Proteomics plays a vital role in biomedical research in the post-genomic era. With the technological revolution and emerging computational and statistic models, proteomic methodology has evolved rapidly in the past decade and shed light on solving complicated biomedical problems. Here, we summarize scientific research and clinical practice of existing and emerging high-throughput proteomics approaches, including mass spectrometry, protein pathway array, next-generation tissue microarrays, single-cell proteomics, single-molecule proteomics, Luminex, Simoa and Olink Proteomics. We also discuss important computational methods and statistical algorithms that can maximize the mining of proteomic data with clinical and/or other ‘omics data. Various principles and precautions are provided for better utilization of these tools. In summary, the advances in high-throughput proteomics will not only help better understand the molecular mechanisms of pathogenesis, but also to identify the signature signaling networks of specific diseases. Thus, modern proteomics have a range of potential applications in basic research, prognostic oncology, precision medicine, and drug discovery.

INTRODUCTION

Since the successful completion of the Human Genome Project that mapped the whole human genomes, a massive number of genomic markers have been identified and are being applied to medical sciences1. Many of them have been developed as routine tests in the clinic. However, a significant limitation of genomic or transcriptomic profiling studies is that genomic and transcriptomic data, which only provide indirect measurements of cellular states, may not accurately reflect the corresponding protein changes. These data fail to reveal changes in posttranslational modifications (PTMs), including phosphorylation and protein degradation. Therefore, the genomic data alone cannot bring a full picture of the disease mechanisms with comprehensive understanding2. Nowadays, the Human Proteome Project (https://hupo.org/) has been launched to characterize the entire human proteome by advanced proteomic techniques, which is the next major challenge3. The guidelines on interpreting proteomic data have also been published and recently updated4.

Proteomics, as the combination of proteome experimentation and data analysis, analyzes protein composition, structure, expression, modification status, and the interactions and connections between proteins at an overall level5. It offers complementary information to genomics and transcriptomics. It is also essential for generating a map of the complex, interconnected pathways, networks, and molecular systems, which directly control the major life activities such as cell proliferation, differentiation, senescence, and apoptosis. With the substantial improvement of experimental technology over the past decade6, the proteomics methods have been evolved from conventional methods, such as immunohistochemistry (IHC) staining, western blot, and enzyme-linked immunosorbent assay (ELISA), to high-throughput methods such as tissue microarray (TMA), protein pathway array and mass spectrometry7. Those high-throughput proteomics techniques not only decrease analysis time but also increase the accuracy and depth of proteome coverage. With the advents of bioinformatics and modern multi-analytes “omics” technologies (Supplementary Fig. 1), proteomics holds a great promise for uncovering the molecular mechanisms that underlies diseases towards the discovery of novel biomarkers8 and can be used as specific diagnostic assays, prognostic predictors, and therapeutic targets to enhance personalized medicine further9,10.

In this review, we will discuss the advances in high-throughput proteomic techniques, statistics and algorithms, progress in applying proteomics to disease diagnostics, current challenges and future perspectives.

HIGH-THROUGHPUT PROTEOMIC TECHNIQUES

With the rapid development of high-throughput technology6, several new technologies are widely used in proteomics and metabolomics in recent years. Regardless the specific technique, these global proteomic approaches (Fig. 1) can be divided into three phases, namely discovery, network-analysis and clinical proteomics. Discovery is the initial phase to identify the amino acid sequence and unknown protein structure with qualification11. We then in network-analysis phase build the global signaling networks and investigate the relations among the known proteins to explore the potential biomarkers with verification. Finally, in the
clinical proteomics phase\(^{12}\), we develop clinical assays related to the productization of the biomarker or panel fitting the clinical flow. The commonly used high-throughput proteomic techniques include mass spectrometry, protein pathway array, next generation tissue microarrays and Luminex and will be discussed in details below.

