Advances in the proteomic discovery of novel therapeutic targets in cancer

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Abstract: Proteomic approaches are continuing to make headways in cancer research by helping to elucidate complex signaling networks that underlie tumorigenesis and disease progression. This review describes recent advances made in the proteomic discovery of drug targets for therapeutic development. A variety of technical and methodological advances are overviewed with a critical assessment of challenges and potentials. A number of potential drug targets, such as baculoviral inhibitor of apoptosis protein repeat-containing protein 6, macrophage inhibitory cytokine 1, phosphoglycerate mutase 1, prohibitin 1, fascin, and pyruvate kinase isozyme 2 were identified in the proteomic analysis of drug-resistant cancer cells, drug action, and differential disease state tissues. Future directions for proteomics-based target identification and validation to be more translation efficient are also discussed.

Keywords: proteomics, cancer, therapeutic target, signaling network, tumorigenesis

Introduction

The use of proteomics as a systems biology tool in cancer research continues to expand in scope and depth, as it evolves rapidly into a universally applicable method for the investigation of practically any biological process. Proteomics is particularly attractive to cancer research because the complexity of tumorigenesis, cancer progression, tumor relapse, and metastasis often involves large protein networks. Indeed, in a little over a decade, applications of mass spectrometry-based proteomics in cancer-related research – ranging from mechanistic investigation to the discovery of novel therapeutic targets – have increased exponentially. This is evidenced by the explosive growth in the number of publications on the subject matter, from 109 in 2000 to 1,349 in 2011 (http://www.ncbi.nlm.nih.gov/pubmed), with the total number of publications containing the keywords “proteomics” and “cancer” reaching over 7,500 to date. A large number of review articles have appeared in the past several years, offering excellent overviews and perspectives on novel proteomic applications in cancer. Many reviews focused on different cancer types, such as breast cancer,1–4 pancreatic cancer,5,6 ovarian cancer,7–9 colorectal cancer,10,11 and glioma.12–14 Others have focused on sample types or subcellular components, such as tissue,15–17 serum,18–20 and secretome.21–23 In addition, other reviews focused on the innovative proteomic methodologies or the various end goals of proteomics in cancer, such as phosphoproteomics,24–27 mechanisms of drug resistance,28–30 and kinome profiling.31–33 Finally, there appear to be more reviews on proteomic biomarker discovery for cancer diagnosis, prognosis, and therapeutic response than on any other area of focus, highlighting the intense activities in the search for new biomarkers that could be translated to clinical applications.30,34–36
Despite the remarkable advances in utilizing proteomics in almost every aspect of cancer research, many challenges remain. Global proteomic approaches, while capable of identifying thousands of proteins (many of which are differentially expressed) are often biased towards soluble and high-abundance proteins, thus easily missing out on many low-abundance proteins that could play critical roles in signal transduction. Most solid tumors are heterogeneous, making it difficult to identify key proteins that could be used as universal targets even in the same type of cancer. While high-throughput proteomic methods can discover “unbiased” cohorts of potential biomarkers and therapeutic targets that regulate disease progression, the sheer number (often in the hundreds, if not the thousands) of hits could be cumbersome, as current functional studies are limited to involving one or a few proteins at a time. Thus, a large number of diagnostic and prognostic biomarkers have been identified by proteomic analysis, but very few have made it to the clinical stage, underscoring the urgent need for the convergence of functional studies and clinical sample validations. Moreover, the progress in drug development has far lagged the pace of proteomic discovery of potential drug targets, as the former generally requires a prolonged developmental process from drug design to in vitro and in vivo tests for efficacy and toxicities, as well as for absorption, distribution, metabolism, and excretion (ADME).

Proteomic approaches have been increasingly used in the preclinical drug discovery process to understand the effects of drug candidates on their protein targets and to shed light on the cellular mechanisms behind the observed phenotype, although many other biomaterials including deoxyribonucleic acid (DNA), ribonucleic acid (RNA), and carbohydrates can also be used as novel drug targets with different approaches. Proteomic studies with the main goal of identifying new druggable targets for the treatment of cancer have been relatively few and are the primary focus of this review. For the convenience of discussion, we will attempt to group various proteomic target discovery studies into several distinct approaches for target identification.

