Evaluation of Multidimensional Chromatography Coupled with Tandem Mass Spectrometry (LC/LC–MS/MS) for Large-Scale Protein Analysis: The Yeast Proteome

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Highly complex protein mixtures can be directly analyzed after proteolysis by liquid chromatography coupled with tandem mass spectrometry (LC–MS/MS). In this paper, we have utilized the combination of strong cation exchange (SCX) and reversed-phase (RP) chromatography to achieve two-dimensional separation prior to MS/MS. One milligram of whole yeast protein was proteolyzed and separated by SCX chromatography (2.1 mm i.d.) with fraction collection every minute during an 80-min elution. Eighty fractions were reduced in volume and then re-injected via an autosampler in an automated fashion using a vented-column (100 μm i.d.) approach for RP-LC–MS/MS analysis. More than 162 000 MS/MS spectra were collected with 26 815 matched to yeast peptides (7537 unique peptides). A total of 1504 yeast proteins were unambiguously identified in this single analysis. We present a comparison of this experiment with a previously published yeast proteome analysis by Yates and colleagues (Washburn, M. P.; Wolters, D.; Yates, J. R., III. Nat. Biotechnol. 2001, 19, 242–7). In addition, we report an in-depth analysis of the false-positive rates associated with peptide identification using the Sequest algorithm and a reversed yeast protein database. New criteria are proposed to decrease false-positives to less than 1% and to greatly reduce the need for manual interpretation while permitting more proteins to be identified.

Keywords: proteome • tandem mass spectrometry • LC–MS/MS • vented column • Sequest criteria

Introduction

One of the greatest challenges facing researchers in the post-genomic era is to identify and quantify all expressed protein components in cells, tissues, and organisms. While two-dimensional gel electrophoresis (2DE) is a powerful technique for protein separation,1–3 it has a number of limitations that have spawned new technologies as alternatives to 2DE.4–12 A powerful alternative technique is the so-called MUDPIT (multidimensional analysis of proteins identification technology) pioneered by Yates and colleagues.13–18 This technique is also termed DALPC for direct analysis of large protein complexes.13,19,20 The acquired tandem mass spectrometry (MS/MS) data are used for database searching21–26 which leads to identification of peptides and proteins. At the heart of the method is the use of two-dimensional (2D) chromatography to separate a peptide mixture prior to analysis by mass spectrometry.

By utilizing peptides’ unique physical properties of charge and hydrophobicity, a complex peptide mixture can be effectively resolved and concentrated prior to sequence analysis by mass spectrometry. Strong cation exchange (SCX) chromatography is generally implemented as a primary separation technique due to its potential for increased loading capacity while reversed-phase (RP) chromatography is a perfect complement as a secondary separation technique because of its ability to remove salts and its direct compatibility with mass spectrometry through electrospray ionization.

2D chromatography can be accomplished by either an online or an offline approach.27 For the online approach, an acidified complex peptide mixture is applied to an SCX chromatography column and discrete fractions of the adsorbed peptides are sequentially displaced directly onto the RP chromatography column using a salt step gradient. Peptides are then eluted and analyzed by tandem mass spectrometry (MS/MS). This approach can utilize as many as 15 (or more) salt “bumps” to fractionate a peptide mixture.13,14 A principal advantage of the online technique is nearly complete automation in an unattended fashion. Essentially, the sample is loaded onto the SCX column and 24 h later the analysis is finished.

The offline approach is performed by applying the acidified complex peptide mixture to the SCX chromatography column followed by a binary gradient to high salt to elute the peptides. Fractions are typically collected every minute into a 96-well plate and reduced in volume, and then each fraction is loaded onto a RP chromatography column automatically via an autosampler and analyzed by LC–MS/MS. The offline technique has several advantages over the online technique: (i)
peptide separation is superior using a linear gradient instead of a step-gradient; (ii) ideally, SCX chromatography is performed with >20% acetonitrile (AcN) in the buffers to linearize peptides, yet only 5–10% AcN can be tolerated with the online approach; (iii) more peptide fractions can be collected with the offline approach (e.g., 96 vs 15); and (iv) user discretion as to which fractions are to be analyzed is available in the offline approach, and interesting fractions can be re-visited (reanalyzed).

We report here the use of an off-line LC/LC–MS/MS strategy to catalog the expressed proteins of the yeast *S. cerevisiae*. One milligram of total protein was proteolyzed and analyzed by LC/LC–MS/MS. A large dataset was generated that allowed us to estimate false-positive rates during database searching using the Sequest program and led to the establishment of new criteria to reduce the false-positive rate to less than 1%.

