Microproteomics: Analysis of protein diversity in small samples

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MICROPROTEOMICS: ANALYSIS OF PROTEIN DIVERSITY IN SMALL SAMPLES

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Received 10 September 2007; accepted 6 December 2007

Published online 12 February 2008 in Wiley InterScience (www.interscience.wiley.com) DOI 10.1002/mas.20161

I. INTRODUCTION

The last decade has seen a resurgence of interest in the study of the proteome—the set of proteins that can be expressed by the genetic material of an organism. Advancements in protein extraction, purification, and identification have been driven by improvements in sensitivity, throughput, and resolution of separations and mass analyzers. However, proteomic approaches have not advanced to the extent of genomic technologies. Efforts to adapt standard methods of tissue sampling, protein extraction, arraying, and identification are reviewed, with an emphasis on those appropriate to smaller samples ranging in size from several microliters down to single cells. The effects of miniaturization on these analyses are highlighted using neuroscience-related examples, as are statistical issues unique to the high-dimensional datasets generated by proteomic experiments. © 2008 Wiley Periodicals, Inc., Mass Spec Rev 27:316–330, 2008

Keywords: 2D gel electrophoresis; imaging mass spectrometry; capillary electrophoresis; microdissection

Proteomics, the large-scale study of protein expression in organisms, offers the potential to evaluate global changes in protein expression and their post-translational modifications that take place in response to normal or pathological stimuli. One challenge has been the requirement for substantial amounts of tissue in order to perform comprehensive proteomic characterization. In heterogeneous tissues, such as brain, this has limited the application of proteomic methodologies. Efforts to adapt standard methods of tissue sampling, protein extraction, arraying, and identification are reviewed, with an emphasis on those appropriate to smaller samples ranging in size from several microliters down to single cells. The effects of miniaturization on these analyses are highlighted using neuroscience-related examples, as are statistical issues unique to the high-dimensional datasets generated by proteomic experiments. © 2008 Wiley Periodicals, Inc., Mass Spec Rev 27:316–330, 2008

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I. INTRODUCTION

The last decade has seen a resurgence of interest in the study of the proteome—the set of proteins that can be expressed by the genetic material of an organism. Advancements in protein extraction, purification, and identification have been driven by improvements in sensitivity, throughput, and resolution of separations and mass analyzers. However, proteomic approaches have not advanced to the extent of genomic technologies. Over the past 30 years, major technological innovations have improved the throughput of genomic sequencing and analysis techniques to the point where sequencing the complete genome of organisms is commonplace. In addition, transcriptomic analysis is feasible in samples as small as single cells. Unfortunately, the analytical capabilities of proteomic technologies lack the speed and robust characteristics of current microarray methods. Given these limitations, why should one pursue proteomic studies? Clearly, the promise implied in proteomics is acquiring global information on changes in protein expression, including post-translational modifications (PTMs) from specific biological tissues under unique physiological conditions. Such information certainly is relevant to biological function, because proteins play an integral role in cell function.

It should not be surprising that changes in gene expression might not correlate well with changes in protein expression (Anderson & Seilhammer, 1997; Futcher et al., 1999); proteins are activated by mechanisms such as precursor cleavages, PTMs, and unique localization, which are not direct consequences of changes in protein production. Furthermore, many physiological responses of organisms are not always genetically mediated. For example, homeostatic responses are relatively uniform despite the genetic variability between specific individuals. One final, but perhaps most important point, the lifetime of proteins can vary by orders of magnitude so that the amounts of two proteins can be surprisingly different even with similar translation rates. Thus, proteomic characterization complements transcriptomic studies, and might be more relevant for the characterization of complex genetic and homeostatic phenomena.

Complexity presents a fundamental challenge in proteomics. Although the genetic code is composed of four bases, the “protein code” consists of over 20 amino acids plus many possible PTMs. The genome for many organisms is known, and does not vary between cells. Although each organism has only one genome, it has many transcriptomes and even more proteomes. Protein expression patterns differ among cell types, and physiological changes alter these expression patterns; these alterations also differ between specific cells of the same type.

Also of concern, tissues of interest can be microscopically small and surprisingly heterogeneous, requiring the dissection and characterization of specific cells from a region of interest. Whereas recent technological developments have made microdissection feasible (see below), extracting, separating, and detecting the wide range of proteins present in extremely small samples present significant issues. Geneticists have overcome these obstacles with the development of the polymerase chain
reaction to amplify DNA. However, analogous methods do not exist for protein amplification.

In the classical proteomic workflow, one obtains the tissue of interest, extracts proteins from the tissue, separates and/or arrays the proteins, identifies proteins of interest, and quantifies changes in protein expression and PTMs. Each step requires greater care and expertise as the sample size decreases. Obligate losses during the extraction and fractionation procedures become more of an issue as one draws closer to the limits of protein detection. Protein diversity decreases in small samples, and fewer proteins will be abundant enough to exceed the sensitivity limit of specific technologies. It also becomes more difficult to evaluate differences between conditions with small sample sizes, because variability increases as one operates closer to the limits of the analytical techniques. Although information content is decreased in smaller samples so that only higher abundance compounds are detected, useful knowledge can still be gained, even from a single cell (Rubakhin, Greenough, & Sweedler, 2003; Hummon, Amare, & Sweedler, 2006; Rubakhin et al., 2006; Neupert et al., 2007; Predel et al., 2007; Rubakhin & Sweedler, 2007).

