SWATH MS data

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Targeted proteomics by selected/multiple reaction monitoring (S/MRM) or, on a larger scale, by SWATH (sequential window acquisition of all theoretical spectra) MS (mass spectrometry) typically relies on spectral reference libraries for peptide identification. Quality and coverage of these libraries are therefore of crucial importance for the performance of the methods. Here we present a detailed protocol that has been successfully used to build high-quality, extensive reference libraries supporting targeted proteomics by SWATH MS. We describe each step of the process, including data acquisition by discovery proteomics, assertion of peptide-spectrum matches (PSMs), generation of consensus spectra and compilation of MS coordinates that uniquely define each targeted peptide. Crucial steps such as false discovery rate (FDR) control, retention time normalization and handling of post-translationally modified peptides are detailed. Finally, we show how to use the library to extract SWATH data with the open-source software Skyline. The protocol takes 2–3 d to complete, depending on the extent of the library and the computational resources available.

INTRODUCTION

Most proteomic analyses involve one or several of an array of MS methods. To date, the most frequently used method is data-dependent acquisition (DDA), because of its unmatched capabilities to identify the protein components of a sample (see **Box 1** for brief explanations on relevant proteomics terminology). DDA-based methods, also referred to as shotgun proteomics, have been widely used to identify and quantify proteins. However, in highly complex proteomic samples, the semi-stochastic nature of DDA leads to some curtailments in the consistency of quantification across many samples, particularly for lower abundant peptide species¹. More recently, mainly driven by the demands of translational research and systems biology projects, the need to generate data that allow the comparative relative or absolute quantification of sets of proteins reproducibly and accurately across sample cohorts numbering tens to hundreds of samples has been recognized. At present, the method of choice for such studies is targeted MS in which definitive assays are used to quantify predetermined sets of proteins across samples at a high degree of reproducibility. The gold-standard targeting MS technique is SRM, also referred to as MRM². In 2012, we introduced SWATH MS³ as a next-generation targeting method that largely maintains the favorable performance characteristics of S/MRM such as quantitative accuracy, dynamic range and reproducibility, while substantially extending the number of quantifiable peptides from the range of tens to hundreds (with scheduling) per sample injection for S/MRM to thousands or tens of thousands per sample injection for SWATH MS. Thus, SWATH MS supports the accurate relative quantification of large fractions of a proteome in a single injection³.

SWATH MS is a variant of the class of data-independent acquisition (DIA) methods that record fragment ion spectra of all ionized species of a sample^{4,5}. For SWATH MS data acquisition,

a high-resolution quadrupole time-of-flight MS cycles through a series of fixed precursor isolation windows that collectively cover the entire *m/z* range of MS-suitable peptides and acquires composite fragment ion spectra from all the precursor ions contained in a specific window at a given time. The window size and dwell time are chosen such that the cycle time is short enough to allow each peptide to be fragmented ~8–10 times across its chromatographic elution profile. A SWATH MS data set therefore constitutes a complete digital record of all ionized species above the detection limit where the fragment ion spectra of individual peptides are represented in a convoluted, but highly structured, manner. The quality of these digital maps mainly depends on the precursor isolation window width, fragment ion resolution, dwell time and cycle time. To identify and quantify peptides in such SWATH MS fragment ion maps, we have devised a targeted data analysis strategy³ that is supported by software tools such as OpenSWATH⁶, PeakView (AB Sciex), Spectronaut⁷ or Skyline⁸, and it is modeled after the automated identification of peptides by S/MRM⁹. In essence, these tools identify peak groups that uniquely associate with the targeted peptide within the comprehensive SWATH MS signal map, and then compute a probability that the targeted peptide has been correctly identified. The peak groups consist of the signals of specific fragment ions derived from the target peptide (transitions) integrated over chromatographic time. The set of transition signals that identifies a target peptide with the highest sensitivity and specificity constitutes a definitive assay for the detection of that peptide, and it has to be determined before the analysis.

A high-quality library of assays is a prerequisite for SWATH MS and similar targeting MS methods¹⁰. Such an assay library is typically built from compendia of fragment ion spectra (spectral libraries), and it contains the exact MS coordinates for each



Box 1 | Terminology

Centroid/profile mode	Raw data (profile mode) is peak-picked (centroided) to produce a peak list of precursor ion masses (MS1) and of fragment ion masses (MS2), which can be used in peptide identification by database searching.
Data-dependent acquisition (DDA)	A mode of operation of a tandem mass spectrometer in which a fixed number of the most abundant precursor ions (e.g., top 20) in every MS1 survey scan are selected for fragmentation and subsequent recording of an MS2 scan. This strategy is commonly referred to as ‘shotgun proteomics’.
Data-independent acquisition (DIA)	A mode of operation of a tandem mass spectrometer that uses a fixed duty cycle to acquire MS2 spectra from mixed populations of precursor ions, which have been cofragmented using isolation windows ranging from tens to hundreds of m/z units. Comprehensive MS2 spectral coverage over a large mass range can be achieved by iterating over sequential precursor isolation windows in a single duty cycle.
Decoy	Additional peptide or protein sequences concatenated to the main protein sequence database, which are used to estimate the FDR in database searching. The decoys should be representative of the target proteins in number and composition. They are typically generated by pseudo-reversal, reversal or scrambling of the target protein sequences, and they should not be identical to target proteins or peptides.
False discovery rate (FDR)	An estimate of the number of false positive identifications contained in a database search result at a given score threshold. FDR can be estimated at the PSM, peptide and protein levels.
Peptide-spectrum match (PSM)	A confident assignment by a database search engine of a peptide sequence to a single MS2 spectrum acquired in DDA mode.
Selected/multiple reaction monitoring (S/MRM)	A mode of operation of a triple quadrupole mass spectrometer in which the first quadrupole is fixed on the precursor m/z of a given peptide, the precursor is fragmented in the collision cell and the third quadrupole is fixed on a fragment ion. The instrument cycles through a fixed list of Q1/Q3 pairs (transitions; see below), and intensities are recorded over chromatographic time. It is considered the gold standard for peptide quantification.
Transition	A pair of masses that represent the precursor ion and a single fragment ion from that same precursor. Multiple transitions are measured in an S/MRM experiment to unambiguously identify and quantify a peptide.
SWATH MS	An instance of the DIA strategy in which highly multiplexed MS2 spectra are collected from wide precursor windows that are designed to cover the m/z range expected for tryptic peptides in a cycle time that is short compared with the elution time of a peptide (e.g., 32 windows of 25 m/z width acquired at a dwell time of 100 ms per window). Quantitative data are extracted in a targeted manner on the basis of prior knowledge of mass spectrometric and chromatographic behavior of peptides using an assay library.
Indexed retention time (iRT)	A normalized retention time space typically calibrated using synthetic peptides that are spiked into every sample measured.
Spectral library	A collection of MS2 spectra with high-confidence peptide sequence assignments.
Consensus spectral library	A spectral library in which MS2 spectrum entries with a redundant peptide sequence assignment have been collapsed into a single entry.
Assay library	A set of coordinates used for targeted extraction of SWATH/DIA data, which typically includes the peptide sequence, the precursor m/z and charge state, the most intense fragment ions m/z and charge states, relative fragment ion intensities and iRT.

targeted peptide. For each peptide, these coordinates consist of (i) the peptide precursor m/z , (ii) the m/z for a selection of its fragment ions together with their relative intensities and (iii) the chromatographic retention time of the peptide in a normalized retention time space. Ideally, the peptides in the assay library cover all proteins of interest for a particular study, or even an entire proteome.