**Experimental Section**

**Apparatus.** An LCQ Deca ion trap mass spectrometer was purchased from Thermo Finnigan (San Jose, CA). Accessories for capillary liquid chromatography were from Upchurch Scientific Inc. (Oak Harbor, WA). Capillary tubing containing a borosilicate frit (Integrafrit) was from New Objective Inc. (Cambridge, MA). Magic C18AQ resin (5 μm, 200 Å) was from Michrom BioResources (Auburn, CA).

**Reagents.** Sequencing grade modified trypsin was purchased from Promega (Madison, WI). High purity acetonitrile (AcN), methanol, and acetic acid (HOAc) were from VWR (Bridgeport, NJ), and heptafluorobutyric acid (HFBA) was from Pierce (Rockford, IL).

**Procedure. Preparation and Digestion of Total Cell Lysate from *S. cerevisiae*.** Strain S288C was grown to log phase in YPD medium. The cells were collected by centrifugation and lysed with glass beads in lysis buffer (20 mM Tris, pH 8.0, 10 mM NaF, 10 mM NaCl, 0.1% deoxycholic acid, 0.5 mM EDTA, and protease inhibitor cocktail from Roche). The lysate containing 1 mg of proteins was denatured by adding urea to 8 M and SDS to 0.1%, albumin (BSA) as standard. The lysate containing 1 mg of proteins was denatured by adding urea to 8 M and SDS to 0.1%, albumin (BSA) as standard. The lysate containing 1 mg of proteins was denatured by adding urea to 8 M and SDS to 0.1%, albumin (BSA) as standard.

**Two-Dimensional Liquid Chromatography with Tandem Mass Spectrometry.** The trypic peptides were acidified with TFA to 0.1%, supplemented with AcN to 25%, and then loaded onto a 2.1 mm × 20 cm polysulfonethyl A column (Poly LC Inc., Columbia, MD) connected to a Beckman Gold HPLC. Fractions were collected every minute during an 80-min gradient from 5% to 35% solvent B (solvent A: 5 mM phosphate buffer, 25% AcN, pH 3.0; solvent B: the same as A with 350 mM KCl; flow rate: 0.2 mL/min). All collected fractions (n = 80, 0.2 mL each) were reduced in volume to ~0.1 mL to remove AcN by vacuum centrifugation and then analyzed by RP-LC–MS/MS in a completely automated fashion using a 100 μm i.d. vented column as described. We loaded as many as 29 fractions onto a single V-column. In other work, we used a single V-column to analyze more than 100 samples (data not shown). Peptides were eluted for each analysis during a gradient from 10% to 30% buffer B (buffer A: 0.4% acetic acid, 0.005% HFBA, 5% AcN; buffer B: the same as A except 95% AcN; flow rate: ~300 nL/min). In general, the amount of sample loaded was adjusted according to its peptide concentration: about 50% of the total volume for fractions with OD214 < 1.2 and 25% for all others (OD214 at least 1.2, fractions 14–27). Gradient length was largely dependent on the peptide concentration estimated by its OD214 absorbance: 60 min (fractions 1–8 and 67–80, OD214 < 0.2); 90 min (fractions 42–66, OD214 between 0.2 and 0.5); 120 min (fractions 28–41, OD214 between 0.2 and 0.5), or 150 min (fractions 9–27, OD214 > 0.5). Peptide ions were detected in a survey scan from 400 to 1700 amu (3 μs scans) followed by five data-dependent MS/MS scans (5 μs scans each, isolation width 3 amu, 35% normalized collision energy, dynamic exclusion for 3 min) in a completely automated fashion on an LCQ-DECA ion trap mass spectrometer (Thermo Finnigan, San Jose, CA).

**Data Processing.** All MS/MS spectra were searched using the Sequest algorithm. The database utilized was a composite of the known yeast ORFs (6139 proteins) and a second database containing 6139 bogus proteins created by precisely reversing the order of the amino acid sequence for each protein so that the C terminus became the N terminus. This is similar to a database created by Moore et al. A total of 162 000 MS/MS spectra were searched against the database using the following criteria: no requirement for trypsin digestion, variable modification of methionine (+16 Da) and cysteine (+57 Da). The reason for the addition of the reverse-database is to provide an estimate of the false-positive rate of peptide identification by Sequest under various parameters.