**Current proteomic techniques for target discovery**

While mass spectrometry (MS)-based proteomic approaches remain the most comprehensive and versatile tool in large-scale proteomic profiling, several non-MS-based techniques, such as reverse phase protein microarrays (RPMA) and peptide arrays have recently gained popularity. RPMA is a technology platform designed for the quantitative analysis of specific phosphorylated, cleaved, or total (phosphorylated and nonphosphorylated) forms of cellular proteins from complex mixtures in multiple samples simultaneously. One microarray can accommodate a range of hundreds to thousands of samples that are printed in a series of replication. RPMA is performed using either a primary or a secondary labeled antibody by chemiluminescent, fluorescent, or colorimetric assays. Multiplexing is also achieved by simultaneously probing multiple arrays spotted with the same lysate with different antibodies and can be implemented as a quantitative calibrated assay. RPMA has now been utilized for potential drug discovery/validation, as well as for advancements in understanding the disease both in the clinic and in the laboratory, and RPMA is currently being integrated into human clinical cancer trials.

The quantitative methods adopted in MS include techniques based on the stable isotope labeling of proteins/peptides, as well as label-free methods. In comparison with labeling methods, label-free methods provide higher dynamic ranges of quantification and are versatile tools that are used to estimate changes in protein abundances between different samples. The utility of MS-based proteomic analyses and their applications in drug target identification have been increasingly recognized over the past decade due to their high sensitivity, specificity, and throughput. For example, gel-free isotopic labeling methods, such as SILAC (stable isotope labeling with amino acids in cell culture), iTRAQ, and isobaric tandem mass tags (TMTs) (Figure 1) can now routinely quantify several hundreds of (to a few thousand) proteins in a single analysis with high reproducibility. Continued innovations and improvements in the instrumentation and bioinformatics tools will drive MS-based proteomics applications in drug discovery, complemented by streamlined and focused non-MS-based methods for targeted and repeated assays, such as protein arrays. The former will play key roles in the discovery stage, where quantitative proteomic analysis is undertaken, which compares differentially expressed proteins for potential drug targets. The array methods will be more cost effective and convenient to use for the validation of studies on drug targets and their effects.

**Target identification in resistant cancer population**

One of the greatest challenges in cancer therapy is drug resistance that occurs either intrinsically or is acquired after a certain period of treatment. Resistance to targeted cancer drugs involves complex and diverse molecular adaptations...
by cancer cells under the selective pressure of therapeutic regimens. For example, the specific target protein in tumors may undergo mutations to become inaccessible to drugs.\textsuperscript{62–64} In addition, a subset of cancer stem cells may be resistant to the cytotoxicity of drugs that target the bulk of more differentiated cancer cells.\textsuperscript{65–68} Moreover, chemotherapeutic agents often induce adaptive changes in the regulatory networks and signaling pathways, leading to the independence of cancer cells on the targeted protein and to the emergence of drug-resistant disease clones.\textsuperscript{69} Indeed, such diverse mechanisms of drug resistance can transform cancer cells under the selective pressure of therapies, which made it extremely difficult to combat tumor recurrence; this is inevitably followed by metastatic progression of the disease and death. Current cancer treatment typically consists of a monotherapy regimen targeting a specific protein or signaling pathway, (for example, selective estrogen receptor modulators [SERMs] for the estrogen receptor, Herceptin for Her2, and gefitinib for epidermal growth factor receptor [EGFR]). However, tumors that are initially responsive eventually develop resistance to therapy via a variety of mechanisms. Switching to alternative drugs after the emergence of resistant disease clones following first-line treatment often achieves little in delaying cancer progression. Resistance can develop to just one drug, or tumors may be cross-resistant to several structurally dissimilar and functionally distinct agents, a phenomenon known as multidrug resistance.\textsuperscript{70} Thus, there is an urgent need to better understand the resistance mechanism and to identify novel targets for a therapeutic intervention that can be more effective in blocking the progression of cancer cells already resistant to the previous treatment.

Here, proteomic approaches hold the promise to identify individual proteins and interactive signaling networks that act individually or collectively in conferring drug resistance. The comparison of protein expression between drug-resistant tumor tissues or cell lines with drug-sensitive tumor tissues or cell lines is one of the most promising tools for drug target identification. Targeting these proteins and signaling pathways may either resensitize tumor response to the original treatment, or it may help overcoming acquired drug resistance by blocking alternative survival signaling induced by the initial treatment. More importantly, proteomic characterization of a resistant phenotype can provide a more complete picture of signaling adaptation, as multiple drug targets for combinatorial therapeutic solutions can be identified to combat resistance and achieve longer-term improvement in disease outcomes. While advances in this application have been limited, the following examples demonstrate that proteomic
approaches are inherently suitable for the discovery of targets for resistant phenotypes, and we expect an increasing number of protein targets to be validated by functional studies followed by therapeutic development.