**Mass spectrometry**

Mass spectrometry (MS) has been developed as one of the most essential and popular tools to identify proteins and their isoforms, and quantify posttranslational modifications, either via the fragments directly or the specific proteolytic activity responsible for their formation\(^{13-15}\). The most significant effect of MS is to discover and detect an intact protein or a subset of composite or surrogate peptides as MS-based quantitative proteomics that traditional immunoassays find incredibly challenging or impossible. MS can be combined with multiple separations and pre-fractionation techniques to identify the target protein/peptide and improve identification accuracies and yields\(^{16}\). For example, two-dimensional polyacrylamide gel electrophoresis (2D-PAGE) is based on electrical charge and molecular weight, while liquid chromatography (LC) based on polarity, electrical charge, and protein molecular weight. For an example of 2D-PAGE, mixtures of proteins are separated by the electrical charge as isoelectric point (pI) in the first dimension and further separated by molecular weight in the second dimension on 2-D gels. The protein samples from different resources, which were labeled by different cyanine dyes such as Cy2b, Cy3, and Cy5 as reporter fluorophores, can be processed in the same 2D-PAGE to purify the target protein and enhance the detection accuracy\(^{17}\). After being digitalized 2D-PAGE by fluorescence scanner and the image analysis\(^{18}\), the interesting or significant spots in the gel are cut out and enzymatically digested to peptides for MS as matrix-assisted laser desorption/ionization-time of flight (MALDI-TOF) MS analysis where each digest yields a peptide mixture that can be analyzed by bottom-up experiment. Although 2D-PAGE has traditionally been used as a standard procedure for proteomics research, gel-based techniques tend to be labor-intensive and time-consuming, and are therefore not suitable for high-throughput proteomics. By contrast, LC or high-performance liquid chromatography (HPLC) allows continuous separation of thousands of proteins from complex mixtures and can be combined with MS as LC-MS for increased throughput\(^{19-21}\). Among them, Reversed-phase liquid chromatography (RPLC) is the most commonly used LC-based separation platform. It is characterized by the distribution of compounds between a water-containing mobile phase and a relatively nonselective stationary phase and other chromatography formats can be added prior to the RPLC separation to improve the dynamic range of measurement\(^{22}\).

According to different strategies of processing, MS-based methods can be divided into top-down, bottom-up, and shotgun approaches. In the top-down proteomics method, a full-length protein, which can be subsequently fragmented inside the MS and the masses of the fragments be recorded, is directly sent for MS analysis\(^{23}\). By contrast, proteins are enzymatically or chemically digested into peptides that serve as input to the MS equipment in bottom-up proteomics techniques. Moreover, shotgun proteomics is a particular case of bottom-up proteomics where the whole proteins in a complex mixture, such as serum, urine, and cell lysates, are cut into peptides and followed by multidimensional HPLC-MS, which aims to generate a global profile of protein mixtures as genome "shotgun" sequencing\(^{24}\). On the other hand, the separation of peptides prior to MS is not necessarily needed in the bottom-up strategy. Then MS data is matched to identify the target proteins and their associated modifications in the protein sequence database by data-dependent discovery engines\(^{25}\), which can be divided into peptide scoring, protein scoring, and finally protein inference\(^{26}\).

**Protein pathway array**

Human diseases, especially cancers, are often a complicated biomedical process attributable to complex protein-based signaling network pathway alterations that control cell behaviors, such as apoptosis, invasion, and metastasis\(^{27}\). Uncovering the underlying changes in multidimensional protein signaling networks not only aids in understanding the molecular mechanisms of pathogenesis but also identifies the characteristic signaling network signatures that are unique for the type or stage of the diseases\(^{28-32}\). Measuring many proteins simultaneously is of great importance to the theory of protein-protein interactions (PPIs) in the signaling network, which is a big challenge to conventional immunoassays such as western blot. Therefore, high-throughput proteomic tools were increasingly used for biomarker discovery in basic, translational and clinical research.