Over the past years, we and others have developed software tools for the generation of spectral libraries (SpectraST¹¹, X!Hunter¹² and Bibliospec¹³). They were originally devised for searching DDA data sets by spectral matching. Analogous spectral libraries have also been used for targeted proteomics by S/MRM^{8,14–16} or SWATH MS^{3,17}, ideally built from fragment

ion spectra generated on the same type of instrument used for targeting. To eliminate the need for assay generation for each experiment, our group has spearheaded the development of publicly accessible assay libraries for the entire proteome of *Saccharomyces cerevisiae*¹⁸, *Mycobacterium tuberculosis*¹⁹ and *Streptococcus pyogenes*²⁰, as well as for disease-relevant human subproteomes, including the human glycoproteome²¹ and a set of cancer-associated proteins²². Most of these assay libraries were optimized for S/MRM, and they are available through the SRMAtlas database (<http://www.SRMAtlas.org>). More recently, we developed an assay library optimized for SWATH MS, which covers >10,000 human proteins annotated in the UniProtKB/SwissProt database²³.

PROTOCOL

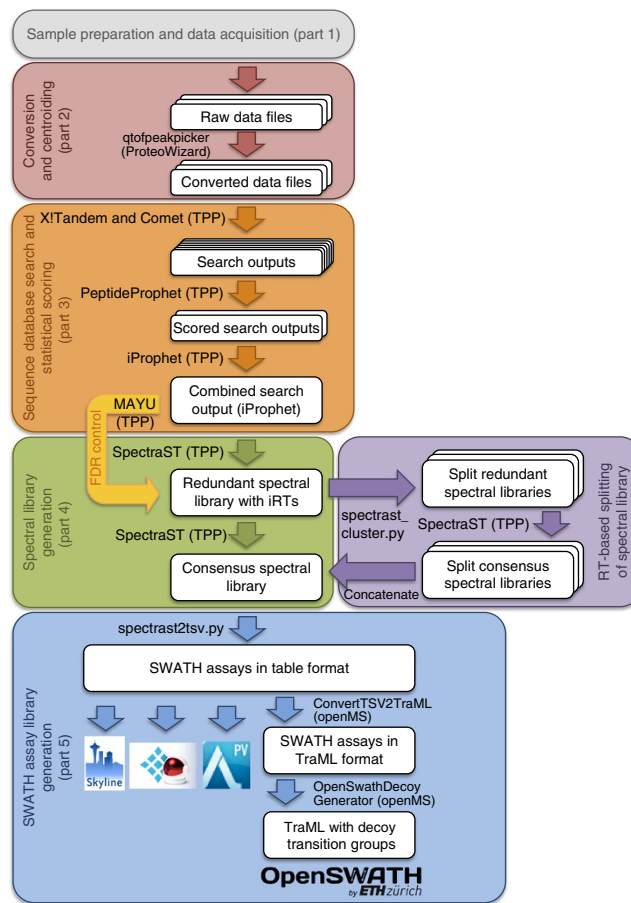
Figure 1 | Workflow for SWATH assay library generation. The library building workflow starts with the selection of representative samples and fragment ion spectra acquisition (part 1), followed by centroiding and conversion of the raw files into an open format (part 2). The centroided fragment ion spectra are searched against a protein sequence database to establish PSMs (part 3). The confidently assigned spectra are then converted into a spectral library, and all retention times are normalized and converted into iRTs (part 4). To address complications when building libraries for post-translationally modified peptides, an optional subroutine has been developed to account for potential errors in site localization of modifications. After consensus library generation, the most intense fragment ions of each peptide precursor are selected (part 5). Optionally, the resulting assay library in table format can be converted into TraML format, and decoy transition groups can be added if required for downstream analysis. RT, retention time.

In this paper we describe a step-by-step protocol and an integrated, openly accessible computational pipeline to generate high-quality assay libraries for targeted MS. All required software tools are freely available through the TPP²⁴, ProteoWizard²⁵ and OpenMS²⁶ software suites, or they are provided as a Python package together with this protocol. For the purpose of user-friendliness, we implemented the protocol on a Windows platform. The computational pipeline described here allows maximal control over each step of the library building process, and it is suitable for large, organism-wide assay libraries, as well as for experiment-specific assay libraries generated from as little as a single DDA data set. The conceptual workflow and considerations to be made at each step are, however, generic, and other tools such as the integrated ProteinPilot-PeakView pipeline, Spectronaut (supporting MaxQuant²⁷ search engine outputs) or Skyline (supporting various search engine outputs) might be used instead. These integrated pipelines allow less control over the workflow, but nevertheless they might provide a suitable alternative for researchers who wish to avoid data handling by command-line tools and Python scripts.

The assay library building workflow described here is optimized for SWATH MS. However, in combination with dedicated analysis tools, it is also applicable to other targeted MS techniques, including S/MRM. Moreover, many of the considerations described here are also valid for building libraries in the context of spectral library searching of DDA^{28,29} or DIA³⁰ data sets.

Protocol overview

The purpose of this protocol is to build assay libraries that support targeted analysis of SWATH MS proteomic data sets. The protocol covers the acquisition of high-quality fragment ion spectra in DDA mode, the assignment of peptide sequences to these spectra, their conversion into spectral libraries and the compilation of the final assays from the spectral libraries. The steps required for custom SWATH assay library generation are outlined in **Figure 1**, and they are described in detail in the following paragraphs. In the PROCEDURE we provide detailed step-by-step instructions, including a detailed list of all files produced during the workflow (**Supplementary Note 1**) and guides to the installation and use of all software tools (**Supplementary Notes 2 and 3**), using a prototypical data set consisting of three samples from a yeast osmotic shock time-course experiment³¹. In **Box 2**, we provide a checklist that summarizes the most important points of the protocol.



Part 1a (Step 1 of the PROCEDURE): selection of representative samples for the generation of spectral libraries

Peptides can only be identified by targeted MS if they are included in the assay library. To cover proteins that are expressed in specific biological conditions, the samples used to generate those assay libraries should therefore represent the entire biological space to be quantified by SWATH MS. In addition, because SWATH MS analysis is more sensitive in identifying and quantifying peptides compared with DDA in side-by-side analyses on the same instrument³, it might be worth fractionating the samples used to generate the assay library before the analysis, e.g., by using isoelectric focusing by off-gel electrophoresis³² or SDS page¹⁰. The resulting fractions are then subjected to DDA, preferably on the same type of instrument that is also used for the subsequent SWATH MS analyses.

As an alternative to the use of fragment ion spectra of native peptides, assay libraries can be built from or supplemented with synthetic peptides¹⁶ or recombinant proteins³³, or they can be computationally predicted. We have previously shown that unpurified unlabeled synthetic peptides produce spectra that are indistinguishable from those derived from natural endogenous peptides¹⁸. By analyzing pools of 100–1,000 synthetic peptides, high-quality fragment ion spectra can be produced very efficiently for large numbers of target proteins, even if they have never been observed from a natural source^{18,19}. A number of approaches and tools have been described to predict those peptides of a protein that are most suitable for targeted analysis (PeptideRank³⁴,

Box 2 | Library generation checklist

This box is meant to summarize critical considerations to be made during SWATH assay library generation.

DDA data acquisition

- The samples might be prefractionated (e.g., by isoelectric focusing using off-gel electrophoresis) to increase coverage.
- The samples contain reference peptides for retention time normalization.
- The DDA instrument parameters are optimized for high-quality fragment ion spectrum acquisition (i.e., longer acquisition/dwell time/trap filling, shorter dynamic exclusion).
- The DDA collision energy, including ramping, mimics the one to be used to fragment that same precursor in SWATH acquisition.
- The spectra in DDA files are centroided with a suitable converter, optimally using fragment ion peak areas instead of peak height for centroiding.
- The DDA files are converted to centroid mode without de-isotoping.

DDA database search and spectral library generation

- When multiple search engines are to be used, the DDA data files are converted to the various input formats with consistent spectrum indices.
- The multiple search engine results are aggregated using adequate tools (e.g., iProphet).
- The protein FDR of the raw spectral library is tightly controlled to be $\leq 1\%$.
- The retention times in the raw spectral library are aligned to reference values (e.g., iRT) before the consensus library generation.
- Optional (mainly recommended for assay libraries with post-translational modifications): Precursors in the raw spectral library are split into as many clusters as needed on the basis of their normalized retention times before consensus library generation.