One example comes from a comparative proteomic analysis of colon cancer stem cells and differentiated tumor cells, which uncovered a prominently upregulated protein, baculoviral inhibitor of apoptosis protein (IAP) repeat-containing protein 6 (BIRC6), one of the IAPs that may play a crucial role in the chemoresistance of colon cancer stem cells. Subsequent knockdown of the gene resulted in resensitization of the cells to chemotherapy. Results from this proteomic study suggest that BIRC6 could be used as a potential therapeutic target to eradicate colon cancer stem cells contributing to colon cancer recurrence. In other cases, an iTRAQ–MS-based proteomics analysis was used to identify proteins associated with the development of docetaxel resistance by comparing docetaxel-sensitive PC3 cells and docetaxel-resistant PC3-Rx cells developed by docetaxel dose escalation. Functional validation experiments were performed using recombinant protein treatment and small interfering (si)RNA knockdown experiments. The findings from this study suggested that macrophage inhibitory cytokine 1 should be further investigated as a potential biomarker and therapeutic target for acquired resistance to docetaxel treatment. However, extensive functional studies and follow-up validations are required before these proteins can be proven as viable drug targets.

Similarly, a quantitative proteomic investigation of hepatocellular carcinoma, a type of liver cancer known for its resistance to chemotherapy, implicated phosphoglycerate mutase 1 (PGAM1) as a potential therapeutic target. Clinicopathological analysis indicated that the overexpression of PGAM1 was associated with 66.7% of hepatocellular carcinomas, and is strongly correlated with poor differentiation and decreased survival rates ($P<0.01$). In addition, short hairpin (sh)RNA-mediated repression of PGAM1 expression resulted in significant inhibition in liver cancer cell growth both in vitro and in vivo. Using a proteomics approach, elevated levels of prohibitin 1 (PHB1) and glutathione S-transferase pi (GSTπ) were found to be associated with paclitaxel resistance in discrete subcellular fractions of two drug-resistant sublines relative to their sensitive sublines. Immunoﬂuorescent staining and fractionation studies revealed an increased level of PHB1 on the surface of resistant cell lines. Transiently silencing either PHB1 or GSTπ gene expression using siRNA in the paclitaxel-resistant cancer cell sublines partially sensitized these cells toward paclitaxel. Intriguingly, silencing PHB1, but not GSTπ, resulted in the activation of intrinsic apoptosis pathway in response to paclitaxel. Similarly, stably silencing either PHB1 or GSTπ significantly improved paclitaxel sensitivity in A549TR cells both in vitro and in vivo. This study suggests that PHB1 is a mediator of paclitaxel resistance, and this resistance may depend on the cellular localization of proteins. Based on these preliminary functional studies, PHB1 is proposed as a potential target for therapeutic strategies for the treatment of drug-resistant tumors. Again, we have yet to see follow-up investigations of the feasibility of such novel, yet unproven, therapeutic targets.

In our recent proteomic study of letrozole-resistant breast cancer, a TMT label was used for a quantitative comparison of protein expression profiles of the resistant versus sensitive MCF-7 cells overexpressing aromatase. This study identified fascin, among other significantly upregulated proteins, as a promising therapeutic target for the inhibition of metastasis of hormone-resistant breast cancer cells that have acquired the enhanced capacity of migration and invasion. Retrospective clinical validation confirms fascin overexpression as an independent indicator of decreased survival and poor prognosis. A subsequent drug discovery effort by our group led to the discovery of a series of thiazole compounds as potent inhibitors of migration and invasion of metastatic cancer cells by binding to fascin.

In summary, a resistant cancer population, whether inherent or acquired, is a major problem that reduces the activity of conventional and/or molecularly-targeted cancer drugs. The success of identifying drug targets in resistant cancer cells by proteomic approaches depends on reliable drug-resistant tumor tissues or cell lines. It is also critical that proteomics be combined with data from other approaches in an attempt to investigate mechanistic pathways in more detail, as well as to validate potential targets in a clinical setting.