Protein pathway array (PPA)\(^{27}\), a gel-based high-throughput platform, employed antibody mixtures to detect antigens in a protein sample which can be extracted from biopsy or tissue. In this approach, microdissection of tumor tissue could be applied to maximize the proportion of the proteins from tumor tissue instead of the surrounding benign tissue\(^{27}\). Then, the immunofluorescence signals of antibody-antigen reactions are converted to numeric data as the value of protein expression by Quantity One (https://www.bio-rad.com/en-us/category/image-lab-software-suite?ID=52911579-0715-48f4-b3de-766b9222582) from Bio-Rad. The biomarkers and proteomic networks can be explored and trained after data normalization and appropriate statistic modeling. PPA has been applied to many diseases such as essential thrombocythemia\(^{29}\) and papillary thyroid carcinoma\(^{30}\). Its high-throughput protein profiles in a robust quantitative manner provided an advantage over traditional methods.

**Next generation tissue microarrays**

Immunohistochemistry staining, as one of the traditional and reliable research methods, uses an enzyme-linked chromogenic substrate for detection and requires microscopic examination\(^{35}\). It remains a time-consuming and subjective process and produces a
qualitative or semiquantitative assessment of protein expression because of the nonspecific stain or background noise. As technology advanced and high-throughput demand increased over the last decade, the TMA gradually began to be widely used in both research and clinical fields. TMA contains many small representative tissue cores of formalin-fixed paraffin-embedded (FFPE) or frozen blocks from hundreds of different cases assembled in an array fashion on a single histologic slide, and therefore allows a large-scale antibody-based molecular analysis of multiple samples at the same time. Therefore, it is a practical and valuable tool to confirm and verify new biomarkers generated from PPA or MS proteomics methods. Thus, it is often used in an independent cohort and identifies the location of the target proteins in the cell membrane, cytoplasm, or nucleus. Since digital pathology with multiple smart microscopes has been developed rapidly in the past year, a new approach of TMA, next-generation tissue microarrays (ngTMAs), was recently created. It allows annotations to be placed directly on the digital slides for a higher and high throughput without major compromise on quality. Due to its improved sensitivity and rapid, large-scale detection capabilities, ngTMAs has become a powerful tool to improve the quality of TMAs used in clinical and translational research but could be more widely used.

**Multiplex bead- or aptamer-based assays**

Proteomics plays a critical role in clinical practice, although there are gaps and limitations to translate proteomics from basic molecular research to clinical use. Multiplex bead- or aptamer-based assays have been developed but have various sensitivity and specificity. Therefore, caution and in-house validation studies must be used before the assay is applied to clinical samples.

Luminex bead-based array system is increasingly used in protein profiling applications in recent years but have various sensitivity and specificity. It makes the detection of proteomic biomarker panel reliable, fast and able to cope with dynamic changes in the variety of clinical practices. Luminex uses different, flexible fluorescent-labeled beads that are spectrally distinguishable and coated with a different capture antibody or probe to identify the antigen or mutation in samples. It is able to detect up to 500 analytes (FLEXMAP 3D Platform: https://www.luminexcorp.com/flexmap-3d/) in a single sample using a 96-well plate or 384-well plate. For proteomics usage, megaplex microspheres are tagged to allow fluorescent detection and can be used in the development of the multiplex immunoassays by labeling multiple target antibodies. After microsphere activation and conjugation reactions, a panel of beads-antibody complexes is mixed and incubated with samples to capture protein analytes in the sample. Then the sandwich structure of bead-antibody-antigen complexes is passed and counted by a flow cytometer using different fluorescent of beads. Therefore, the high-throughput Luminex system has a great potential for fast multiplexed analysis of panels of genetic, proteomic, metabolic biomarkers associated with disease diagnosis, prognosis, and therapeutics in patients.