Assay library/transition list generation

- Fragments smaller than 350 m/z or bigger than 2,000 m/z are filtered out.
- Fragments with m/z in the precursor SWATH window are filtered out.
- Only fragments with mass accuracy within $\pm 0.05 m/z$ of the expected mass are used.
- The most intense y - and b -ion fragments fulfilling the above criteria are selected.
- In the case of a library containing assays for C-terminally heavy isotope-labeled peptides, no b -ions must be included.
- Fragments with neutral loss may be considered if the library was acquired on the same instrument.
- All assays should have the same number of fragment ions.

PeptideSieve³⁵, CONSeQuence³⁶, ESPPredictor³⁷, Detectability Predictor³⁸ and STEPP³⁹), although their predictors are less accurate than empirical assessment of optimal peptides³³. Fragment ion spectra can also be entirely computationally predicted using physicochemical models of peptide fragmentation or by machine-learning approaches. Predicted fragment ion spectra, however, are also expected to less faithfully match empirical spectra of native peptides⁴⁰. For library building, empirical data, either from native or synthetic peptides, are therefore preferable over computationally predicted spectra.

Generally, it has been shown that very extensive SWATH assay libraries, for example, those resulting from fractionation before MS analysis and/or addition of synthetic peptides, lead to more peptide and protein identifications during SWATH data analysis without impairing quantification accuracy²³. It is, however, important to note that such extensive assay libraries require more stringent FDR control during the SWATH data analysis, because the increased search space results in higher numbers of false-positive identifications (see also **Box 3** and ref. 23). Incidentally, the presence of an assay in the library does not guarantee that the corresponding peptide can be detected—i.e., if the peptide is not present or is below the limit of detection in the SWATH analysis of a given sample.

Part 1b (Step 2): working with retention time reference peptides

Because the chromatographic retention time of the targeted peptides is an essential component of the final peptide assay, it is recommended to work with retention time reference peptides⁴¹,

which are spiked into all samples that are used for library generation. This will allow effective peptide retention time normalization and retention time–based splitting of fragment ion spectra, to determine most accurate retention times for each targeted peptide. Alternatively, endogenous retention time reference peptides can be used for retention time normalization (ref. 10 and Parker *et al.*, unpublished data). As with spike-in reference peptides, endogenous reference peptides need to cover a large retention time range of the sample, and they need to be well detectable over all samples. All downstream steps are identical and the quality of the alignment, as well as the resulting data, is very comparable between the workflow with spike-in and endogenous retention time reference peptides. As a reference scale for the retention time normalization, either the unit-less iRT scale⁴¹ or the retention times in minutes from any previously acquired MS injection can be chosen. The main advantage of using the iRT scale is that it is a defined reference, and therefore it facilitates transferability between instruments and laboratories.

Part 1c (Step 3): acquisition of fragment ion spectra

Optimal performance of the assay library for the targeted identification of peptides in SWATH MS data sets is achieved if the spectra used to generate the spectral libraries reflect as closely as possible the relative fragment ion intensities in the SWATH MS maps. To ensure optimal portability of the assays, it is therefore highly recommended to use the same type of instrument for library generation as for SWATH MS analysis⁴². If no TripleTOF 5600+ mass spectrometer is available to generate the library, an alternative instrument with beam-type collision cell or ion trap–type collision

Box 3 | Considerations for controlling the spectral library FDR

Estimating and controlling the FDR in shotgun proteomics has been the subject of many studies, and standard methods using model-based^{53,60} and decoy-based⁴⁶ approaches, or hybrids of these, are now well established in the field. As the scale of proteomics projects has grown, and the scanning speed of mass spectrometers has increased, it has become apparent that methods that deal specifically with robustly estimating FDRs in very large-scale data sets are required. Such methods have been developed and implemented in the MAYU software⁵⁴ with a particular focus on estimating the PSM level, peptide level and protein level FDR in large-scale DDA data sets. As the creation of very large spectral libraries for use in targeted SWATH data analysis workflows is actively being pursued, a discussion of the effect of error rates in spectral libraries built for targeted analysis of SWATH/DIA is justified.

Previous studies have emphasized the importance of high-quality spectra when constructing spectral libraries and suggested that errors introduced at this stage might be propagated into the results of a spectral library searching strategy for DDA data¹¹. A question that has not been directly addressed is whether errors in spectral libraries will be propagated into targeted analysis of SWATH data, or whether the consistent fragment ion spectrum sampling in chromatographic time will be able to resolve errors that are introduced at the spectral library level. This question can only be answered by considering the source of error in the spectral library. For example, if the error arises because of coisolation of multiple peptide species (or other species), then a mixed (or chimeric) spectrum will result with the potential to match to a peptide in the sequence database with a high score. If such a library spectrum is then used as the basis for targeted analysis of SWATH data, it is improbable that a high score will be produced because the fragment ions are very unlikely to perfectly co-elute and, as such, the error from the spectral library will not be propagated to the SWATH data analysis results.

However, there is a second type of spectral library error that is more problematic. That is, a fragment ion spectrum in the DDA data could be produced from a single peptide precursor and still match to the wrong sequence in the database search. If this is the case, SWATH data, and targeted analysis thereof, will match faithfully to the library spectrum with perfect co-elution of fragment ions, thereby propagating the original error into the SWATH analysis results. If the first type of error is predominant in the DDA data, then a moderate FDR in the spectral library would be well tolerated for downstream analysis and propagation to SWATH results would not be an issue. However, if the second type of error is more frequent, a more conservative FDR threshold in the library creation would be required. To our knowledge, a systematic investigation into which type of error predominates in DDA data has not been performed, but it remains an open question worthy of further study. With this uncertainty, researchers may choose the threshold depending on the downstream analysis question, but perhaps for large-scale libraries that could be distributed for use by many labs a more conservative threshold is warranted. In any case, robust methods for estimating FDR at PSM, peptide and protein levels, such as provided by MAYU, should be used during the library creation process.

cell operable in higher-energy collisional dissociation (HCD) mode may be used instead, as they generate fragmentation patterns that are similar to those generated by the TripleTOF 5600 instrument⁴³. If data sources other than those from a TripleTOF 5600 instrument are used, it is recommended to ensure that the relative fragment ion intensities do not exceed 30% variation between DDA and SWATH MS measurements, as a larger difference would impede the use of relative fragment ion intensities as a peptide identification score during SWATH MS data analysis⁴⁴.

The optimization of the MS acquisition settings to generate fragment ion spectra for library generation will be described in detail elsewhere (S. Bader, A. Keller, D. Shteynberg, L. Hood and R. Moritz, unpublished data; cited with permission from the authors). Here we suggest the following generic settings for the acquisition on a TripleTOF 5600: (i) increase the fragment ion spectrum accumulation times to 150 ms to maximize the quality of the spectrum. (ii) Record more than one fragment ion spectrum of the same precursor by reducing the dynamic exclusion time to 20 s, which is approximately half of a chromatographic peak width (typically 30–60 s). This will increase the chance that a second fragment ion spectrum is recorded from the same sample at higher peptide precursor signal intensity than the first fragment ion spectrum. Further, if the first spectrum is contaminated with fragments of a second, concurrently fragmented precursor, the background would be expected to be changed for the second spectrum. (iii) Aim for the highest similarity possible between the relative intensities of the fragment ions in the library and in the SWATH MS measurements by using the same collision energy settings for

both modes of operation. Specifically, regardless of the charge state of the selected peptide precursor, we recommend using a collision energy that reflects most closely the settings used in SWATH MS data acquisition, for example, applying a collision energy according to the equation of a doubly charged peptide (slope 0.0625, intercept -3.5), ramped ± 15 V from the calculated collision energy over the MS2 accumulation time (with an upper limit of 80 V). The specific instrument acquisition settings recommended in this protocol are summarized in **Supplementary Table 1**.