**Target identification from drug action (chemical proteomics)**

Chemical proteomics is a technique that identifies proteins enriched or isolated as a result of interacting with or binding to a chemical probe, usually a small molecule drug that is fixed on a solid support. Therefore, these affinity-based enrichment techniques, in combination with MS, have enabled the direct determination of protein-binding profiles of small molecule drugs under physiological conditions and represent one of the most direct approaches to screen for drug–protein interactions. The major drawback encountered in the affinity-based chemical proteomics is the
presence in the pulled-down extract of nonspecifically bound proteins. Among these, a large number of proteins that bind nonspecifically to most conventional affinity matrices have been reported. It is, therefore, strongly recommended that green fluorescent protein be used as the tag of choice because it shows minimal nonspecific binding to mammalian cell proteins, it can be quantitatively depleted from cell extracts, and it allows for the integration of biochemical protein interaction data with in vivo measurements using fluorescence microscopy. In addition, it is also necessary to perform appropriate negative control experiments to distinguish nonspecific interactions from specific interactions. Chemical proteomic approaches also include global proteomic profiling of cellular samples treated with a biologically active compound without enrichment steps. In pharmacology, chemical proteomics have been utilized to determine the specificity of drugs and their analogs, for anticipated as well as unknown targets that may also bind to the probe. These activity-based probes can specifically target diverse sets of enzyme families and provides direct information about the activation state of identified proteins. Several kinds of chemical probes have been used in proteomic studies across a multitude of enzyme classes such as hydrolases, proteases, kinases, phosphatases, histone deacetylases, glycosidases, and oxidoreductases. Thus, chemical proteomic approaches not only identify protein targets for drugs that exert known biological activities in vitro or in vivo, but they also can discover previously unknown targets for drugs of known modes of action (MoA). These off-target proteins may be additional druggable targets, but they may also account for the side effects and toxicities. Identification of protein targets that are responsible for toxicities may prove valuable in early drug development to minimize failure in clinical trials. In addition, chemical proteomics can identify novel drug targets by broad-action chemical probes (for example, nonselective kinase inhibitors such as Staurosporine [Life Technologies, Carlsbad, CA, USA]). In the following, we present a few examples in which chemical proteomics were used effectively for target discovery and validation.

Tyrosine kinase inhibitors represent a large group of targeted therapeutics for cancer. Gefitinib (Iressa™; AstraZeneca, London, UK) is the first selective inhibitor of the EGFR tyrosine kinase domain. Gefitinib used as monotherapy is an effective treatment for patients with locally advanced or metastatic nonsmall cell lung cancer (NSCLC) with EGFR mutations. Gefitinib has also been tested in clinical trials in other tumors, including head and neck squamous cell carcinoma (HNSCC), as monotherapy, or in combination with other chemotherapies or radiation, but it has shown limited clinical efficacy with response rates of 10%–15%. Using cleavable isotope-coded affinity tagging (cICAT)-based liquid chromatography–tandem mass spectrometry (LC–MS/MS) method, Chen et al identified the tyrosine phosphorylation levels of 21 proteins between control and epidermal growth factor-treated A431 human cervical cancer cells. Of these, endofin, DCBLD2, and KIAA0582 were validated to be novel tyrosine-phosphorylation targets of epidermal growth factor signaling and responsive to gefitinib therapy. Pernas et al observed that gefitinib sensitivity correlated with phospho (p)-AKT and p-STAT3 activation in HNSCC cell lines and tumor specimens, thus p-AKT and p-STAT3 could serve as potentially useful drug targets for further development of novel therapeutic agents for HNSCC. More recently, Wu et al used quantitative chemical proteomics to identify several kinases including NEK9, LYN, JAK1, WEE1, and EPFA2, which are involved in cell survival and the proliferation of HNSCC cell lines. These findings may lead to new therapeutic options for HNSCC patients.

For inhibitors acting on multiple tyrosine kinases, chemical proteomic studies could help uncover additional unknown targets and their interacting signaling network. For example, as an inhibitor of multiple tyrosine kinases, dasatinib targets Bcr-Abl, SRC family kinases, c-Kit, and platelet-derived growth factor receptor kinase, a cell-permeable kinase probe (DA-2) was employed to profile potential cellular targets of dasatinib; a number of previously unknown dasatinib targets were identified, including several serine/threonine kinases (PCTK3, STK25, elt-2A, PIM-3, PKA C-α, and PKN2). According to another report, nearly 40 different kinase targets of dasatinib were discovered using quantitative phosphoproteomics. These include receptor tyrosine kinases (ephrin receptors, discoidin domain receptor 1, and EGFR) and nonreceptor tyrosine kinases (FRK, BRK, and ACK). These results provided a system-level view of dasatinib action in cancer cells and suggested functional targets and rationales for therapeutic strategies.