The main focus of this review is to describe recent developments that have made “microproteomics” a reality. Sampling issues are covered first, followed by a discussion of new methods that have been developed to improve the sensitivity and selectivity of conventional sampling techniques, such as laser capture microdissection (LCM). Approaches to protein separation, including multidimensional separations and smaller gels, are described. Improvements in mass spectrometric resolution and sensitivity are also presented, as well as innovative statistical methods for analyzing and cataloging the complex data generated by these experiments. Approaches to characterize preselected proteins such as with immunoaffinity approaches or GFP-labeling are not covered. Protein arrays based on antibodies or other selective capture agents are an exciting area undergoing rapid development, and have been the focus of several recent reviews (Borrebaek & Wingren, 2007; Lu & Liu, 2007; von Eggeling, Melle, & Ernst, 2007). In this review, mass-spectrometric-based identification methods are highlighted. We briefly discuss recent developments, such as mass spectrometric imaging (MSI), which bypass steps in the conventional proteomic workflow and allow small samples to be characterized. As the dynamic range of these cutting-edge technologies improves, these “shorter” approaches may provide attractive alternatives for the proteomic analysis of small samples.

II. TISSUE SAMPLING

The first and most critical issue facing anyone trying to analyze small samples is how to acquire the optimal specimen of interest. If a poor quality sample is obtained, or if proteins are degraded or contaminated with other unwanted tissue types, then even the best protein identification techniques will not rescue the experiment. There are many possible techniques that can be employed; choosing the appropriate one depends on experimental requirements. For example, if anatomical precision is not required, then freehand microdissection can be used. Flow cytometry is useful for identifying cells that express specific biomarkers, but anatomic orientation and resolution is lost. These limitations are also shared by some emerging technologies, such as dielectrophoresis-activated cell sorting (Hu et al., 2005c), use of optical tweezers (Grier, 2003), and immunomagnetic separation (Safarik & Safarikova, 1999). Several approaches have been developed to reproducibly obtain small samples of defined anatomical regions. An excellent method for microscope-aided microdissection in the CNS is described by Cuello and colleagues (Cuello & Carson, 1983), which relies on the differential light transmittance of myelinated and unmyelinated tissues in fresh brain slices. Because myelination decreases light transmittance, myelinated tissues appear dark when transilluminated and unmyelinated tissues appear lighter. With 2–300 μm sections used, this method can provide a surprising degree of anatomical detail and precision. It is important to maintain temperatures as close to freezing as possible while ensuring that ice crystals do not develop during the microdissection process. If they do, much of the anatomical resolution inherent to transillumination is lost. Further, if temperatures rise, then significant protein degradation can occur. The addition of protease and phosphatase inhibitors to bath solutions should also be considered to help improve the consistency of results.

Another useful sampling technique, the micropunch method, was developed by Palkovits and Brownstein (1983). Punch microdissection entails making small punches to remove tissue from anatomically defined regions of fresh or frozen samples, and can also be aided by the use of a loupe or dissecting microscope. Although anatomical delineation is not as precise as with transillumination, the ability to immediately snap-freeze specimens and keep them frozen throughout the dissection process is a huge advantage for proteomic studies. Micropunch and transillumination both require substantial amounts of technical skill and practice to obtain the requisite precision and reproducibility.

When examining invertebrate neurons, individual cells can be in the range of 20–500 μm in diameter, a size that allows direct sampling of these cells, from microdissection to subsequent protease treatments in preparation for matrix-assisted laser desorption/ionization mass spectrometry (MALDI MS) (Li, Garden, & Sweedler, 2000). As shown in Figure 1, individual neurons can be divided into subsections such as soma and neurites, and each assayed for their neuropeptide content (Rubakhin, Greenough, & Sweedler, 2003). Similarly, microdissection and/or isolation of mammalian cells (e.g., red blood cells, pituitary cells) have been demonstrated to be effective for direct analysis using mass spectrometry (MS). Single cell analysis of pituitary cells using MALDI MS has allowed positive identification of 14 peptides derived from pro-opiomelanocortin prohormone (Rubakhin et al., 2006). Similarly, using capillary electrophoresis (CE) coupled to an electrospray ionization-Fourier transform ion cyclotron mass spectrometer, Hofstadler et al. (1995) were able to identify α- and β-chains of hemoglobin from a few red blood cells. Furthermore, MALDI MS has been shown to be robust in analyzing peptides in samples as small as 2 μm dense core vesicles (Rubakhin et al., 2000). Analysis at the single cell or organelle level can provide a comprehensive view of how an individual cell processes protein products with fewer confounding factors.
A relatively recent improvement in tissue sampling is the development of LCM by Emmett-Buck and colleagues (Emmett-Buck et al., 1996). This technique uses a finely-focused laser to either cut around cells of interest or adhere them to a plastic film, depending on the instrument. Slices of ethanol-post-fixed tissues from 8–40 μm thick can be dissected, depending on the instrument and exact technique employed. Several investigators have adapted LCM to proteomic studies. As shown in Figure 2, we demonstrated that LCM and the required ethanol postfixation (30 sec in 70% ethanol) does not interfere with two-dimensional gel electrophoresis (2D)E-based protein separation (Moulédous et al., 2002). However, we found that conventional histological staining methods in brain (creosyl violet, hematoxylin/eosin, and toluidine blue) as well as some more unusual stains (chlorazol black E, Sudan black B) did interfere with 2DE-based protein separation (Moulédous et al., 2002). In some tissues, such as kidney and esophagus, conventional tissue-staining techniques can be used to provide anatomic detail without interfering with protein recovery and identification (Banks et al., 1999; Emmett-Buck et al., 2000). However, we have found that in brain, higher stain concentrations are required and interfere with protein recovery.