Part 2: conversion and centroiding of fragment ion spectra

Database search engines that establish PSMs generally require raw instrument output data (profile spectra) to be converted into a peak list format (centroided spectra) in a vendor-independent open format such as mzML⁴⁴ or mzXML⁴⁵. Both the conversion and centroiding (peak picking) process are typically performed by a single tool, the converter. The available centroiding algorithms slightly differ in the way they extract intensities from profile peaks in precursor (MS1) and fragment ion (MS2) spectra. For the purpose of assay library generation, it is important that the converter yield fragment ion intensities that match, as close as possible, those extracted from SWATH MS data. For conversion and centroiding of TripleTOF 5600 fragment ion spectra, we tested three different converters: ProteinPilot (AB Sciex), msconvert (with 'prefer vendor' setting; developed by AB Sciex) and qtofpeakpicker, which has been developed by our group and is, like msconvert, also distributed through ProteoWizard (see **Supplementary Note 2** for algorithm details). A comparison of



TABLE 1 | Effect of increasing data set size on score threshold and FDR.

iProphet probability threshold	PSM FDR (%)	Peptide FDR (%)	Protein FDR (%)	Protein identifications	Data set
0.9774	0.08	0.20	1.0	2,162	Yeast 3 DDA files (this protocol)
0.9171	0.16	0.40	2.0	2,249	
0.3983	0.37	1.04	5.0	2,414	
0.9994	0.07	0.18	1.0	11,102	Human 331 DDA files (Rosenberger <i>et al.</i> ²³)
0.9970	0.19	0.38	2.0	11,537	
0.9809	0.44	1.00	5.0	12,203	

the results obtained if sets of DDA files were converted by either of these converters indicates that the qtofpeakpicker (using peak areas) yielded the highest level of reproducibility of fragment ion spectra across replicate DDA runs (**Supplementary Fig. 1a**) and achieved best portability of the derived assays to the corresponding fragment ion intensities obtained by SWATH MS (**Supplementary Fig. 1b**). The numbers of identified peptides and proteins from a database search after conversion with any of the peak pickers are slightly higher for the qtofpeakpicker than those achieved for the other two peak pickers tested (**Supplementary Fig. 1c**). Examples for the converter-dependent variability of relative abundances of fragment ions in centroided MS2 spectra are given in **Supplementary Note 4**. In summary, these data show that different centroiding algorithms can cause surprisingly large intensity differences for even the most prominent peaks of a fragment ion spectrum. To ensure the highest-possible assay quality when accurate relative fragment ion intensities might be crucial for downstream use, a tool that maintains these relative fragment ion patterns, such as the qtofpeakpicker described above, should be selected for the conversion and centroiding of raw instrument files into a search engine-compatible and vendor-independent open format.

Part 3: sequence database searching and statistical scoring of PSMs

Spectral libraries are built from fragment ion spectra that are assigned with high confidence to a peptide sequence. To establish this match, centroided fragment ion spectra are subjected to sequence database searching. At this stage, it is important that the protein sequence database (typically in FASTA format) contain the sequences of the retention time reference peptides to allow for retention time normalization at a later step. To control the FDR of the PSMs, the protein sequence database also needs to contain a decoy entry for every protein⁴⁶. Even though protein sequence reversal is, owing to its simplicity, the most commonly used method to generate decoy peptides, decoys most precisely reflecting target peptides are generated by pseudo-reversal of target peptide sequences⁴⁶. This latter method was thus used for

this protocol. To maximize the number of PSMs and the discrimination between true and false assignments, the search output of multiple search engines may be combined⁴⁷. In general, we recommend using search engines that are maximally orthogonal in their search algorithms, as this results in highest numbers of identifications⁴⁸. The optimal parameters for search engines, such as the number of tolerated tryptic termini, missed cleavages, precursor mass tolerance and variable modifications, depend on the specific biological sample, experimental setup and purpose of the library. In this protocol, searches were done for fully tryptic peptides. Although semi-tryptic peptides might originate from biologically relevant proteolytic cleavage by endogeneous proteases, several publications have reported that most of those peptides may originate

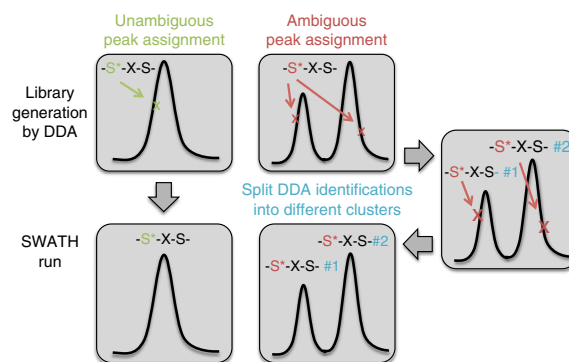
from nonspecific trypsin/chymotrypsin activity^{49,50} and/or in-source fragmentation⁵¹. A discussion on how various parameter settings affect the underlying search has been provided by Eng *et al.*⁵². Notably, the targeted SWATH data analysis does not change upon inclusion or exclusion of semi-tryptic peptides, as these will be extracted in the same manner as fully tryptic peptides. **Supplementary Table 2** contains the main parameters used for the protocol case study. The PSMs from each search engine are scored using PeptideProphet⁵³ and subsequently combined and rescored using iProphet, a tool that integrates evidence from multiple identifications of the same peptide across different experiments and search engines and thus improves the discriminating power between correctly and incorrectly assigned PSMs⁴⁷.

As dozens or hundreds of DDA runs might be combined to generate a comprehensive SWATH assay library, it is important to thoroughly control the FDR of the final data set both at the level of PSMs and at the level of inferred proteins. The MAYU software⁵⁴ has implemented a robust method to estimate the FDR of such large-scale DDA data sets at PSM, peptide and protein levels, and it can be applied to the iProphet output. **Table 1** shows the FDR at PSM, peptide and protein levels for the data set associated with this protocol generated from three DDA files in yeast in comparison with a recently published large-scale human library²³ generated from 331 DDA files, highlighting the requirement for increased stringency in larger data sets. In some cases, the iProphet probability threshold to achieve a certain protein FDR appears to be low; however, this is unproblematic as the iProphet probability is intended to be interpreted at the peptide level and not at the PSM level. The iProphet probability is used as a ranking, and the FDR estimated by MAYU is controlled on the basis of decoys. How the MAYU-estimated FDRs change with the applied iProphet score cutoff is shown in **Supplementary Figure 2** for the data set described in this protocol. A discussion of the effect of error rates in spectral libraries for targeted analysis of SWATH/DIA, in particular with respect to error propagation from assay library to SWATH identifications, is provided in **Box 3**.



PROTOCOL

Figure 2 | Splitting peptide identifications with distant elution times. During a DDA search, it may happen that multiple fragment ion spectra are assigned to the same peptide precursor, even though they span a wide retention time segment and might not come from the exact same molecular species. This is not a rare event, especially in the context of post-translationally modified peptides in which the modification cannot be unambiguously assigned to a certain amino acid. This figure depicts such ambiguous peak assignment on the example of a phospho-peptide containing a phosphorylated serine (S*) in presence of a second, unphosphorylated serine (S). Fragment ion spectra recorded at distant retention times can be clustered apart during the SWATH assay library generation. The distinct SWATH assays might then be used to resolve the correct assignment on the level of SWATH MS data. See **Supplementary Note 5** for examples.



Part 4: generation of a spectral library with aligned retention times

SpectraST is a software tool that compiles all fragment ion spectra assigned to a specific peptide sequence above a certain quality threshold (e.g., iProphet probability) into a spectral library format^{11,28}. At this step, it is advisable to transform all retention times into a normalized retention time scale⁴¹ before the consensus spectra are computed. This is accomplished by establishing a linear correlation between experimental retention times and unit-less absolute retention time values (iRTs) for retention time reference peptides identified in each DDA run. The resulting correlation curves are then used to convert the retention time for all the other peptides identified in the corresponding DDA runs into iRT scale.