Another widely used platform is Meso-scale Discovery ( MSD) assay (https://www.mesoscale.com/) which may be multiplex, single-plex or ultrasensitive. It has been used mostly for cytokine detection in the mice with type 1 diabetes or radiation treatments, and the astrocytes with neuronal networks. It has also been compared with other platforms or detection assays. For human cytokine profiling, the MSD assay is more sensitive than Luminex assay but less specific, while a recent study shows low or no significant correlations for detecting most of the cytokines (except interleukin 6) among Luminex xMAP®, MSD V-Plex® and Quanti- kine assays. A study of 38 epileptic children shows a freeze-thaw cycle results in consistent measurements in 46% (6 of 13) of the analytes using Luminex high-sensitivity assay, 11% (1 of 9) using Luminex standard-sensitivity assay, and in no analytes using MSD assay. Therefore, the Luminex high-sensitivity assay appears to have better precision than the other 2 assays for epilepsy research. For detecting plasma Alpha-Synuclein in Parkinson’s disease patients, MSD assay has a smaller effect size than Quanterix assay but correlated well with Biolegend.

One of the widely used bead-based multiplex assays is Simoa® (Single Molecule Array, owned by Quanterix). It covers 6 disease areas, is customizable, and includes 109 oncology, 26 neurology, 19 immunology, 13 cardiology and 45 infectious disease assays as of May 2022. Their platform can be used to detect 6 to 10 biomarkers in a single test, and can detect as low as 1 fg/mL of proteins. As a highlight of its performance, Simoa® had the highest sensitivity and precision in a comparison of platforms’ performance in post-traumatic stress disorder and Parkinson’s disease, as well as the lowest variation and highest effect size in a 3-platform comparison on Parkinson’s disease.

Antibodies are the primary detection tool in the bead-based assays but face challenges in high-throughput platforms. To meet the challenge, protein-binding reagents are produced such as slow off-rate aptamer and have been commercialized as the SOMAscan® assay. The aptamer-based SOMAscan® assay can assess expression of 1,000 to 9,000 antigens in a single test and has an impressive dynamic range (8 orders of magnitude), a great sensitivity (lower detection limit, 40 fM) and a high precision (median coefficients of variance = ~5%). In a study on embryonic stem cells, SOMAscan has a higher reproducibility, a higher sensitivity and a larger dynamic range than nano LC-MS/MS and RNA sequencing, but fewer features to detect. The SOMAscan’s results are overall comparable to those of nano LC-MS/MS and RNA sequencing. In a study on patients with end-stage renal disease, SOMAscan correlated very well with ELISA in detecting 2 of the 3 targeted proteins, but not in the last one. However, compared with antibody-based Olink platform, SOMAscan® assay shows a wide range of correlation in assessing protein expression in 2 cohorts of chronic obstructive pulmonary disease and thus should be used with caution. Moreover, despite the greater coverage and overall good correlation, SOMAscan did not reveal bigger odds ratios of the proteins linked to acute kidney injury than those revealed using one of the immunoassays including MSD (electro-chemiluminescence platform), Access (paramagnetic-chemiluminescence platform) and Unilc (chemiluminescence platform) and Biochip (multiplexed ELISA platform).

**Proximity extension assay (Olink)**

Proximity extension assay, as one of proximity-dependent ligation assays, is based on oligonucleotide–linked antibody pairs that have slight affinity to each other. When these oligonucleotide-linked antibodies are brought in proximity, the two unique oligonucleotides linked to the antibodies will be extended by a DNA polymerase and amplified exponentially later. Quantitative real-time PCR is often used to amplify and quantify the oligonucleotides in the sample. Thus, oligonucleotides can serve as a unique surrogate marker of specific antigens which the antibodies recognize. As described in its original reports, the 5 specific assay steps include: 1. Oligonucleotide-linked antibody pairs are added into the sample; 2. The probe pairs bind to the antigen and subsequently the probe oligonucleotides are brought into close proximity; 3. The oligonucleotides form pair-wise binding of matching probe-pairs; 4. The matching probe-pairs are amplified using universal primers. The process is termed as pre-amplification due to the lack of specific primers; 4. The matching probe-pairs are digested using uracil-DNA glycosylase and unbound universal primers are removed; 5. The pre-amplified probe-pairs (DNA templates) are quantified using specific primers and quantitative real-time PCR. Multiplex and 96-multiplex detection methods have been developed and also show very robust signal.
high sensitivity and specificity. The Olink assay has been applied to several clinical fields with great success, including coronavirus disease 2019, traumatic brain injury and renal diseases. It can simultaneously quantify over 3,000 proteins in a miniscule amount of sample (e.g., a few microliters).