An alternative to direct sample staining is the technique of “navigated” LCM (Wong et al., 2000; Moulédous et al., 2003a). In this method, a stained section is used to guide the dissection of an adjacent, unstained section of tissue. Accurate image registration is necessary to ensure the success of this technique, which can be accomplished by the use of a variety of image analysis programs. Such fine dissections permit detailed questions to be addressed; for example, what are the proteomic changes seen in a specific type of cell expressing a specific receptor? This issue can be elucidated by using “immuno-LCM” to harvest cells expressing specific markers (Moulédous et al., 2003b). However, conventional avidin-biotin chemistry interferes with protein extraction and 2DE. Craven and colleagues also showed that immunogold staining in renal tissue resulted in the loss of 60% of total protein (Craven et al., 2002). Fortunately, fluorescently tagged antibodies do not significantly reduce protein recovery, nor do they interfere with subsequent analyses (Moulédous et al., 2003b). Moreover, immunostaining does not significantly change the percentage of protein coverage for proteins identified with peptide mass fingerprinting (PMF) after 2DE. Thus, with the aid of a fluorescent-capable LCM, more detailed and precise studies can be undertaken. The tradeoff is that much more starting tissue is required to perform proteomic studies on specific subsets of cells.

Several groups have combined LCM with the technique of differential in-gel electrophoresis (DIGE; also see section VII), a technique to reduce measurement variability in 2D gels (Zhou et al., 2002; Lee et al., 2003; Greengauz-Roberts et al., 2005; Wilson et al., 2005; Kondo & Hirohashi, 2006; Sitek et al., 2006). One recent study demonstrated the ability to detect 900 protein spots on 2D gels generated from less than 1 μg of kidney glomeruli using LCM followed by DIGE (Sitek et al., 2006). Other modes of protein separation have also been effectively coupled with LCM. One recent study captured approximately 1500 cells using LCM, then subjected the sample to nano-LC followed by FT-MS identifications (Mustafa et al., 2007). Using this approach, 4 proteins that were uniquely expressed in the blood vessels of brain tumors were identified. Several investigators have also shown that proteins can also be identified by direct mass spectrometric analysis of laser-captured tissues using MALDI-MS and SELDI-MS (Xu et al., 2002; Bhattacharya, Gal, & Murray, 2003; Kwapiszewska et al., 2004; de Groot et al., 2005; Guo et al., 2005; Kriegl et al., 2005). Caprioli and colleagues demonstrated that meaningful spectra could be obtained from samples containing as few as 10 laser-captured cells (Xu et al., 2002).

III. PROTEIN EXTRACTION

Once tissue samples have been obtained, the next step is to extract the proteins from the tissue. Although this step is perhaps the most critical one in performing a successful proteomics experiment, it has been given the least attention by current investigators. All extractions are aimed toward an elusive and as yet unachieved goal, the complete extraction of all proteins from a sample. The wide range of physicochemical properties of cellular proteins and their differing solubilities make it difficult to efficiently extract all proteins, which range in size from small cytoplasmic signaling molecules to lipophilic, megadalton membrane receptors. However, the introduction of new techniques and optimization of classical methods have improved extraction efficiencies for small samples. Extractions have been categorized from “gentle”, which consists of osmotic, chemical, or enzymatic treatments or mild grinding, to “vigorous”, which are active mechanical cellular disruption methods (Scopes, 1994). In order to achieve as
complete an extraction as possible, most investigators use a combination of cellular disruption and treatments with strong detergents and chaotropes. Although mechanical cell disruption might improve extraction efficiencies over simple chemical extraction, there is no method of cell disruption that is clearly superior for small samples. The only limitation is the size of the probes or vessels available for sample processing. Ultrasonication, homogenization, freeze-thawing, pressure-cycling, and bead mills have all been used (Schutte & Kula, 1990a,b; Hopkins, 1991; Melendres et al., 1991; Belo et al., 1996; Raynie, 2004; Butt & Coorssen, 2006; Flanagan et al., 2006; Smejkal et al., 2006). We have had success in using ultrasonication with small-diameter probes. Methods such as bead mills and pressure cycling, an extension of the old “french press” technology, are currently difficult to use with the smallest samples. However, they do have potential advantages for certain types of tissues and extractions that warrant re-examination when the equipment has been appropriately miniaturized. Care must be taken to avoid any heating of the sample during cell disruption, which can degrade proteins prior to subsequent analyses.

Chemical solubilization, the second phase of the process, has also improved markedly over the past several years. Solubilization has three major components: (a) minimize interactions between proteins as well as between proteins and other substances (e.g., nucleic acids, lipids); (b) remove contaminants; and (c) prevent protein precipitation during the process of protein separation (Rabilloud, 1996). The chemical strategies employed are dependent on (a) whether the proteins are needed in their native or a denatured conformation; (b) which nonprotein substances need to be removed from the sample; and (c) the subsequent separation technology to be used. The goal of denaturation is to eliminate protein interactions by disrupting noncovalent bonds as well as covalent disulfide bonds. This will disrupt secondary and tertiary protein structure and so is not desirable when examining protein complexes. However, denaturation improves the ease and effectiveness of extraction. A wider variety of solvents and

FIGURE 2. The effect of tissue fixation and LCM on 2D gel-based protein separation. Samples of the rat brain region called the striatum were used to determine whether fixation or dissection method affected protein separation using 2-D gel electrophoresis. Three conditions were tested: samples were either manually dissected without fixation, fixed in ethanol and then manually dissected, or fixed and dissected using LCM. Twenty-five microgram of total protein was loaded onto immobilized pH gradient (IPG) strips and separated by 2-D gel electrophoresis as described. A: Unfixed, unstained, manually dissected sample. B: Ethanol-fixed, manually dissected sample. C: Ethanol-fixed, LCM sample. D: Number of spots detected on each gel. Results are expressed as mean ± SEM of three independent experiments. No statistically significant difference was observed between any of the treatment groups. Reprinted with permission from Mouledous et al. (2002), copyright 2002 Association of Biomolecular Resource Facilities.


constituents will remain unidentified (Bishop, 1976), and has perhaps an order of magnitude higher chemical diversity due to chemical modifications; however, the highest resolution separation methods currently available can only distinguish a few thousand distinct species (Anderson, 1995). Therefore, sample complexity should be reduced or many constituents will remain unidentified. A commonly suggested first step is fractionation of cells to focus on a specific organelle or region (e.g., synaptic cleft) of interest. Taken to an extreme, a tissue can be fractionated so that only a single compound will be isolated via immunohistochemical approaches. However, also application of the emerging field of nanotechnology to the extraction and separation of proteins suggests that more thorough coverage of the proteome from samples as small as a single cell may soon be possible (Hellmich et al., 2005; Ivanov et al., 2006). Recently, the detection of single proteins, some of low abundance, from individual cells has been demonstrated (Cai, Friedman, & Xie, 2006; Huang et al., 2007).