When the mechanisms of biologically active compounds against cancer cells are not fully understood, chemical proteomics could help elucidate the modes of action by mapping out protein networks perturbed by the drug treatment. In this aspect, a wealth of proteomic discoveries have seldom been translated into validated drug mechanisms, most of which stayed as preliminary findings that could lead to confirmed targets by in-depth functional studies. For example, celecoxib (Celebrex®; Pfizer, Inc, New York, NY, USA), originally a widely prescribed nonsteroidal anti-inflammatory drug, has recently been shown to have anticancer properties with...
uncertain mechanisms, ranging from modulating the expression of Bcl-2 family members and mitochondria-mediated apoptosis,\textsuperscript{95,97} to inhibiting nuclear factor kappa B,\textsuperscript{98,99} Akt,\textsuperscript{100} and Stat3\textsuperscript{101} signaling pathways. Proteomic analysis of human oral squamous cell carcinoma found that celecoxib treatment induced ten- to 20-fold overexpression of heterogeneous nuclear ribonuclear protein C.\textsuperscript{102} Similarly, global proteomic profiles of colorectal cancer cells before and after treatment with celecoxib revealed significant alterations among multiple proteins involved in diverse cellular functions ranging from glycolysis, protein biosynthesis, DNA synthesis, messenger RNA processing, protein folding, phosphorylation, redox regulation, to molecular chaperon activities.\textsuperscript{103} However, none of the proteomic studies has been conclusive in the mechanistic interpretation of numerous alterations in protein expression.

The Wnt-β-catenin signaling pathway is a developmental signaling pathway that plays a critical role in regulation, differentiation, proliferation, and apoptosis; thus, aberrant Wnt-β-catenin signaling is widely implicated in numerous cancers.\textsuperscript{104–106} However, identification of targeted Wnt-β-catenin pathway inhibitors in cancer patient treatment has been hampered by the limited number of pathway components that are amenable to small molecule inhibition. Recently, Huang et al\textsuperscript{107} used an iTRAQ approach to identify a small molecule, XAV939, that selectively inhibits β-catenin-mediated transcription. Interestingly, XAV939 also inhibits the poly-adenosine diphosphate-ribosylating enzymes, tankyrase 1 and tankyrase 2, which interact with a highly conserved domain of axin and stimulate its degradation through the ubiquitin–proteasome pathway.\textsuperscript{107} Tankyrases, involved in fundamental cellular processes such as telomere homeostasis and Wnt signaling, are potential telomere-directed anticancer targets.\textsuperscript{108,109} Therefore, XAV939 is not only a useful tool in understanding Wnt-β-catenin signaling, but also a potential drug that targets Wnt-β-catenin and telomerases.

Chemical proteomics aimed at characterizing the effects of drug candidates could sometimes lead to the discovery of new drug targets. For example, sulforaphane (SFN) is known to have antimicrobial and anticancer properties in experimental models.\textsuperscript{110–112} SFN can modulate multiple cellular targets involved in cancer development including DNA protection, inhibition of cancer cell proliferation, induction of apoptosis, inhibition of neoangiogenesis, progression of benign tumors to malignant tumors, and metastasis.\textsuperscript{112,113} Mastrangelo et al\textsuperscript{114} identified serotonin receptors as the novel targets of SFN by proteomic analysis in Caco-2 colon cancer cells. This finding may not only shed light on the serotonin-mediated signaling pathways in colon cancer, but it may also lead to the development of potential novel therapeutic agents targeting serotonin. In another study, phosphoglucomutase 3 was identified by proteomic analysis and may contribute to SFN-induced cell death in the LNCaP prostate cancer (PCa) cells, which make phosphoglucomutase 3 a potential molecular therapeutic target for PCa.\textsuperscript{115} While these are very early studies on the discovery of druggable protein targets in the relevant disease, they do open up new venues for the development of novel treatment regimens.

Although the field of chemical proteomics has proven its value in identifying novel drug targets, several challenges still remain to be overcome. First, the chemical probes must be carefully designed to covalently attach to proteins of interest and allow purification and/or identification. Second, the spectrum of available activity-based probes needs to be broadened in order to target additional enzyme classes. In addition, the development of high-throughput and gel-free assays, in conjunction with activity-based probes, will be required to enhance the experimental value of chemical proteomics. The continued success of chemical proteomics depends on the design of novel probes or new probe classes that can specifically target diverse sets of enzyme families, as well as on an unbiased assessment of the full spectrum of drug–target interactions and their molecular MoA.