**Results**

A flowchart of the experiment is presented in Figure 1. Whole yeast extract (1 mg protein) was trypsinized into peptides in the presence of SDS and urea. The resulting peptides were
separated by strong cation exchange (SCX) chromatography with fraction collection. Each fraction \((n = 80)\) was collected and analyzed by automated reversed-phase (RP) chromatography coupled with tandem mass spectrometry using a vented column approach. In total, more than 162,000 MS/MS spectra were automatically collected and searched against the composite database (see the Experimental Section). More than 26,800 peptides were identified (7537 unique peptides, see the Supporting Information).

As an example, the analysis of fraction 36 is shown in Figure 2. The off-line SCX analysis (Figure 2A) resulted in the collection of 80 discrete fractions. The LC–MS/MS analysis of fraction 36 is shown in Figure 2B. Figure 2C shows a single MS survey scan during LC–MS/MS analysis at time 109.6 min. Many peptides can be seen coeluting. The mass spectrometer sequentially selected five peptide ions for further sequence analysis by collision-induced dissociation. (D) MS/MS scan of precursor ion \(m/z 1010.7\). This was one of the five peptide ions chosen for analysis which was later matched to the peptide amino acid sequence shown. Some \(b\)- and \(y\)-ions derived from the peptide ion are also indicated.

Sequest matches an MS/MS spectrum with a peptide sequence using the following steps. (i) All theoretical peptides with molecular masses isobaric to that of the precursor ion for an MS/MS spectrum are extracted from a database. (ii) Each peptide is given a preliminary score by examining the number of predicted fragment ions from the database peptide that match the acquired fragment ions in the MS/MS spectrum. (iii) The 500 best-matching peptides undergo a more rigorous ion-matching algorithm that generates a Sequest cross-correlation score (\(X_{corr}\)). A list of the best-matching peptides is returned to the user with the top-scoring peptide being considered the best candidate. The difference of \(X_{corr}\) scores between the top peptide and the runner-up is represented by the function termed delta-correlation score (\(\Delta C_n\)).

In most large-scale peptide sequence analysis studies where Sequest is utilized, the results of the search for each MS/MS spectrum are filtered according to some established criteria to achieve a list of peptide matches. The criteria utilized include three parameters: state of tryptic ends (fully- or partially tryptic), \(X_{corr}\), and \(\Delta C_n\). Although this data processing procedure is sophisticated, the standard for accepting matched peptides is rather empirical, and a large number of spectra require manual interpretation. This dataset presented a unique opportunity to evaluate the rate of false-positives achieved by varying these three parameters because of the size of the dataset. More than 162,000 MS/MS spectra and more than 20,000 spectra easily matched to correct peptides. Hence, the
HPLC pump, two six-port valves, one micro-tee, one micro-cross, a vented RP column and a LCQDECA ion trap mass spectrometer. SCX according to following formula: % fal = \( \frac{100 \times n_{\text{rev}}}{n_{\text{total}}} \), where % fal is the estimated false-positive rate, \( n_{\text{rev}} \) is the number of peptides identified (after filtering) from the reverse-database, and \( n_{\text{total}} \) is the total number of identified peptides from both databases found from the reverse-database and dividing the result by the rate can be estimated by doubling the number of peptides same for both databases. Therefore the overall false-positive acids), it can be assumed that the rate of false-positives is the from the reverse-database is defined as a false-positive (incorrect) orientation. The reverse-database was positioned when the same threshold of Xcorr was applied, fully tryptic peptides had a much lower false-positive rate than partially tryptic peptides. Based on the Table 1, we determined new criteria (highlighted in the Table 1) with the goal of an overall estimated false-positive rate of less than 1%: \( \Delta C_n \) score > 0.08; in the case of fully tryptic peptides, Xcorr should be larger than 2.0, 1.5, or 3.3 for charge states of +1, +2, +3, respectively; partially tryptic must have an Xcorr score of > 3.0 (+2 state) or > 4.0 (+3 state). We next compared these new criteria with those previously delineated by Yates and colleagues.14 Our dataset was processed (filtered) according to both criteria (Figure 5A). Our criteria resulted in an approximately 3-fold lower estimated false-positive rate (10.6% vs 30.8%) for identified proteins before manual validation, which decreased the requirement of manual interpretation. To further analyze the false-positives in identified proteins, the proteins were categorized by the number of unique peptides contributing to their identification (Figure 5B). The false-positive rate among proteins identified by a single peptide was 26% (188 out of 716). This rate decreased dramatically when two peptides were utilized in the identification to 1.4% (4 out of 279). Finally within this very large dataset, there were no false-positives detected for proteins identified by three or more peptides. Therefore, we manually confirmed proteins identified by one or two peptides instead of all proteins matched by four or fewer peptides.14 A total of 216 (12.6% of 1720) proteins were discarded, which was consistent with the predicted false-positive rate (10.6% in Figure 5A). In our dataset, proteins identified by a single peptide after manual interpretation represented 33.9% (510 out of 1504) of the entire dataset that was significantly less than that in the dataset published previously (42.1%, 617 out of 1465).14 Furthermore, while 858