IV. SEPARATION OF PROTEIN MIXTURES

There are limits to the number of proteins that can be detected and identified from a single protein mixture. It has been estimated that the average cell expresses 10–15,000 unique proteins (Hastie & Bishop, 1976), and has perhaps an order of magnitude higher chemical diversity due to chemical modifications; however, the highest resolution separation methods currently available can only distinguish a few thousand distinct species (Anderson, 1995). Therefore, sample complexity should be reduced or many constituents will remain unidentified. A commonly suggested first step is fractionation of cells to focus on a specific organelle or region (e.g., synaptic cleft) of interest. Taken to an extreme, a tissue can be fractionated so that only a single compound will be isolated via immunohistochemical approaches. However, Subcellular fractionation methods require relatively large amounts of starting material, and it can be difficult to obtain a completely pure subcellular fraction (Pasquali, Fialka, & Huber, 1999). Nevertheless, these techniques could be used to obtain small amounts of a specific organelle needed for study (e.g., mitochondria, synaptosomes) when enough tissue is initially available. Several recent examples highlight the ability to perform proteomic measurements on synaptic fractions (Collins et al., 2006; Burre, Zimmermann, & Volknandt, 2007; Moron et al., 2007). Other specialized capillary-scale separations can be used for high sensitivity characterization of small-volume organelle sampling (Olson, Ahmadzadeh, & Arriaga, 2005). Another approach to decrease sample complexity is sequential extraction. For example, a hypotonic lysis buffer with mild detergent can be used to extract cytoplasmic proteins followed by a buffer that contains strong detergents and chaotropes to extract less soluble (membrane) proteins. Although each step results in some sample loss (Molloy et al., 1998; Herbert, 1999), sequential extraction provides a straightforward approach to fractionate smaller tissue samples (100–250 µg) with less protein loss. We have found that with samples under 50 µg total protein content, wide-ranging single-stage protein extractions appear to provide the best compromise between detectable protein diversity and sample loss.

Given these issues, what are the choices available to separate these protein/peptide mixtures? Most available methods are based on chromatographic or electrophoretic techniques, or some combination of the two. Liquid chromatography (LC) has become one of the most common methods to separate proteins, allowing their identification via MS. With a reverse phase column, peptides, normally in the presence of an ion-pairing reagent, separate based on hydrophobic interactions with the stationary phase. Similarly, ion-exchange columns separate analytes based on charge, and size-exclusion columns based on molecular size. The peak capacity of a separation is an indication of how many different analytes can be separated in a single separation (Giddings, 1991). Normally, the peak capacity (maximum number of theoretical resolution planes) of LC is several hundred, so that an LC separation interfaced to a mass spectrometer can be used to identify several hundred peptides (some degree of peak overlap is acceptable with MS detection). Because many samples have greater complexity, multi-dimensional chromatographic separation methods have been developed. These combined systems have vastly improved peak capacities (Giddings, 1991; Davis & Blumberg, 2005) and allow much more complex samples to be studied; thus, they are well suited to proteomic experiments. In these methods, the mixture is separated in stages, with subsequent separations based on differing properties. For example, the analytes can be separated by charge, pH, hydrophobicity, size, and ability to bind to specific stationary phases, to name a few. The number of different combinations of techniques that have been used is quite large (Evans & Jorgenson, 2004). One specific method of analysis that involves several stages of separation followed by MS analysis has been termed “MudPIT” (multidimensional peptide identification technology) (Washburn, Wolters, & Yates, 2001). MudPIT involves the combination of two chromatographic approaches (strong cation exchange and reverse phase) directly coupled to a mass spectrometer. Protein samples are first digested into component peptides, applied to the chromatographic system, and then eluted in cycles before being introduced to the second stage. Each cycle involves eluting peptides from the ion exchange (SCX resin) with a specific salt concentration, followed by a solvent gradient of increasing hydrophobicity to separate peptides on the reverse phase column; every subsequent cycle begins with an increased salt concentration. Often, the separation by cation exchange resin is performed offline, and the fractions are individually processed through reverse-phase liquid chromatography coupled to the mass spectrometer. Although this offline method is time consuming, there is an increase in the number of peptides identified compared to the direct SCX-reverse phase-high performance LC; for example, more than 5000 peptides were characterized in a single experiment by Li et al. (2005b). In general, such methods require large amounts of starting material as there tend to be protein losses with each fractionation step. Hence, MudPIT is not currently useful for the analysis of smaller samples.
Electrophoretic separations for proteins are among the older separation modes (Tiselius, 1937), and are often used to simplify complex mixtures. Electrophoretic techniques have usually been associated with the use of a gel matrix to assist with the separation (Andrews, 1989). Simple one-dimensional separations by molecular weight can be used, as well as 2DE, as pioneered by O’Farrell and Klose in the 1970s (Klose, 1975; O’Farrell, 1975). Conventional 2DE first separates proteins by their isoelectric points and second by molecular weights. Under ideal circumstances, electrophoresis can be used to resolve hundreds to thousands of individual proteins. However, scaling laws favor miniaturization, so that small 2DE separations have long been known to have advantages for volume-limited samples and for high separation efficiencies (Ruchel, 1977; Poehling & Neuhoff, 1980). We have had good success with “minigel” formats (first dimension isoelectric focusing strips 7–11 cm wide, and second dimension sodium dodecyl sulfate polyacrylamide gel electrophoresis, or SDS-PAGE gels) for microproteomic applications.