**Nanopore based single-molecule proteomics**

Nanopore has been increasingly used for DNA or RNA sequencing and tried on proteomics. Its early application to proteomics was seen in sequencing peptides of mycobacterium. Peptide-oligonucleotide conjugates and measurements with nanopore-induced phase-shift sequencing were used and seemed able to sequence short peptides. Later, addition of helixase was found effective to reduce the reading error rate to 30 rereads per million. It is also proposed to combine nanopore with other techniques such as fluorescence labeling and protein-fragmentation for better readouts. The major challenges of nanopore based single-molecule proteomics are low efficiency and lack of sufficient sensitivity for detecting PTM.

**STATISTICS AND ALGORITHMS**

Traditional statistics methods, such as Student’s t-test and one-way analysis of variance (ANOVA), have various biases and may be time-consuming to handle big data. Therefore, new high-throughput approaches or machine learning-based algorithms are needed to process big data that are generated from multi-omics. Machine learning can be divided into supervised learning and unsupervised learning approaches generally. In terms of supervised learning, it applies a “labeled” training set to train a model and predict a qualitative or quantitative output, such as classification and regression. By contrast, unsupervised learning has an unlabeled output set and enables the algorithms to determine and identify the natural patterns with shared similarities in an unknown dataset, such as cluster analyses. Artificial intelligence and digital pathology are involving rapidly and will play an even more important role in research, pathology and medicine while some traditional statistic tools remain important such as normalization and batch effect removal.

**Normalization**

The most common and necessary form of big data pre-processing phase is normalization, which is being used to centralize and rescale all of the data as a whole numerical matrix to improve their numeric stability, overall performance and model fit. All machine learning-based statistic models, such as distance-based cluster analysis, regression, and principal component analysis, are susceptible to unscaled data distribution. For example, the most commonly used formula of normalization is Z-score, which is also called the standard score. Z-scoring the data centers the raw data by subtracting the mean (average) of a group of values of expression of genes or proteins first to reduce the influence of an extreme outlier that could affect the mean of a dataset with a small number of samples and then divides each data variable by the standard deviation (SD) to scale the data variable. Furthermore, common housekeeping genes and proteins, including
GAPDH (glyceraldehyde-3-phosphate dehydrogenase) and beta-actin, whose expression is considered the same in all samples, can also be used to normalize array data as a house-keeping gene. Besides, another preferred treatment for the ratio data is to log the data but not to Z-score the data.

Significance analysis of microarrays

Significance Analysis of Microarrays (SAM) (https://statweb.stanford.edu/~tibs/SAM/) is a supervised learning program for large-scale gene or protein expression data mining developed by the Stanford University Statistics and Biochemistry Labs. SAM, a Microsoft Excel add-in package, is a widely used high-throughput permutation-based approach to identify differentially expressed proteins between sets of samples in abundance proteomics data using modified t-statistics (q-value) which measures the strength of the relationship between protein abundance and disease outcome. Unlike the regular t-test for small sample size, SAM algorithm is an excellent fit for big data to minimize the number of false positives and negatives by permuting the columns of the protein abundance and automatic imputation of missing data via the nearest neighbor algorithm. Furthermore, one of the SAM’s valuable features is that it gives estimates of the False Discovery Rate using data permutations, which is the proportion of proteins likely to have been identified by chance as being significant (Fig. 3).