**Target discovery based upon differential disease state tissues (tissue proteomics)**

Proteins and their spatiotemporal distribution changes play a central role in biological processes, including cancer initiation and progression. Analysis of a specific homogeneous cell type from tumor tissues could reveal molecular changes that take place in tumorigenesis since the concentration of disease-related proteins are likely to be much higher within or near the tumor areas. Recently, large-scale protein identification and comparative quantitation of highly complex protein mixtures have been achieved with proteomic strategies based upon differential disease state tissues, known as tissue proteomics.\textsuperscript{115,116–120} Remarkable advances in tissue proteomics have been propelled by the rapid development of efficient methodologies and techniques in innovative sample preparation, sophisticated MS instrumentation, and powerful bioinformatics tools. The goals of tissue proteomic research focus on early or more accurate diagnosis, improvement of therapeutic strategies, and better evaluation of prognosis and/or prevention of a given disease, as well as on the
identification of novel drug targets based upon the differential protein expression between “control” and “case” groups for the disease. A major hurdle in tissue proteomics analysis is the variability observed among the tissue samples due to the heterogeneity of tumor tissues containing cancer cells, as well as inflammatory, vascular, and connective tissue cells. Detailed proteomic analyses of these clinically valuable samples require meticulous preparation procedures, which is a major current focus of tissue proteomics. So far, tissue proteomics has uncovered large numbers of proteins with altered expressions in tumor tissues by analyzing both fresh–frozen biopsy samples and archived tumor tissues stabilized with formalin fixation and paraffin embedding (FFPE). However, very few, if any, of these proteins have actually become novel diagnostic biomarkers or therapeutic targets. As illustrated in the examples below, most tissue proteomic studies remain as preliminary investigations that identified differentially expressed proteins in diseased tissues. While these proteins could be potential prognostic markers or therapeutic targets, there is a general lack of in-depth functional and validated studies that follow up on these initial screening results.

Tissue proteomics has been employed quite frequently in the study of lung cancer for molecular mechanism elucidation as well as for novel drug target discovery.121–126 Peng et al123 studied the protein profile changes between human pulmonary adenocarcinoma tissues and paired surrounding normal tissue with two-dimensional (2D) gel electrophoresis and ESI-Q-TOF (electrospray ionization, quadrupole, time-of-flight) MS/MS instruments. Thirty-two differentially expressed proteins (>2-fold change; \( P < 0.05 \)) were identified in pulmonary adenocarcinoma when compared to normal tissues. Knockdown of pyruvate kinase isozyme 2 (PKM2), one of the overexpressed proteins, led to a significant suppression of cell growth, to the induction of apoptosis in vitro, and to tumor growth inhibition in vivo. Moreover, the shRNA-expressing plasmid targeting cofilin-1, another overexpressed protein, significantly inhibited tumor metastases and prolonged survival in vivo. This tissue proteomic analysis and validation indicated that PKM2 and cofilin-1 could be potential therapeutic targets for pulmonary adenocarcinoma. It should be noted that small molecule inhibitors of PKM2 were identified and shown to inhibit cancer cell glycolysis and increase cancer cell death following the loss of growth factor signaling,127 which was consistent with the proteomics-driven conclusion. Cofilin-1 has long been associated with increased tumor metastasis due to its role in regulating cytoskeleton dynamics,128 and the inhibition of cofilin-1 by small molecule inhibitors has been demonstrated to enhance actin depolymerization.129 In another example, Carretero et al124 performed an integrated genomic and proteomic study for the identification of genes and phosphoprotein status associated with Lkb1 loss and progression to invasive and metastatic lung tumors in primary and metastatic de novo lung cancers. Phosphoproteomic analysis determined that two key modulators of focal adhesion dynamics, Src and focal adhesion kinase, are upregulated by Lkb1 loss during NSCLC progression. Moreover, the combined inhibition of Src, phosphoinositide 3-kinase, and MEK1/2 resulted in a synergistic tumor regression. These results point towards a mechanism underlying the increased propensity for metastases seen in Lkb1-deficient lung tumors, and they identified the Src signaling pathway as a molecularly targetable pathway for the treatment of LKB1-deficient NSCLC in humans. Therefore, development of therapeutic Src inhibitors, including dasatinib and saracatinib, will be a valid therapeutic strategy for the treatment of LKB1-deficient NSCLC.