The multiple steps in the 2DE process can lead to protein losses (Zhou et al., 2005). Although 2DE has had a reputation for being slow, difficult, and poorly reproducible, these issues have been largely resolved with the development of immobilized pH gradient strips that provide more consistent separations in the first dimension, as well as electrophoretic devices capable of running multiple gels quickly and consistently (Gorg et al., 2000). A renewed interest in radiolabeling strategies (Vuong et al., 2000) and the development of more sensitive fluorescent protein stains (Patton, 2002) have markedly improved detection sensitivity. There have been efforts to directly analyze samples from gel-based separations or through an intermediate transfer to a polyvinylidene fluoride membrane using MALDI time-of-flight MS; however, this approach has yet to be widely applied (Gusev, 2000).

Capillary electrophoresis (CE) is a method that combines several features of column chromatography and electrophoresis. In this method, analytes are introduced into a buffer (conducting electrolyte) and injected onto an open capillary. When a potential is applied, the analytes migrate along the capillary with their velocity depending on their electrophoretic mobility and electroosmotic (buffer) flow (Jorgenson & Lukacs, 1983). During electrophoretic separations one applies a current across a solution, which generates heat. This heat tends to disturb the separation process. In CE, heat dissipation is more effective as the capillary diameter is reduced, so that smaller capillaries can be used with higher voltages to improve and speed up separations. Like LC, CE can be directly interfaced with mass spectrometers. However, combining CE with an MS instrument presents several challenges, because the mass of the material in a nanoliter-volume (or smaller) analyte band is difficult to characterize with MS. Reviews of how these challenges have been overcome include Simpson and Smith (2005), Banks (1997), and others (Oehnseorge, Neususs, & Watzig, 2005).

There are several operational modes of separation by CE that have been interfaced with a mass spectrometer, notably, capillary zone electrophoresis (CZE) and capillary isoelectrofocusing (CIEF). CZE is similar to LC in that proteins are injected onto the column, and the components are separated into discrete bands that elute from the capillary and are subsequently introduced into the mass spectrometer. CIEF is similar to the gel-based approach because the separation is based on the pI (Shen & Smith, 2002). A direct comparison of LC, CZE, and CIEF for analyses of yeast cytosol digest showed that the CE methods have a high resolving power, separation efficiency, and shorter run times compared to LC (Shen et al., 2000). However, CIEF allows more material to be introduced into the capillary (Cooper, Wang, & Lee, 2004). Smaller sample volumes suggest that CE would be an ideal tool for microproteomics. However, these small sample bands mean that a smaller dynamic range of proteins can be detected. Taken together, these characteristics suggest that CE separation would be well-suited for the last stage of a comprehensive multistage separation, or as a sole technique for the smallest samples.

V. IDENTIFICATION OF PROTEINS USING MASS SPECTROMETRY (MS)

As mentioned earlier, one challenge in sequencing proteins arises from the complexities inherent in having 20 amino acid building blocks and a variety of PTMs. This complexity is even greater because protein translation levels differ both temporally and spatially. Gel- and immunohistochemical-based methods to identify proteins have been used successfully in a wide variety of samples. Edman degradation has also been a powerful approach. However, the throughput of these methods is low. Although MS technologies have been utilized for decades, it was only with the development of ionization methods that allow large protein molecules to be introduced into mass analyzers has MS been widely applied to proteomics. Specifically, ESI and MALDI revolutionized protein characterization (Tanaka et al., 1988; Hillenkamp et al., 1991; Karas, Bahr, & Gießmann, 1991). In MS-based methods, much smaller sample amounts are required to sequence a peptide. Amounts can be in the attomole or even zeptomole range under ideal circumstances.

All mass analyzers require the ionization of a protein molecule in the gas phase; therefore, inefficiencies in the vaporization and ionization processes can often limit overall system performance. Because it is easier to ionize and characterize peptides, proteins are often cleaved into complex peptide mixtures prior to measurements. In this so-called “bottom-up” approach, a protein is digested into its component peptides with enzymes such as trypsin or chymotrypsin, and the peptides characterized with MS. Once the peptide sequences are confirmed, they are matched to precursor proteins using various databases, in an approach called peptide mass fingerprinting (PMF) (James et al., 1993). This “bottom up” approach has been widely used.

Global proteomics by means of PMF normally requires the detection of multiple peptides to unambiguously identify the protein; measuring the peptides with sufficiently high mass accuracy can permit proteins to be identified based on fewer peptides. This method also uses accurate mass information and the retention times from a list of previously identified peptides to decrease the sequencing time spent on those peptides by the mass spectrometer. In a report that demonstrated the utility of this approach to proteomics, 61% of the predicted peptides in the Deinococcus radiodurans proteome were identified (Lipton et al., 2002).
A newer protein identification strategy is the “top-down” approach, which uses extremely accurate masses of intact proteins obtained from a Fourier transform mass spectrometer as well as masses obtained from their subsequent CID fragmentation to identify proteins (Kelleher, 2004; Han et al., 2006). A major advantage of the “top-down” approach is its robustness for analyzing modified proteins (Pesavento et al., 2004). The “bottom-up” approach will miss PTMs located on peptides that are not identified via MS.