Even though modern mass spectrometers display a reasonably high level of reproducibility in repeat recordings of fragment ion spectra of the same peptide across replicates⁴², the consolidation of multiple fragmentation observations of the same peptide precursor ion into a single consensus spectrum provides a more accurate fragmentation pattern than any single spectrum (best replicate) for that precursor^{11,55} (**Supplementary Fig. 3**). The consensus spectrum, therefore, is the optimal representation of the fragment ion spectrum of a targeted peptide.

To avoid combining ambiguously assigned fragment ion spectra (e.g., spectra matching to the same sequence but acquired at significantly different normalized retention times into a wrongly averaged consensus spectrum), we developed a strategy to split and process fragment ion spectra from precursors of different retention times for consensus library generation (**Fig. 2** and **Box 4**). These considerations are particularly important in the context of isobaric peptides with post-translational modifications that may be assigned by the search engine to incorrect amino acid residues owing to ambiguities in the site localization. However, we also observe the phenomenon of distant retention times for identical identifications in data sets that had not been searched for post-translational modifications (**Supplementary Note 5**).

Part 5: generation of a SWATH assay library from a consensus spectral library

Once a consensus spectral library has been generated (or downloaded from the web), the most intense fragment ions need to be retrieved for each precursor. The number of fragment ions should be high enough to ensure specificity of identification within a SWATH MS map, but not too high, as less-intense transitions introduce noise into the extracted data, reduce specificity and may adversely affect the target identification and limit of detection. In previous studies, it has been recommended to

use six transitions per peptide precursor⁹. Optionally, simulations can be used to estimate the appropriate number of transitions required to achieve unique ion signatures for the targeted peptides in a given proteome background⁵⁶. Peptide precursors represented in the library with fewer fragment ions than required to achieve high specificity should not be considered. We recommend using the same number of transitions for all assays because different numbers of transitions per precursor may result in mixed statistical distributions for the target identifications in automated peak scoring if this is not accounted for. With regard to ion types, we found that it is acceptable to include γ - and b -ions, as well as common neutral losses, if the library was recorded on the same instrument used to record the SWATH MS data⁴².

Transitions with fragment ion masses below 350 m/z should be excluded from the library, as they are typically less specific and thus more noisy than transitions of fragment ions with higher m/z values. Furthermore, also transitions with fragment ion mass falling within the isolation window of their precursor m/z ratios should be excluded from the assay library, as those are typically highly interfered with incompletely fragmented precursors from the same SWATH window. Incidentally, this is the only difference in the process of building a library for S/MRM acquisition and for SWATH MS extraction. As this filtering step makes the assay library dependent on the specific instrument setup, it is therefore desirable to publish not only the final assay library as a transition list but also the consensus spectral library.

In case a library is to be constructed that contains assays for isotopically light and heavy peptides (e.g., labeled with heavy arginine or lysine at the C terminus of each peptide), it is important to consider that, depending on the labeling strategy, b -ion transitions from light and heavy precursors might not be distinguishable. This is because these fragments might not carry the isotopic mass difference, and the chance is high that the precursor m/z of the light and the heavy form of a peptide fall into the same SWATH window. For libraries containing C-terminally isotope-labeled heavy peptides, we thus recommend to only include γ -ions in the SWATH assay library.

Different SWATH data analysis software tools accept different formats of assay libraries. The library formats can be divided into two classes: (i) a simple table in tab-separated (.tsv) or comma-separated (.csv) format in which each row contains a transition and columns contain information to specify this transition, and (ii) a transition list in TraML format⁵⁷. Although the table format is easy to read and manipulate, the TraML format is well

Box 4 | Splitting peptide identifications with distant elution times

On an HPLC system, any given peptide is expected to elute within a single peak at a characteristic, well-defined retention time. However, in a DDA workflow it may happen that multiple identifications of a given peptide actually span a rather large time segment, eventually longer than that covering the average peptide chromatographic peak width within the HPLC condition used. In such cases, questions may arise for consensus library generation whether those multiple fragment ion spectra should be globally combined into a single assay or whether they should be clustered apart and processed independently. Despite having the same peptide identification, those fragment ion spectra could indeed originate from different isobaric peptide sequences (falsely assigned) or different peptide conformations, and therefore a global consensus merging would erroneously combine those different fragmentation spectra, yielding a single peptide assay with both incorrect relative intensities and incorrect retention time approximation (**Fig. 2**). For a best-replicate-based nonredundant spectral library, intensity and retention time would be correctly retrieved for one spectrum cluster, but all information on the other spectrum cluster(s) would be lost.

This phenomenon is quite frequently observed with peptides carrying post-translational modifications, in which it can be challenging for a search engine to accurately locate the modification on the peptide sequence (L.C.G. C. Ludwig, A. Maiolica, U. Toprak, P.N. and R.A., unpublished data). In this library generation protocol, we present an optional extended subroutine to handle such cases. The additional steps consist of (i) the separation of clusters of fragment ion spectra with elution time beyond a user-defined threshold (e.g., two iRT units for our HPLC setup), (ii) the generation of consensus spectra for each of these clusters independently and (iii) the merging of all those consensus spectra into a final consensus spectral library. The resulting assay library thus contains multiple assays (with different relative intensities and retention times) associated with the same peptide sequence. The uniqueness of the peptide assays is provided either by using a unique assay identification number (openswath option in Step 37) or by using an incremented protein name (peakview option in Step 37) for each assay pointing to the same peptide sequence. By using this pipeline, several compelling cases of peptide identifications were found in the provided data sets even without searching for post-translational modifications (**Supplementary Note 5**).

It should be noted that the quantification of those multiple chromatographic peaks matching to identical peptide sequences is not trivial, and it is beyond the scope of this protocol to describe the detailed downstream analysis steps for such cases.

defined, and thus contains unambiguous information and is the format endorsed by the HUPO Protein Standards Initiative. OpenSWATH requires libraries in TraML format and containing pre-computed decoy transition groups, facilitating the discrimination between true and false signals and error rate estimation during SWATH data analysis⁶. Decoys need to represent the targets well, but at the same time they have to be different from the target assays. Decoys based on shuffled sequences have been shown to be best suited for the purpose of modeling the targets⁶. However, decoys based on full reversal of peptide sequences have been successfully used as well, and they enable the generation of decoy transition groups for even highly repetitive or palindromic peptide sequences²³. Other software tools, such as PeakView, Spectronaut and Skyline, require the assay library to be in table format, and they do not require decoy transition groups to be provided with the library. **Supplementary Tables 3–6** summarize the assay library formats required for the currently available SWATH analysis software tools. Both TraML and table formats are supported by the tools described in this protocol, and the hereby generated SWATH assay libraries can thus be directly

used with all major SWATH analysis software suites, including OpenSWATH, Skyline, Spectronaut and PeakView.

For the dissemination of SWATH assay libraries, the SWATHAtlas database (<http://www.SWATHAtlas.org>) provides a suitable platform. To allow more flexibility for the user, as mentioned above, we advise publishing both the redundant and consensus spectral libraries in sptxt format, as well as the final assay list in the csv/tsv format and/or in TraML format.

Several other databases provide assay or spectral libraries for download, such as SRMAtlas (<http://www.SRMAtlas.org>) and PeptideAtlas (<http://www.peptideatlas.org/speclib/>) and NIST (<http://chemdata.nist.gov/dokuwiki/doku.php?id=peptidew:download>). However, the libraries provided through these databases might not have been acquired on a TripleTOF mass spectrometer, and the relative intensities of fragment ions may thus not reflect the relative intensities of the subsequent SWATH MS measurements very well. Furthermore, it is important to understand whether these libraries contain retention time information at all, if the retention times have been normalized or, most optimally, if they contain iRT values.