Proteomic analysis of PCa tissues associated with multistage tumor progression also provides a valuable source of clinically relevant biomarkers and novel therapeutic targets.130,131 For example, Ummanni et al132 recently reported the differential protein expression patterns from histologically characterized PCa tumor tissues and surrounding benign tissues of individual PCa patients based upon 2D differential gel electrophoresis coupled with MS. The study identified 118 protein spots differentially expressed in cancer (n=24) when compared to benign (n=21) prostate tissues adjacent to cancerous tissues. Analysis of these gel spots by matrix-assisted laser desorption/ionization–time-of-flight MS/MS revealed 79 unique proteins. Moreover, system biology analysis of proteomic results revealed several novel drug targets of PCa development and/or progression including eIF4A3, DDAH1, ARG2, Prdx3, and Prdx4, although functional validation of individual targets have yet to be performed. It is evident that proteomic analysis of multistage PCa tissues could provide new insights into PCa progression and potentially lead to the design of novel diagnostic and therapeutic strategies.

Efficient peptide/protein extraction approaches are crucial to the success of tissue proteomic analysis. Two major strategies of mining proteomic information from FFPE archive tissue samples have been developed.133–138 One strategy aimed to recover full-length proteins by heat treatment in suitable buffers with consequent reversion of the formaldehyde-induced cross-links. The operation conditions have been optimized by various labs since Shi et al139 initially established this method. Application of high temperatures and the
addition of detergent sodium dodecyl sulfate were indicated as two critical conditions for enhanced protein extraction yields.\textsuperscript{140–142} Because the protein was deposited in large amounts as insoluble, densely packed aggregates, applications of 40,000 psi pressure were reported to recover 96\% of proteins from a tissue surrogate model, compared with a 26\% recovery rate at 14.7 psi.\textsuperscript{138,144} Another major strategy is based upon the direct proteolytic digestion of intact FFPE archive tissue samples, followed by liquid chromatography–MS/MS characterization of the complex peptide mixture.\textsuperscript{145} Tissue solubilization can be achieved in various buffers including sodium dodecyl sulfate–dithiothreitol,\textsuperscript{145} radioimmunoprecipitation assay,\textsuperscript{146} acetonitrile ammonium bicarbonate,\textsuperscript{147} Tris-HCl\textsuperscript{148} under different temperatures, and pH conditions for the optimal yield of peptides. As a successful example, Hwang et al\textsuperscript{147} characterized 428 prostate-expressed proteins from FFPE archive tissue samples for the discovery of PCA biomarkers and potential drug targets using the shotgun approach. Results to date demonstrate that direct trypsin protein digestion is an effective sample preparation strategy for proteomic analysis of FFPE archive tissues.

Approximately 70\% of diagnosed breast cancers express the estrogen receptor (ER+) whereas ER– breast cancers are not well differentiated and clinically tend to be more aggressive.\textsuperscript{149–152} A global proteomic characterization and quantitative comparison of ER+ and ER– breast tumors were recently performed on fresh–frozen breast tumor tissues.\textsuperscript{153} The study identified 2,995 unique proteins including a number of receptor tyrosine kinases and intracellular kinases that are abundantly expressed in both ER+ and ER– breast cancer tissues. Using a label-free quantitative approach, 236 proteins were found differentially expressed between ER+ and ER– breast tumors. Compared with ER– breast tumors, 141 proteins were selectively upregulated, while 95 proteins were downregulated in ER+ tumors. Molecular function analysis of gene ontology showed that the dehydrogenase, reductase, cytoskeletal proteins, extracellular matrix, hydrolase, and lyase categories were significantly enriched in ER+ breast tumors, whereas selected calcium-binding proteins, membrane traffic proteins, and cytoskeletal proteins were enriched in ER– tumors. Biological process and pathway analysis indicated that proteins related to amino acid metabolism, proteasome, and fatty acid metabolism were overexpressed in ER+ tumors, while proteins related to the glycolysis pathway were overexpressed in ER– tumors. Given the clinical challenges in treating ER– breast cancer, such in vivo findings of differentially expressed proteins in ER– tumors are especially significant because of their pathological relevance. These proteins may serve as potential therapeutic targets for ER– breast cancer if functionally validated. Recently, Cabezon et al\textsuperscript{154} used a systematic 2D gel-based proteomic profiling strategy, applied to the analysis of 78 fresh triple-negative breast cancer (TNBC) tissue biopsies, in combination with a three-tier orthogonal technology (2D polyacrylamide gel electrophoresis/silver staining coupled with MS, 2D Western blotting, and immunohistochemistry) approach, they identified and validated one specific protein, Mage-A4, which was expressed in a significant fraction of TNBC and Her2-positive/ER– lesions. The existence of immunotherapeutic approaches specifically targeting this protein, or Mage-A protein family members, provides novel management options for TNBC and Her2-positive/ER– patients bearing Mage-A4 positive tumors.