VI. INTEGRATING TISSUE SAMPLING AND CHARACTERIZATION IN A SINGLE STEP: MASS SPECTROMETRIC IMAGING (MSI)

Usually, scientists utilize immunohistochemical techniques to determine protein localization in a tissue, and MS to identify proteins of interest. However, over the last 30 years several forms of probe-based MS have been adapted to protein localization studies. Recently, Caprioli and others have pioneered the adaptation of MALDI MS to image proteins in their anatomical context in tissue samples (Spengler, Hubert, & Kaufmann, 1994; Stoeckli, Farmer, & Caprioli, 1999; Stoeckli et al., 2001; Todd et al., 2001; Altelaa et al., 2005; Caldwell & Caprioli, 2005; Crecelius et al., 2005; Jurchen, Rubakhin, & Sweedler, 2005; McDonnell et al., 2005; Rubakhin et al., 2005; Monroe et al., 2006). In MALDI MSI, a tissue section is prepared and matrix is applied. The laser is focused onto a discrete point, and a mass spectrum acquired. At each point, the amount of tissue sampled is in the nanoliter-volume range; thus, this approach is inherently microproteomic. The laser is rastered across the sample to obtain spatially resolved mass spectra, creating a “spectral image” of the tissue sample. Because imaging requires the acquisition of thousands of spectra, MSI is heavily dependent on instrumental and data-storage capabilities. One advantage of this approach is that because minimal sample handling is required, protein losses are reduced. However, the depth of proteomic coverage is less than when conventional separations are performed. Also, the procedures used to collect and prepare samples for MSI are critical, especially when analyzing phosphoproteins or neuro peptides, both of which can be quickly modified or degraded if sample preparation is suboptimal. New, innovative ionization methods such as desorption electrospray ionization (DESI) are also being applied to MSI. In contrast to MALDI, DESI permits images to be created from tissue slices under ambient conditions (Cook et al., 2006). This method performs well, but its upper mass range must be extended before its application to biomedical proteomics is practical. Because MSI is a rapidly changing field, readers are encouraged to examine recent publications devoted to this technology, such as (Heeren & Sweedler, 2007).

VII. QUANTIFICATION OF PROTEIN EXPRESSION DIFFERENCES

Often, some of the most important information gained from a proteomic study is the determination of differences in protein expression between experimental conditions. Although one frequently uses the term quantification to indicate the absolute amount of a material (in concentration or mass), in most proteomic measurements, the term implies relative abundance, such as the percent change between two treatment groups. Quantification is usually done after the sample is fractionated, but can be performed either before or after protein identification. The advantage of quantifying before identification is that efforts can then be focused on identifying only those proteins that are differentially expressed. Quantification can follow either a gel-based separation, by comparing differences in the intensity of corresponding spots, or after MS measurement, by evaluating differences in the intensities/areas of corresponding mass spectral peaks. Because of the expense and effort required for many proteomic measurements, care must be taken to ensure that the experimental design balances the running of samples from the different groups to avoid any systematic bias. There are numerous case studies that illustrate the dangers of design flaws that can lead to systematic bias in proteomic studies (e.g., see Baggerly et al., 2004a; Baggerly, Morris, & Coombes, 2004b; Baggerly, Coombes, & Morris, 2005a; Baggerly et al., 2005b; Hu et al., 2005a). This problem can be avoided by applying standard experimental design principles such as blocking and randomization, as described in standard textbooks (e.g., Box, Hunter, & Hunter, 2005), within the laboratory setting.

How is quantification performed? Gel-based methods usually involve fluorescent and some visible stains, as well as radiolabeling. The relationship between spot intensity and protein amount is linear for these stains over several orders of magnitude. Silver stains do not have as linear a relationship, but are applied less frequently for quantification. DIGE is a method that can reduce variability by running control and treatment samples together on the same gel, as well as a pooled sample from the two groups. Dyes with different spectral characteristics are used to detect and quantify spots from the different groups. The most common set of dyes are the Cy dyes (Friedman, 2006). Although these approaches are well validated, areas that can be improved when working with the smallest samples include the use of more efficient (saturation) protein labeling, use of more sensitive dyes, and reducing the effect that dyes have on changing protein migration.

After scanning the gels for a given experiment, a key quantitative challenge is to detect spots on the gels while filtering out artifacts, and to match the spots across gels and quantify them. Assuming N gels and p protein spots in the set, the goal of this step is to construct an N-by-p matrix that contains the protein expression values for each spot for each gel, which can be surveyed to identify those proteins related to the factor of interest. Various commercial and noncommercial methods have been developed to accomplish this goal, but unfortunately, these methods encounter numerous difficulties that limit their effectiveness, especially for larger studies. Various errors, such as spot detection, spot matching, and spot boundary estimation errors, are typical and reduce the probability of finding protein expression differences in the data. These problems tend to accumulate with larger studies, and encourage some investigators to perform smaller studies that are underpowered to find realistic-sized differences (Fig. 3).

In response to these problems, we have developed an approach for preprocessing 2DE data called Pinnacle (Morris,
Clark, & Gutstein, 2007). The name derives from the fact that, unlike most existing methods, this method performs spot quantification using pixel intensities at peak spot intensity values in the horizontal and vertical dimensions (pinnacles) rather than spot volumes. The underlying philosophy is to keep the preprocessing as simple as possible in order to avoid any extra bias and variance that can result from the propagation of errors that characterize more complex methods. The steps of the Pinnacle method are as follows:

1. Align the 2DE gel images.
2. Compute the average gel, averaging the staining intensities across gels for each pixel in the image.
3. Denoise the average gel using the two-dimensional undecimated discrete wavelet transform (UDWT).
4. Detect pinnacles on the denoised average gel, which are pixel locations that are local maxima in the horizontal and vertical directions with an intensity above some minimum threshold (e.g., the 75th percentile for the average gel).
5. Perform spot quantification on the individual gels by taking the maximum intensity within a stated tolerance of each pinnacle location.

Assuming \( p \) pinnacles are detected on the average gel, one obtains an \( N \times p \) matrix of spot intensities that can be surveyed for potential biomarkers.