MATERIALS

REAGENTS

- iRT retention time peptides. The retention time peptides are typically a set of 10–20 synthetic peptides, which span a wide range of hydrophobicities and thus liquid chromatography (LC) retention times, which are converted into the unit-free iRT scale⁴¹. The peptides are spiked into each sample, and they allow normalizing retention times over various MS runs and different LC setups⁴¹. Any set of peptides covering a wide range of hydrophobicity can be used. For this protocol, we used the iRT-kit from Biognosys
- One or several peptide samples (see INTRODUCTION and Step 1 of the PROCEDURE for details.)

EQUIPMENT

- TripleTOF 5600+ mass spectrometer (AB Sciex)
- Nanoflow high-performance liquid chromatography (HPLC) system
- Computer: PC with Microsoft Windows 7 (Microsoft), ≥4 GB of RAM, sufficient hard disk space (for the protocol case study ≥40 GB)

Software

- Microsoft Windows 7 (Microsoft)
- MS Excel (Microsoft)
- ActivePerl (x86)
- OpenMS 1.11—nightly build (32-bit)



PROTOCOL

- TPP 4.7 (polar vortex) revision 1
- Anaconda (32-bit) for Python 2.7 (Continuum Analytics)

EQUIPMENT SETUP

▲ **CRITICAL** Detailed instructions, including screenshots, for the installation of each software module are provided in **Supplementary Note 3**.

ActivePerl (x86) Install ActivePerl from <http://www.activestate.com/activeperl/downloads> with default settings.

TPP 4.7 (polar vortex) rev 1 Install TPP from <http://sourceforge.net/projects/sashimi/files/Trans-Proteomic%20Pipeline%20%28TPP%29/> with default settings. (Note: This will also install ProteoWizard's msconvert and qtofpeakpicker, as well as the Apache http server, which is required for the TPP web interface.) See the TROUBLESHOOTING section for more details.

OpenMS (version 1.11, 32-bit) For the most updated version (nightly build), install from http://ftp.mi.fu-berlin.de/OpenMS/nightly_binaries/. OpenMS will ask to install Microsoft .NET Framework 3.5 and 4.0. See the TROUBLESHOOTING section for more details.

Anaconda (32-bit) Install from <https://store.continuum.io/cshop/anaconda/>, which includes a python interpreter and all required python libraries for

Python 2.7. (Select installation for all users to automatically set the location to C:\Anaconda.)

msproteomictools python package Open the command-line prompt: Open the Start Menu (Windows icon in lower left corner) and type 'cmd' in the search field. Right-click the cmd.exe file and select 'Run as administrator'. To install the package, type: 'C:\Anaconda\Scripts\pip.exe install msproteomictools' Note that alternative ways to install the msproteomictools package are described in **Supplementary Note 3**.

X!Tandem and Comet parameter files (.params), yeast protein sequence file (yeast.fasta), iRT-kit (iRT.txt) and SWATH windows (swaths.txt) Download them from ProteomeXchange⁵⁸ (data set identifier: PXD001126) and move to C:\Inetpub\wwwroot\ISB\data.

DDA raw files: nseivse_L120327_001, 010 and 016 (.wiff, .wiff.mtd .wiff.scan) Download these nine files from ProteomeXchange⁵⁸ (data set identifier: PXD001126) and copy them to C:\Inetpub\wwwroot\ISB\data (the wiff.mtd file can be omitted).

SWATH raw files: nseivse_L120412_001, 010 and 016 (.wiff, .wiff.mtd .wiff.scan) Download these nine files from ProteomeXchange⁵⁸ (data set identifier: PXD001126) and copy them into a folder on your computer.

PROCEDURE

Part 1: sample preparation and data acquisition ● **TIMING** a few hours to several days

▲ **CRITICAL** To exemplify the workflow for SWATH library generation, we provide three TripleTOF 5600+ DDA files from a recent SWATH MS study³¹. The three files represent samples from an osmotic shock time course (0, 60 and 120 min) in yeast. All output files from each step of the procedure are listed in **Supplementary Note 1**, and the parameters of all software tools and commands are described in more detail in **Supplementary Note 2**.

1| Prepare peptide samples with a final concentration of 0.5–1 µg/µl. The method of preparation of these peptide samples from biological specimen is highly dependent on the sample type and has been described in many instances in the literature (e.g., human²³ and yeast⁵⁹); however, essentially any sample preparation method compatible with standard shotgun or targeted proteome analysis should be compatible.

2| Spike iRT peptides into your sample at a ratio of 1:20.

▲ **CRITICAL STEP** The presence of iRT retention time reference peptides is crucial to generate a high-quality SWATH assay library and to perform subsequent SWATH MS data analysis. As iRT reference peptides, either a set of synthetic peptides (as described in this protocol, see also the Reagents section) or, alternatively, a set of well-detectable endogenous peptides spanning a large retention time range can be used.

? **TROUBLESHOOTING**

3| Inject 1–2 µg of your sample onto a nano-HPLC instrument coupled to a TripleTOF 5600+ MS operating in DDA mode (on the TripleTOF 5600 this is called information-dependent acquisition). Please refer to **Supplementary Table 1** for specific instrument parameters.

Part 2: conversion and centroiding of the raw data ● **TIMING** a few hours

4| Once the data are acquired, copy the files into the folder C:\Inetpub\wwwroot\ISB\data for conversion into a vendor-independent format and centroiding.

5| Open the Start Menu (Windows icon in lower left corner) and type 'cmd' in the search field. Right-click the cmd.exe file and select 'Run as administrator'.

6| In the command-line window that opens, type the following command to change to the directory that contains the data:

```
cd C:\Inetpub\wwwroot\ISB\data
```

! **CAUTION** Many commands to be entered in the command line window are case-sensitive, and it is therefore advised to control the spelling carefully.

7| Run the following command on each file to convert and to centroid the profile data:

```
qtofpeakpicker --resolution=20000 --area=1 --threshold=1
--smoothwidth=1.1 --in nselevse_L120327_001.wiff --out
nselevse_L120327_001.mzXML
```

The --area option causes the converter to use the area of a peak as intensity instead of the peak apex. The converted files will be located in C:\Inetpub\wwwroot\ISB\data.

? TROUBLESHOOTING

8| Reduce fragment ion spectrum complexity by keeping only the top 150 peaks:

```
msconvert nselevse_L120327_001.mzXML --mzXML --filter
"threshold count 150 most-intense" --outfile
nselevse_L120327_001_c150.mzXML
```

This filtering leads to a much smaller file size, and, as a consequence, most software tools described in this protocol will run faster. If the library generation workflow is performed using a powerful computing infrastructure, this step can be omitted.

Part 3: database searching and controlling FDR ● TIMING a few hours to several days

9| X!Tandem database search (Steps 9–19) To start an X!Tandem search, navigate to the TPP web interface using a web browser, such as Internet Explorer: http://localhost/tpp-bin/tpp_gui.pl. You should also be able to simply double-click the TPP icon that has been generated on your Desktop during the installation.

10| Enter the user name and password as 'guest' and click on the 'Login' button.

11| In the 'Home' tab, select 'Tandem' as the analysis pipeline from the drop-down menu.

12| Go to the 'Analysis Pipeline' tab and then to the 'Database Search' tab.

13| In the 'Specify mz[X]ML Input Files' section, click the 'Add Files' button and select the three converted and reduced mzXML files.

14| In the 'Specify Tandem Parameter File' section, click the 'Add Files' button and select the 'xtandem.params' file. This file defines the settings to be used for the search. All settings defined here will overwrite the settings in the default parameter file. Please consult **Supplementary Table 2** for a list of parameters that deviate from the default values.

15| In the 'Specify a sequence database' section, click the 'Add Files' button and select the 'yeast.fasta' file. This is a protein sequence database containing all annotated yeast proteins, the iRT retention time reference peptides (concatenated to a single protein) and a pseudo-reversed decoy peptide for each target peptide. The names of the 'proteins' containing the decoy peptides start with 'reverse_' followed by the target protein name. This tag will be used several times again during the course of this protocol (Steps 28, 29, 30, 33 and 35).