Clustering and discriminant analyses

Hierarchical clustering analysis (HCA) has been used to cluster the big data by forming a mathematical model based dendrogram. Several optimized mathematical formula-based models are created to measure the distance between data points, including Manhattan (L1) distance, Euclidean (L2) distance, Pearson correlation, and others. The Euclidean distance is the most commonly used but is vulnerable to outliers in non-normal distribution data especially, but might be inferior to Pearson correlation in analyzing proteomic data. Manhattan distance requires the strict normalization. Pearson correlation is a scale-invariant of the similarity measure, etc. It must be noted that the choice of distance measure impacts the performance of HCA and thus should be decided with caution. Moreover, there are different principles which can be calculated to measure the distance between clusters, such as average distance, minimum distance way, and maximum distance ways. The average distance way uses the average of all data points in one cluster to map to the closest one of the other clusters. Both distance measure and its calculation formula determine which samples and clusters are grouped together. Based on these 2 metrics, the model is optimized to keep the distance between the data points within one cluster as close as possible in the numerical matrix, but keep the distance between the data points in different clusters as far as possible. Besides, clustering results are also affected by both input data and selected variables, such as feature distributions and biomarkers. For example, the clustering results will be significantly different if samples and biomarkers are added, deleted, and/or replaced. Therefore, essential variables (biomarkers), sample selection criteria and study goals should be clearly defined prior to a HCA for robust and reproducible analysis. In addition, HCA can be divided into one way and two-way HCAs. Two-way HCA indicates that the data is clustered using the X-axis (samples) and Y-axis (biomarkers) at the same time (Fig. 4a) comparing with one way, which means either axis clusters the data according to study design.

Additionally, there is a particular clustering analysis, called Grid Analysis of Time-series Expression (GATE) (Fig. 4b), to analyze and visualize high-dimensional biomolecular according to time series. GATE, as an integrated computational software platform, uses a correlation-based clustering algorithm to arrange time series or continuous-time points on a two-dimensional hexagonal array. It dynamically colors individual hexagons according to the
expression level of genes or proteins to create animated movies of systems-level molecular regulatory dynamics. Furthermore, GATE allows interactive interrogation of movies against a wide variety of knowledge datasets, such as Protein interaction hubs, Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways, Kinase enrichment analysis (KEA), WikiPathways pathways, etc, to infer potential regulatory control mechanisms from patterns of correlation. Those dynamic protein-protein interactions and clustering are able to allow investigating the continuous changes of cell lines or animals at different time points and dosages of treatment, the snapshots of the disease progression, and the different stages of cancer.

In contrast to clustering analysis that classifies known samples, (predictive) discriminant analysis classifies unknown samples based on what the algorithm learned and built in the training set\(^\text{95}\). For example, support vector machines (SVMs), which are not data-type dependent, can be applied to linearly separate the numerical or categorical data, and to identify the potential biomarkers as classifiers\(^\text{96}\). All samples need first to be divided into two groups as the training and validation sets. Then SVMs are trained in the training set for the most optimized algorithm, will be tested it later in the validation set and will finally produce the prediction rate by comparing the predicted with true values. The results will be affected by both input data (samples) and selected variables (also known as features or factors). In light of two data sets, both the training and validation sets must include all of the types of patient samples to cover any related clinical situations, such as stages, grades, histologic classification and complication, to eliminate the false negative and positive in the clinical practice. For example, the SVM algorithm may not recognize any “new” cases that have not been included in the training set, even if it is as simple as common sense for researchers or physicians. Besides, the samples in the training set need strict rule-in and rule-out criteria as well as keep the samples as many and diverse as possible to achieve the most accurate classification. In addition, the validation set can be the same or part of the training dataset as internal validation for retrospective evaluation, which is an option for a small sample size or population. However, an external validation cohort is recommended for prospective evaluation and increases reproducibility, generalizability and scientific rigor of the study. Several issues and problems of discriminant analysis must be noted and avoided during the analysis such as predictive versus descriptive discriminant analyses and linear versus quadratic models\(^\text{97,98}\). In summary, the main aim of (machine learning-based or not) discriminant analysis is to devise a computationally effective statistic model to classify multiple groups of subjects and identify the potential classifiers with a higher prediction rate.
Kaplan-Meier (K-M) curve and survival analysis