This method is significantly quicker and simpler than the typical spot detection algorithms used in 2DE, which perform spot detection on individual gels with complex spot definitions, quantifying spot volumes, and matching spots across gels. This usual approach is time-consuming and error-prone, and is especially problematic for larger studies, because these errors propagate as the number of gels increases. It takes a great deal of time-consuming hand-editing to try to fix these errors. Further, the process of estimating spot boundaries is also difficult and error-prone, and leads to increased coefficients of variance when spot volumes are used to quantify the proteins. Simulation studies have demonstrated that the Pinnacle method leads to more reliable and precise quantifications (Morris, Clark, & Gutstein, 2008). Statistical principles suggest that performing spot detection on the average gel leads to greater sensitivity and specificity, because true protein spots are reinforced across gels whereas artifacts and noise average out. As a result, this method has the potential to reliably detect and quantify fainter spots, thus increasing the realized dynamic range of the technology (Fig. 4).

VIII. MASS SPECTROMETRIC-BASED QUANTIFICATION

The ability to detect a protein via MS depends on multiple variables, including vaporization/desorption and ionization efficiency (which depend on sample constituents besides proteins), and losses during separation and sample handling. Thus, it is difficult to directly compare peaks between treatment groups unless these variables are constant. Although a direct correlation between peak intensities from two samples is challenging, evolving methods use parallel MS experiments for a direct comparison of the peptide peaks (Wiener et al., 2004). Perhaps the most common method for quantification involves tagging the proteins with labels that differ only by stable isotopes (Fricker et al., 2006). Using such methods, proteins in two or more proteome samples are tagged with different isotopic tags that vary in their molecular weight but are chemically nearly identical; thus, any sample or ionization differences between the two can be corrected, and their peak intensities directly correlated to their concentrations. Several different chemical tags using H/\(^{2}\)H, \(^{12}\)C/\(^{13}\)C, \(^{16}\)O/\(^{18}\)O, or \(^{14}\)N/\(^{15}\)N isotopes have been reported (Leitner & Lindner, 2004; Righetti et al., 2004).

One can avoid the requirement for derivatizing peptides/proteins with the isotopic label by using in situ labeling. For

![FIGURE 3. Comparison of yeast cytosol tryptic digest (at 5 mg/mL) using three different CE separation modes. A: Capillary isoelectric focusing (CIEF); B: capillary zone electrophoresis (CZE); and C: reverse phase LC. The concentration used for CIEF was 16-fold lower than the samples used for CZE and LC. Reprinted with permission from Shen et al. (2000), copyright 2000 American Chemical Society.](image-url)
example, isotopic labeling of proteins in cell cultures is achieved by using stable isotope-enriched amino acids in the culture media (known as SILAC) (Ong, Foster, & Mann, 2003). This method is used for cell cultures under different conditions and allows the quantification of the expression levels of hundreds of proteins, thus promising insights into cell signaling pathways and also several interconnected signaling pathways.

Methods similar to Pinnacle discussed above have also been developed to detect and quantify peaks in one-dimensional MS data, such as MALDI MS and surface-enhanced laser desorption-ionization (SELDI) MS (Coombes et al., 2005; Morris et al., 2005). Stand-alone software is freely available for applying the method to MS analysis (Karpievitch et al., 2007). The steps are as follows:

1. Align the spectra on the time scale by choosing a linear change of variables for each spectrum in order to maximize the correlation between pairs. We have found that alignment can be done much more simply and efficiently on the time scale rather than the m/z scale.

2. Compute the mean of the aligned raw spectra.

3. Denoise the mean spectrum using the UDWT.

4. Locate intervals that contain peaks by finding local maxima and minima in the denoised mean spectrum.

5. Quantify peaks in individual raw spectra by recording the maximum and minimum height in each interval, which should contain a peak. This quantification method implicitly removes the baseline artifact.

6. Calibrate all spectra using the mean of the full set of calibration experiments.

Assuming that a total of $p$ peaks are detected on the average gel, an $N$-by-$p$ matrix of peak intensities is obtained for the $N$ spectra in the study that can be surveyed for potential biomarkers.

A key component of these approaches as described for two- and one-dimensional data is that we perform peak (or spot) detection on the average spectrum (gel), rather than on individual spectra (gel); a number of advantages are obtained (Morris et al., 2005). First, it avoids the difficult and error-prone peak-matching
step that is necessary when using individual spectra. Eliminating peak-matching improves the accuracy of the data, and results in no missing data. Second, it tends to result in greater sensitivity and specificity for peak detection, because averaging across \( N \) spectra reinforces the true signal while weakening the noise by a factor of \( \sqrt{N} \). Averaging enables us to better detect real peaks down near the noise region of the spectra, and decreases the chance of flagging spurious peaks that are noise. The simulation study presented in Morris et al. (2005) demonstrates that using the average spectrum results in improved peak detection, with the largest improvement being for low-abundance peaks of high prevalence. Third, it speeds preprocessing time considerably, because peak matching is by far the most time-consuming preprocessing step.

IX. DETERMINATION OF STATISTICAL SIGNIFICANCE

Given an \( N \times p \) matrix that contains relative quantifications for \( p \) features (peak/spot quantifications) for each of \( N \) spectra/gels, the next analysis step is to identify which features are significantly associated with the factor of interest, and that might indicate potential biomarkers. Significance should be determined with appropriate statistical tests, not simply ad hoc criteria such as fold-change, because statistical tests appropriately account for the variability of the measurements when making the determinations, whereas simple use of fold-change does not. The particular test to use depends on the number of groups and the experimental design of the study. For example, if the goal is to find proteins that are differentially expressed between two experimental conditions, then a \( t \)-test (or nonparametric rank-sum test if normality cannot be assumed) can be done for each \( p \) feature, and then all features with sufficiently small \( P \) values flagged as significant.