Figure 3. Automated nanoscale microcapillary LC–MS/MS using a vented column. (A) Integrated scheme including an autosampler, HPLC pump, two six-port valves, one micro-tee, one micro-cross, a vented RP column and a LCQDECA ion trap mass spectrometer. SCX fractions were first loaded onto a sample loop in valve 1 that was connected by dashed lines. Valves 1 and 2 were then switched (connected by solid lines), which closed the micro-tee (flow restrictor; split) and opened the micro-cross (vent). At this stage, the sample was transferred to the trap region of the vented-column and desalted by extensive wash at high flow rate (up to 5 μL/min). After sample loading at high flow rate, valve 2 was again switched (dashed-line connection), which opened the flow-restrictor (split) and closed the vent. Peptides bound to the trap were eluted during a RP gradient and ionized by electrospray with voltage (1.8 kV) applied via a gold wire to one arm of the cross. All 80 fractions were analyzed by this same procedure. (B) A standard peptide bradykinin (monoisotopic m/z of 530.8 for doubly charged ion) was sequentially titrated from 10 fmol to 1 pmol using a 100 μL i.d. nanoscale vented-column. Selected ion chromatograms for the doubly charged peptide ion (m/z 530.3–533.3) are shown. The amount of the peptide loaded is also indicated as well as the strength of signal recorded.
proteins were found in both datasets, 607 proteins were unique to the published dataset and 646 proteins were found only in our dataset (Figure 5C).

We also examined the elution profile of the peptides with different solution charge states. At the pH utilized for the separation (pH 3.0), only lysine, arginine, histidine, and the amino-terminus contribute to the peptide solution charge state. Peptides with different solution charges were well separated mainly based on their ionic charge (Figure 6A) but also dependent on hydrophobicity (Figure 6B) during the elution.

**Figure 4.** Evaluation of the effect of ΔCn on the false-positive rate of peptide identifications. A total of 162,000 MS/MS spectra were searched against the chimeric forward- and reverse-yeast database. Sequest analysis results were filtered for fully tryptic peptide ends and a charge state of +2. The effect of increasing the ΔCn cutoff at different Xcorr values is shown. A false-positive was defined as the identification of a peptide from the reverse-database after filtering. The estimated false-positive rate was calculated as described in the text. The estimated percent of false-positives at an Xcorr > 1.5 and a ΔCn of > 0.08 was less than 1% (170) with an n of more than 20,000.

**Table 1.** Evaluation of the False-Positive Rate Using Different Xcorr Cutoff Thresholds

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FUL, fully tryptic ends; PAR, partially tryptic ends; NON, non-tryptic; % FAL, the percentage of false-positives; N, the sum of peptides based on the Xcorr cutoff (shown in the first column). All peptides display ΔCn of > 0.08. The established thresholds are shown in boldface type.
The number of peptides identified in each fraction was outlined in Figure 7A. The number of identified proteins increased steadily with the number of fractions analyzed and did not reach a plateau (Figure 7B). Although the slope appears to decrease gradually, if the curve is normalized according to the number of peptide identified in each SCX fraction (Figure 7A), it is clear that a plateau is not being reached (data not shown). To increase the throughput of the method, one strategy is to decrease the redundant MS/MS analysis of peptides from the same fraction. Analyzing every other fraction would reduce the analysis time by 50% and would still allow the identification of 84% of the same proteins. Analyzing only every third fraction resulted in 74% of the same proteins being identified (Figure 7C).

Discussion

Multidimensional chromatography coupled to tandem mass spectrometry (LC/LC-MS/MS) represents a promising alternative for proteome analysis. Multidimensional chromatography is most commonly accomplished by the combination of SCX and RP chromatography because of their orthogonal separation properties. In this report, we present the analysis of expressed yeast proteins by LC/LC-MS/MS techniques. More than 162,000 MS/MS spectra were collected throughout the analysis with a final result of identification of 1504 proteins.