The Kaplan-Meier (K-M) curve is a time-event statistic method to investigate the relationship between the endpoint event and period of time. It can be used to evaluate survival time, disease recurrence, clinical trial, animal study, etc. For the survival analysis, data can be classified into two types, including complete data and censored data according to the endpoint event. Death and disease recurrence are the most commonly used endpoint events. Complete data is defined as the event occurrence during the experimental period. By contrast, censored data includes the subjects who were lost to follow up or experienced a non-qualiﬁed event before the end of the study. The time starting from a deﬁned point (zero time point) to the occurrence of a given event needs to be measured as input data. The higher the censored data ratio is in the study, the less accurate the results generally are. The K-M estimate is the simplest way of computing survival over time. The steep survival curve indicates a low survival rate or shorter survival period. It indicates that there might be confounding factors or effect modiﬁcation in the cohort which can be determined by stratiﬁed analysis and multivariate analysis, if the survival curves of each group cross and inferential analyses show statistical differences.

The two survival curves can be compared statistically by the rudimentary log-rank (Mann–Whitney U) test, which has been widely used, including Breslow and Tarone with different weight functions during computing. But they, usually as univariable analysis, do not allow to test the effect of the other disease-related variables. By contrast, Cox proportional hazards regression model, which is often used as multivariable analysis, can test the effect of other variables while identifying the independent variables of disease. For example, biomarkers can be analyzed alone with other risk factors such as age, gender, smoking history, and stage to determine whether it independently affects the prognosis. Therefore, an in-depth and comprehensive survival study of PPIs or microarray is to perform a log-rank test to identify the biomarkers that have statistically signiﬁcant ﬁrst and then analyze it with other risk factors together using the Cox regression model. The results of those double analyses can be classiﬁed into three categories: (1) Biomarkers have statistically signiﬁcant in both of the log-rank test and the Cox regression model. It means those biomarkers affect the prognosis as independent factors. (2) Biomarkers have statistically signiﬁcant only in the log-rank test but not in the Cox regression model. It means those biomarkers are correlated with risk factors as effect modiﬁcation to impact the disease of interest and may have confounding factors (Cox regression model may reveal them). (3) If biomarkers have statistically signiﬁcant only in the Cox regression model but not in the log-rank test, bias or study errors such as confounding bias need to be considered in the study. Moreover, the number of cases as complete data should be at least ﬁve to ten times greater than the number of variables as multiple secondary endpoints in the Cox regression model to avoid the type I error.

Besides, many popular regression models are being used to analyze the proteomics or microarray-based big data, and their functions are similar to the Cox regression model in varying degrees. For example, the multivariable logistic regression, which is a supervised classiﬁcation algorithm, is used to model the relationship between a set of continuous, categorical, or dichotomous independent variables and a dichotomous outcome as a dependent variable without time variable. Cox regression model incorporates time variable but is not able to process the missing values and censored data during a certain amount of time. The advantages of the logistic regression, in which the underlying concept is quite the same as linear regression, are assumed to be a linear association between the features and dependent variable (also known as outcome or label). However, it does not require that the variables normally distributed in the linear discriminant analysis. In addition, the dependent variable is quantitative in the multiple linear regression, rather than a binary outcome in logistic regression.

Principal component analysis

The main objective of principal component analysis (PCA) is to decrease the dimensionality of the big data by creating a set of new variables, called principal components, to represent the majority of the information within the original dataset. Those new principal components, which may be uncorrelated with each other, are reducing the complexity and noise of the original data. Principal component analysis in the cohort which can be separated by eight proteins. The first principal component is plotted on the X-axis and captures 34.9% of the variance. The second principal component is plotted on the Y-axis and achieves 15.6% of the variance. The scree plot represents the contribution of each principal component in PCA, and each principal component's contribution decreases sequentially.