In this setting, the selection of this \( P \)-value cutoff must be done with care. Because these assays survey hundreds or even thousands of proteins simultaneously, we expect a certain number of them to have small \( P \) values even if none are truly related to the factor of interest. In the absence of true differences, we expect ca. 50 \( P \) values of \(<0.05 \) for every 1,000 protein spots. In statistics, this is called the multiplicity or multiple testing problem. One classical approach to deal with multiplicities is Bonferroni, which would use a cutoff of 0.05/\( p \) to determining significance, where \( p \) is the number of features considered. This cutoff would control the experiment-wise error rate at 0.05, and mean that we expect that the probability of at least one false positive result is 0.05.

Because this criterion strongly controls the false positive rate, but results in a large number of false negatives, it is widely considered too conservative for exploratory analyses like these. Other methods control the false discovery rate, or FDR (Benjamini & Hochberg, 1995). Controlling the FDR at 0.05 means that if the features declared significant, we expect 5% or fewer of them to be false positives. There are a number of procedures for controlling FDR (Benjamini & Hochberg, 1995; Benjamini & Liu, 1999; Yekutieli & Benjamini, 1999; Genovese & Wasserman, 2002; Storey, 2002; Ishwaran & Rao, 2003; Pounds & Morris, 2003; Storey, 2003; Efron, 2004; Newton et al., 2004; Pounds & Cheng, 2004; Datta & Datta, 2005). Many of these methods take advantage of the property that the \( P \) values that correspond to features that are not associated with the factor of interest should follow a uniform distribution, whereas the distribution of \( P \) values for predictive features should be characterized by an overabundance of small \( P \) values. In most of these methods, one inputs the list of \( P \) values and desired FDR, and receives as output a \( P \)-value cutoff that preserves the desired FDR. Typically, this cutoff is smaller than 0.05, but quite a bit larger than the corresponding Bonferroni bound 0.05/\( p \). Using the FDR of 0.05, we still expect that one out of every 20 flagged proteins are false positives. We must be willing to accept some false positives so that we have sufficient power to capture relevant markers. Thus, it is important to validate any flagged markers (e.g., by Western blotting or multiple reaction monitoring MS).

Instead of performing peak/spot detection and applying statistical tests to each feature, another alternative is to model the spectra or gel images using functional data techniques. One flexible method that has been developed for this purpose is the wavelet-based functional mixed model (Morris, Walla, & Gutstein, 2008). This approach does not depend on peak or spot detection at all, and can search for differentially expressed regions of the spectra or gel images in a way that takes statistical considerations and fold-change into account, and accounts for the multiplicity problem. These methods are more complex and computationally intensive, and just under development, but show considerable promise for the analysis of proteomic data.

Power calculations can be done to assess how large a sample size (number of gels, number of subjects) is needed for a particular proteomics study. In recent years, various methods have been developed to perform sample size calculations for settings where FDR is used to deal with the multiplicity problem (Hu, Zou, & Wright, 2005b; Jung, 2005; Li et al., 2005a; Pounds & Cheng, 2005; Liu & Hwang, 2007). In each case, the calculations depend on the reproducibility and assumed subject-to-subject variability in the data, as well as the magnitude of the effect sizes for the differentially expressed proteins. If preliminary studies in the tissue of interest are available, then these data can be used to estimate the reproducibility (variance across replicate gels) and subject-to-subject variability (variance across gels for different subjects), and the effect size (difference between group means on the log scale) for the set of spots detected in the preliminary data. Absent these preliminary studies, it is difficult to perform a sample size/power calculation.

The use of small biological samples in proteomic analyses is likely to affect the statistical properties of these studies. As we operate closer to the detection limit of the available technology, it is likely that technical variability will increase, potentially reducing the statistical power of the research. This effect can be mitigated by performing larger studies with more replicates and/or subjects. Another possibility is to pool samples. By increasing the amount of total protein per assay, this pooling can help mitigate the effects of analyzing samples near the detection limit, but at the cost of losing protein information for individual samples. Pooling may be an acceptable tradeoff in studies attempt to find proteins differentially expressed across groups, but this pooling prevents the possibility of finding proteins correlated with the subjects’ individual outcomes.
X. CONCLUSIONS

Proteomics is a rapidly evolving field. Technological advances permit us to examine a greater fraction of the proteome from less tissue. Statistical and analytical advances also permit us to determine significant changes in protein expression and PTMs from smaller samples with greater certainty and specificity.

However, more progress must be made. For example, improvements in protein extraction methods as well as the specificity of protein identification techniques to allow us to evaluate lower-concentration proteins, or a greater fraction of the membrane proteins are certainly needed, as are better methods to characterize more complex PTMs such as carbohydrates. Of course, such advances should also be coupled to greater sensitivity so that smaller samples can be assayed.

What options await us on the horizon? Methods such as MSI or DESI that bypass much of the conventional proteomics workflow are intriguing, because they minimize the number of steps needed to identify proteins from tissue. These systems could minimize obligate protein losses from extraction and separation techniques, and so perhaps will provide more efficient protein identification from small groups of cells. Nanotechnology, another rapidly developing area, is poised to make significant contributions to microproteomics as novel miniaturized platforms are applied to protein extraction, separation, and identification. Further evaluation and development of these and other promising and exciting methods are needed to allow researchers to take advantage of the information offered by proteomics. As we move toward an ultimate goal of examining the complete proteome from single cells or small cell groups, the keys to achieving this end will be creativity, flexibility, and a willingness to take risks and innovate past the ultimate limitations of conventional approaches.

ACKNOWLEDGMENTS

The support of the National Institute on Drug Abuse to the UIUC Neuroproteomics Center on Cell to Cell Signaling through Award No. DA018310, is gratefully acknowledged. Work in the authors’ laboratories is also supported by additional NIH grants from NIDA, NIAAA, and NGMS. We thank Marlen Banda for manuscript preparation assistance.

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