The sensitivity of peptide detection in LC-MS/MS is largely dependent on the number of false-positives. A total of 162,000 MS/MS spectra were searched against the chimeric database described in the text that contained yeast ORFs in both the forward and reverse direction. (A) Comparison of the criteria (marked by asterisk) previously described by Yates and colleagues (ref 14) and our new criteria from Table 1. While peptides with ΔC₈ > 0.1 were accepted in the previous criteria, we selected ΔC₈ > 0.08. Ion charge state (+1, +2, or +3) and tryptic state (ful, fully; par, partially; or non, non-tryptic) are indicated. The sum of peptides (n = nrev + nreal) in each category, the number of false-positives (fal = 2nrev) and the percentage of false-positives (%) are also shown. After manual validation, we removed 216 proteins (12.6%) and accepted 1504 proteins. (B) Assignment of proteins according to how many unique peptides used in their identification for this study (1504 proteins) and one previously published (ref 14) (1465 out of 1486 proteins with extractable peptide information). (C) The overlap in the proteins detected in this study and those previously described (ref 14).

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A two-dimensional separation and analysis of proteolyzed yeast protein has already been presented by Yates and colleagues. They reported the analysis of 1486 yeast proteins by a combination of SCX and RP chromatography with peptide sequence analysis by tandem mass spectrometry. It should be noted that their report actually represents the pooled results from several sources. We therefore sought to evaluate the rate of false-positives after filtering database search results. We created a chimeric or composite database that contained protein sequences for all yeast ORFs in the forward direction followed by the reverse-database and 21,071 from the true database) when the reverse-database was often small, it required a very large number of right-answer peptides to provide an accurate estimate. We made use of a vented-column for the automated analysis of the 80 SCX fractions. This technique allowed for effect of analyzing only every other or every third fraction from the 80 SCX fractions on total number of peptides, unique peptide and proteins identified. The number of total peptides (26,815), unique peptides (7,537) and proteins (1,504) were normalized to 100%. Key: 1, all fractions with odd numbers; 2, all fractions with even numbers; 1, 4, fractions 1, 4, 7, ...; 2, fractions 2, 5, 8, ...; 3, fractions 3, 6, 9, ...

214 nm. For most fractions, we loaded only one-half of the sample volume to maximize sensitivity. For several fractions with much higher peptide concentrations, some peak broadening effect was observed (data not shown) and we reduced the loading amount to 25%. To provide the ability for adjustment to the amounts of sample loaded and gradient length, we first analyzed every 8th fraction (fractions 8, 16, 24, 32, 40, 48, 56, 64, 72, and 80).

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214 nm. For most fractions, we loaded only one-half of the sample volume to maximize sensitivity. For several fractions with much higher peptide concentrations, some peak broadening effect was observed (data not shown) and we reduced the loading amount to 25%. To provide the ability for adjustment to the amounts of sample loaded and gradient length, we first analyzed every 8th fraction (fractions 8, 16, 24, 32, 40, 48, 56, 64, 72, and 80).

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In the case of partially tryptic peptides, $X_{corr}$ cutoff values of 3.0 and 4.0 for 2+ and 3+ peptides, respectively, are used. Partially tryptic peptides with a charge state of +1 and nontryptic peptides of any charge state are not accepted. If this dataset were filtered with the previously established criteria we would need to remove ~630 peptides compared to only ~180 by using the new criteria (Figure 5A). Furthermore, the new criteria resulted in identification of more proteins (1538 vs 1419) that were calculated by subtracting the estimated false-positives from the total proteins shown (Figure 5A).

We have evaluated an offline multidimensional chromatography peptide separation approach for the large-scale analysis of expressed yeast proteins. We identified 7537 peptides (1504 proteins) during a completely automated analysis. Our data suggest that additional protein or peptide separations would still be required to maximize the separation capacity to meet the challenge of the complexity of a higher eukaryotic proteome. For instance, enriching for cysteine-containing peptides before two-dimensional LC–MS/MS would decrease the peptide complexity, reduce the number of peptides identified for the same proteins and therefore improve throughput for protein identification. Alternatively, removing the time factor could be accomplished by the deposition of RP-LC elutions directly onto plates for analysis by MALDI ionization.

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Supporting Information Available: Table of unique peptide sequences. This material is available free of charge via the Internet at http://pubs.acs.org.

References

PRO2556V