▲ CRITICAL STEP The sequence database fasta file needs to contain the iRT retention time reference peptides that can be added as separate 'proteins' or concatenated to a single protein if the peptides have tryptic ends.

16| Check the option 'Convert output to pepXML'.

17| Click the 'Run Tandem Search' button.

18| The view switches to the 'All Jobs' tab where all jobs that have been submitted recently are listed. Click 'refresh' in this table to see the current state. While the X!Tandem search is running, the Comet search can already be submitted as well (starting at Step 20).

? TROUBLESHOOTING

19| When the job has been completed, you can go to the folder C:\Inetpub\wwwroot\ISB\data and check the three files that were generated for each mzXML file: a .tandem.params file, a .tandem file and a .tandem.pep.xml file.

? TROUBLESHOOTING

20| Comet database search (Steps 20–27). In the 'Home' tab of the TPP web interface, select 'Comet' as the analysis pipeline from the drop-down menu.



- 21|** Go to the 'Analysis Pipeline' tab and then to the 'Database Search' tab.
- 22|** In the 'Specify mz[X]ML Input Files' section, click the 'Add Files' button and select the three converted and reduced mzXML files (if not selected already from the X!Tandem search).
- 23|** In the 'Specify Comet Parameter File' section, click the 'Add Files' button and select the 'comet.params' file. This file defines the settings to be used for the search. All settings defined here will overwrite the settings in the default parameter file. Please consult **Supplementary Table 2** for a discussion of the parameters.
- 24|** In the 'Specify a sequence database' section, click the 'Add Files' button and select the 'yeast.fasta' file (if not selected already). See Step 15 for more information on the protein sequence database.
- 25|** Click the 'Run Comet Search' button.

26| The view switches to the 'All Jobs' tab where all jobs that have been submitted recently are listed. Click 'refresh' in this table to see the current state.

? TROUBLESHOOTING

27| When the job has been completed, go to the folder C:\Inetpub\wwwroot\ISB\data and check the files that were generated. For each mzXML file, a pep.xml file should have been generated. Add 'comet' to the file names of these pep.xml files to avoid confusion later on: xxx.comet.pep.xml (where xxx is the file name).

28| Score and combine search outputs with PeptideProphet and iProphet (Steps 28–32). To run PeptideProphet on the X!Tandem search results, run the following command:

```
xinteract -OARPd -dreverse_ -Ninteract.tandem.pep.xml  
nselevse_L120327_0*.tandem.pep.xml
```

Please note that this and the following steps can also be done through the TPP web interface (Steps 28, 29, 30 and 33).

29| To run PeptideProphet on the Comet search results, run the following command:

```
xinteract -OARPd -dreverse_ -Ninteract.comet.pep.xml  
nselevse_L120327_0*.comet.pep.xml
```

30| Run iProphet to combine the search outputs of the X!Tandem and the Comet search and to improve discrimination between true and wrong PSMs:

```
InterProphetParser DECOY=reverse_ interact.comet.pep.xml  
interact.tandem.pep.xml iProphet.pep.xml
```

31| To explore which peptides and proteins have been identified by the search engines or to inspect the corresponding spectra, open the TPP web interface as described in Step 9, click on the 'Utilities' tab and then the 'Browse files' tab and select the 'iProphet.pep.xml' link in the file list to open the iProphet output in the PepXML viewer.

? TROUBLESHOOTING

32| To export a spreadsheet of the iProphet results, click on the 'Other Actions' tab and then on the 'Export Spreadsheet' button. A file named iProphet.pep.xls is created in the folder C:\Inetpub\wwwroot\ISB\data, which can be opened with Excel.

33| FDR estimation with MAYU (Steps 33 and 34). To process the iProphet results with MAYU for FDR estimation, run the following command:

```
Mayu.pl -A iProphet.pep.xml -C yeast.fasta -E reverse_ -G 0.01  
-H 51 -I 2 -P protFDR=0.01:t
```

34| Retrieve the minimum iProphet probability at which the protein FDR is <1% by opening the file ending with '_psm_protFDR0.01_t_1.07.csv' in Excel, sorting the column called 'score' and reading the lowest value. (For the case study it equals 0.9774; depending on the computer you ran the above software tools, the value might be slightly different.)

Part 4: spectral library generation ● **TIMING a few hours to 1 d**

35| To generate a spectral library from all acquired spectra above a certain iProphet cutoff and to convert all retention times into iRTs, run the following command after replacing the number following -cP with the cutoff you read out from the MAYU output in the step above:

```
spectrast -cNSpecLib -cICID-QTOF -cf "Protein! ~ reverse_"
-cP0.9774 -c_IRTiRT.txt -c_IRR iProphet.pep.xml
```

! CAUTION The iRT.txt file contains the peptide sequences to be used as iRT retention time reference peptides. This file needs to be adjusted in case reference peptides different from the ones suggested have been used.

▲ CRITICAL STEP The correlation coefficient R^2 of the linear regression should be >0.95. Open the spectrast.log file in a text editor and scroll to the end to see the linear regression equation and the R^2 .

? TROUBLESHOOTING

36| SpectraST consensus library generation. On high-mass-accuracy instruments, it may be useful to restrict the merging of spectra for consensus spectrum generation if they have unacceptably large retention time differences. Here we provide two options for consensus library generation: (Step 36A) a simple option that assumes that all fragment ion spectra are correctly assigned, and (Step 36B) a more sophisticated option that additionally considers retention times when merging spectra (**Box 4**).

(A) Consensus library unsplit

(i) Generate a consensus library by running the following command:

```
spectrast -cNSpecLib_cons -cICID-QTOF -cAC
SpecLib.splib
```

(B) Consensus library split

(i) To split distant, retention time-separated peptide identifications, run the following command:

```
python C:\Anaconda\Scripts\spectrast_cluster.py -d
2 SpecLib.sptxt
```

For the case study, this command results in nine output files.

(ii) To regenerate .splib, .splib and .pepid from the split .sptxt files, run the following command (This is equivalent to running nine times the command: 'spectrast -cNsplit-SpecLib_1 -cICID-QTOF SpecLib_1.sptxt' with adjusted numbers.):

```
FOR %A IN (SpecLib_*.sptxt) DO spectrast -cNsplit-
% ~ nA -cICID-QTOF %A
```

(iii) Generate a consensus library for each spectral library by running the following command (This is equivalent to running nine times the command: 'spectrast -cNcons-split-SpecLib_1 -cICID-QTOF -cAC split-SpecLib_1.splib' with adjusted numbers.):

```
FOR %A IN (split-SpecLib_*.splib) DO spectrast -
cNcons-% ~ nA -cICID-QTOF -cAC %A
```

(iv) Merge the consensus libraries back into a single consensus library ('grep' is a little executable which is installed together with the TPP):

```
grep -hUv ### cons-split-SpecLib_*.sptxt >>
SpecLib_cons_concat.sptxt
```

The splitting will add the tag 'Subgroup_xx_' in front of the protein name so that the different clusters of a peptide can be identified easily.

Part 5: assay library generation ● **TIMING 30 min to a few hours**

37| The last step is to convert a spectral library into an assay library for SWATH MS data analysis—i.e., to extract the most intense fragment ions for each peptide precursor. The SWATH windows can be defined in a simple table, which allows the script to disregard transitions for which the fragment ion falls into the same window as the precursor ion, as these typically result in noisy signals. If you plan to analyze your SWATH data with PeakView follow Step 37A, and for Skyline, Spectronaut or OpenSWATH follow Step 37B. The required input formats for each SWATH analysis software are listed in **Supplementary Tables 3–6**.

(A) For PeakView

(i) Extract most intense transitions from the spectral library:

```
python C:\Anaconda\Scripts\spectrast2tsv.py -l
350,2000 -s b,y -x 1,2 -o 6 -n 6 -p 0.05 -d -e -w
swaths.txt -k peakview -a SpecLib_cons_peakview.tsv
SpecLib_cons.sptxt
```

PROTOCOL

To run this command for the split library, replace the input file at the end of the command to 'SpecLib_cons_concat.sptxt'. The swaths.txt file contains the SWATH windows that are required to ignore transitions with fragment ion m/z values falling into their precursor SWATH windows.