Tissue proteomics for cancer drug target identification is still in its infancy. Although promising results have been reported, there has yet to be a breakthrough that is close to becoming clinically applicable. The challenge will be to perform robustly designed, large retrospective studies including independent validation sets, followed by prospective validation studies to demonstrate the clinical benefits for patients. The development of siRNA in vivo techniques will greatly speed up selection of useful proteins. Characterization of discriminator proteins will provide new molecularly targeted anticancer drugs. Faced with the complexity of cancer tissue proteomics and its great clinical potential, a global coordination of ongoing efforts appears to be crucial.

**Conclusion**

In this review we provided an overview of three major approaches of proteomics employed in discovering potential therapeutic targets for cancer. Target identification in resistant cancer populations is one of the most promising tools for drug target identification, and it can uncover multiple drug targets for combinatorial therapeutic solutions to combat resistance and achieve longer-term improvement in disease outcomes. Chemical proteomics approaches not only identify protein targets for drugs that exert known biological activities, but they can also discover previously unknown targets for drugs of known MoA. The tissue proteomics approach offers the advantage of direct clinical relevance and can complement mechanism-based approaches like chemical proteomics.

Proteomics continues to evolve rapidly both in instrumental advancement and innovative applications in biomedical research, particularly in the field of cancer. The complexity of cancer biology requires not only in-depth functional studies for mechanistic elucidation, but also “unbiased”
Novel therapeutic targets in cancer

System approaches for a global view of interactive signaling networks that reflect the cancer disease stage. Cancer proteomics has uncovered a phenomenal number of cancer-specific proteomic alterations, shedding light on previously unknown mechanisms in tumorigenesis, cancer progression, and metastasis. Perhaps more prominently, proteomics has provided an unmatched wealth of potential biomarkers that could be clinically used for the diagnosis and prediction of treatment responses. The utility of proteomics as a valuable tool for the discovery of new cancer treatment targets has been increasingly recognized in recent years. We note the remarkable progress in several directions when cancer proteomics plays an important role in drug discovery, as illustrated in Figure 2. Novel proteomics methodologies, both MS-based and non-MS-based, continued to be optimized and perfected using high-resolution and high-speed MS, simplified and robust array technologies for tumor tissue, cell, subcellular compartment, and other relevant biological samples. Proteomics has been increasingly used to elucidate the mechanisms of drug resistance in cancer with the hope that novel protein targets could emerge for therapeutic intervention of the resistant disease. In other innovative frontiers, chemical proteomics studies examining drug-induced biological perturbations often reveal new clinically relevant biomarkers and drug targets; in dissecting mechanisms of cancer progression, proteomics also identified previously unknown or unconsidered druggable targets.

However, despite the tremendous progress and potential of proteomics for target discovery, there are serious challenges. A lack of coordinated efforts to follow up with the proteomic discovery of novel therapeutic targets largely accounts for the current gap between drug development (design, synthesis, and optimization) and new drug targets emerging from numerous proteomic studies of cancer. Most of the proteins identified in the

Figure 2 Quantitative proteomic analysis in target discovery, validation, and drug development.

**Abbreviations:** Bio, biological; SILAC, stable isotope labeling by amino acids in cell culture; iTRAQ, isobaric tag for relative and absolute quantitation; TMT, tandem mass tag; prep, preparation.
proteomic analyses of drug resistance have not been validated for their roles in specific disease processes and their potential in clinical use. Functional studies involving in vitro manipulations of gene expression using specific pharmacological inhibitors, antisense RNA, RNA interference, or gene knockout experiments should be an integral part of a proteomic study for target discovery. Clinical validations of candidate biomarkers should also be included in retrospective and prospective studies. Future investigations should place more emphasis on functional studies and on the clinical validation of novel targets for their translation in clinical trials with greater speed and higher success rates. Proteomics is also expected to play a major role in the preclinical and clinical research of targeted and combinatorial therapies.

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References


102. Looking at the literature, the role of targeted chemical proteomics in pharmacology is becoming increasingly prominent. The methodology involves the identification of protein interaction partners using quantitative mass spectrometry and bead proteomes, which can aid in understanding the mechanisms of action of clinical ABL kinase inhibitors and other drugs. This approach has been used in the identification of specific protein interaction partners in cancer cells, providing valuable insights into the pharmacology of drugs.