! CAUTION The swaths.txt file needs to be adjusted to contain the SWATH window scheme that has been (or is to be) used for SWATH data acquisition.

! CAUTION The spectrast2tsv.py script recognizes common amino acid modifications, but if required additional ones can be specified using an additional input table. An example for this can be found in the msproteomicstools folder under analysis\spectra_libs\config_file_examples.

(B) For OpenSWATH, Spectronaut and Skyline

- (i) Extract most intense transitions from spectral library to generate the final SWATH assay library:

```
python C:\Anaconda\Scripts\spectrast2tsv.py -l
350,2000 -s b,y -x 1,2 -o 6 -n 6 -p 0.05 -d -e -w
swaths.txt -k openswath -a
SpecLib_cons_openswath.csv SpecLib_cons.sptxt
```

To run this command for the split library, replace the input file at the end of the command to 'SpecLib_cons_concat.sptxt'. The swaths.txt file contains the SWATH windows that are required to ignore transitions with fragment ion m/z falling into their precursor SWATH window.

! CAUTION The swaths.txt file needs to be adjusted to contain the SWATH window scheme that has been (or is to be) used for SWATH data acquisition.

! CAUTION The spectrast2tsv.py script recognizes common amino acid modifications, but if required additional ones can be specified using an additional input table. An example for this can be found in the msproteomicstools folder under analysis\spectra_libs\config_file_examples.

- (ii) This final SWATH assay library in table format can be directly used with the Skyline and Spectronaut software or further processed for use with OpenSWATH as described below in substeps (iii) and (iv). In the **Supplementary Tutorial**, we describe how to load the library exported in Step 37B(i) into Skyline and how to extract SWATH traces for visualization and data analysis.
- (iii) The OpenSWATH software requires the SWATH assay library to be in TraML format:

```
ConvertTSVToTraML -in SpecLib_cons_openswath.csv -
out SpecLib_cons.TraML
```

- (iv) The OpenSWATH software requires decoy transition groups to be present in the TraML assay library. Add decoy transition groups based on shuffled sequences:

```
OpenSwathDecoyGenerator -in SpecLib_cons.TraML -out
SpecLib_cons_decoy.TraML -method shuffle -append -
exclude_similar
```

? TROUBLESHOOTING

Troubleshooting advice can be found in **Table 2**.

TABLE 2 | Troubleshooting table.

Step	Problem	Possible reason	Solution
2, 35	iRT peptides were not spiked into the sample or the R^2 of iRT calibration <0.95	iRT peptides were not present in the sample or not all iRT peptides were correctly identified/detectable	If no iRT peptides were spiked into the samples or if they are not clearly detectable (or they show a bad correlation), it is possible to use endogenous peptides present in all samples to perform the retention time alignment, as described by Parker <i>et al.</i> , unpublished data. Alternatively, a higher concentration of the iRT peptides could be spiked into the samples to make sure that they can be correctly identified

(continued)



TABLE 2 | Troubleshooting table (continued).

Step	Problem	Possible reason	Solution
7	Error when running commands and during installation of OpenMS/ProteoWizard	A software dependency for the ProteoWizard tools (qtofpeakpicker and/or msconvert) is not present	ProteoWizard ^a requires .NET Framework 3.5 SP1 and 4.0 and Microsoft Visual C++ 2008 SP1 and 2010 SP1 Redistributable Packages to be installed. These should have been installed automatically during the installation of OpenMS ^b ; however, if the step was skipped or resulted in an error, then these packages might need to be installed manually. Go to Control Panel → Programs and Features → Turn Windows features on or off → activate Microsoft .NET Framework 3.5.1 (including both subfolders). In our experience, the error can even be ignored
7, 18, 26	Error when running a command	User does not have permissions to write a file into the target folder	Right-click on the folder and select 'Properties'. Go to the 'Security' tab and allow the users 'Full control'
19	pepXML is not generated for all X!Tandem search outputs	Bug in TPP web interface	Go to the 'Analysis Pipeline (Tandem)' tab and then to the 'pepXML' tab. Select here the .tandem files that have not been converted to pepXML, and click the 'Convert to PepXML' button
26	Comet gives an error	TPP version 4.7.0 comes with older comet version, which is not compatible with current parameter file	Uninstall TPP ^c and install version 4.7.1 as described in the installation part of the protocol
18, 26	Comet or X!Tandem gives an error	Search takes too long and Apache reaches timeout	Click on the Windows button in the lower left corner, and in the search field enter httpd.conf. It should suggest 'Edit the Apache httpd.conf Configuration File'. Scroll down until you see, close to the end of the file, the lines: # Add 5-h timeout Timeout 18000 Replace these two lines with: # Add 5-h timeout—changed to 1 week Timeout 604800 Save and close. Your user needs full rights on that file to be allowed to modify it: right-click on the file and select 'Properties'. Go to the 'Security' tab and allow the users 'Full control'
31	Protein of interest not covered in information-dependent acquisition runs for library generation	Protein is insufficiently abundant or is not expressed under the conditions used for library generation	Libraries that do not cover the proteins of interest could be topped up with synthetic peptides. Crude unpurified synthetic peptides can be purchased from JPT or Thermo for 10–20 EUR/USD per peptide

^aTo get further support for ProteoWizard tools, please consult the website (<http://proteowizard.sourceforge.net>) and email to the support list (support@proteowizard.org). ^bTo get further support for OpenMS tools, please consult the website (<http://open-ms.sourceforge.net>) and subscribe to the support mailing list (<http://sourceforge.net/p/open-ms/mailman>). ^cTo get further support for TPP tools, please consult the Wiki (<http://tools.proteomecenter.org/wiki/index.php?title=Software:TPP>) and subscribe to the spctools discussion group (<https://groups.google.com/forum/#!forum/spctools-discuss>).

● TIMING

Steps 1–3 (part 1), sample preparation and data acquisition: a few hours to several days, depending on the number of samples (excluding preparation of peptide samples)

Steps 4–8 (part 2), conversion and centroiding of the raw data: a few hours, depending on the number of samples

Steps 9–34 (part 3), database searching and controlling FDR: a few hours to several days, depending on the number of samples and search engines to be included



Steps 35 and 36 (part 4), spectral library generation: a few hours to 1 d, depending on the size of the library
 Step 37 (part 5), assay library generation: 30 min to a few hours, depending on the size of the library
 For the specific data set described in this protocol, we required 3 d for sample preparation (part 1, including sample preparation) and 2 d for the bioinformatics part (parts 2–5)

ANTICIPATED RESULTS

The final SWATH assay library generated from the three example yeast injections consists of 101,472 transitions belonging to 16,912 peptide precursors, 15,239 modified peptides, 14,948 stripped peptides and 1,948 unique proteins, and it can be directly used as an input for all currently available SWATH analysis software tools (the numbers correspond to the SWATH assay library without retention time–based splitting).

Accessing ProteomeXchange data.

The DDA and SWATH raw files of the case study have been deposited to the ProteomeXchange Consortium⁵⁸ via the PRIDE partner repository with the data set identifier PXD001126.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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AUTHOR CONTRIBUTIONS O.T.S., L.C.G. and B.C.C. developed the workflow and wrote the manuscript; P.N. developed the tools spectrast2tsv.py and spectrast_cluster.py; G.R. and H.L. developed and implemented the retention time normalization and iRT calibration in SpectraST; W.E.W. developed the qtofpeakpicker; D.A., P.M. and B.M. implemented automated SWATH library import into Skyline; and R.A. directed the project and contributed to writing the manuscript.

COMPETING FINANCIAL INTERESTS The authors declare competing financial interests: details are available in the [online version of the paper](#).

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