Fig. 5 Principal component analysis (PCA). a The PCA mapping was performed using the Partek Genomics Suite (Partek, St. Louis, MO) (https://www.partek.com/partek-genomics-suite/). Patients with different survival status (red represents dead and blue represents alive) were separated by eight proteins. The first principal component is plotted on the X-axis and captures 34.9% of the variance. The second principal component is plotted on the Y-axis and achieves 15.6% of the variance. b The scree plot represents the contribution of each principal component in PCA, and each principal component's contribution decreases sequentially.
dataset while minimizing the loss of information. Technically, the number of new principal components can be equal to the number of variables in the original dataset. However, the contribution of new principal components, which represent the proportion of the original data, decrease sequentially (the first principal component accounts for most of the variability of the original dataset, the second subsequent component accounts for as much of the remaining variability as possible, ... until the last component). Therefore, only the first few principal components are the most representative, and this trend of progressively decreasing variability of each principal component can be visualized as the scree plot (Fig. 5b). This statistical approach to lower the dimensional representations within a data set through principal components is useful for the classification and the compression of a large dataset or big data.

**Ingenuity pathway analysis, gene-set enrichment analysis and circo**

Many analytical methods combined with online databases to analyze proteomics and microarray data, and are more suitable for discovering clinical significance rather than in-depth statistical analysis. Three commonly used computational tools are described and may be useful for some studies.

Ingenuity Pathway Analysis (IPA) was a web-based software application for causal analysis using expression datasets\textsuperscript{104}. It is now owned by Qiagen with >109,000 expression datasets and 8.5 million findings [https://digitalinsights.qiagen.com/products-overview/discovery-insights-portfolio/analysis-and-visualization/qiagen-ipa/]. It can generate hypothetical molecular interactions to understand cellular processes based on knowledge databases such as Biomolecular Interaction Network Database (BIND) database, Biological General Repository for Interaction Datasets (BioGRID) database, Cognia database, DIP database (Database of Interacting Proteins), IntAct database, Molecular INTeraction database (MINT), Munich Information Center for Protein Sequences (MIPS) database, QIAGEN's Ingenuity Knowledge Base, etc. Therefore, IPA can simultaneously visualize and analyze cross-database data of genomics, proteomics, and metabolomics data for signaling networks and canonical pathways from integrated various omics formats. The 2-dimensional signaling network offers a landscape survey of multi-omics (Fig. 6a) in which all upgraded and downgraded genes or proteins are visualized and connected or linked based on the latest database, and can be labeled by either function or pathway. The principle of ranking canonical pathways activity contains the research-based changes of each molecule such as the fold changes from PPA or microarray and the database-based the importance of each molecular in each canonical pathway, which is calculated with the Fisher's exact test as the negative log of this p value (Fig. 6b).

Gene set enrichment analysis (GSEA) ([https://www.gsea-msigdb.org/gsea/index.jsp](https://www.gsea-msigdb.org/gsea/index.jsp)) is another computational method that provides pathway enrichment tools to help interpret datasets\textsuperscript{105}. This approach focuses on cumulative changes in the expression of multiple genes as a gene set, which shares similar biological function, chromosomal location, or regulation, instead of an individual gene to identify pathways\textsuperscript{106}. One similar web-based GSEA tool is Enrich for analyzing human and mouse data\textsuperscript{107} and modEnrich for analyzing fish, fly, worm and yeast data\textsuperscript{108}.
most significant advantage of this GSEA method is that it can catch some pathways, in which several genes change in a small amount but in a coordinated way (Fig. 6c). The results reflect many of the complexities of co-regulation and modular expression by enrichment score (ES), corresponding to a weighted Kolmogorov–Smirnov-like statistic.

Additionally, Circos (http://circos.ca/) is a software package for visualizing omics-based data and information in a circular layout109. It has an online version (http://circos.ca/circos_online) which however was nonfunctional as of January 2022. Circos plot can be created for exploring relationships and contributions between canonical pathways and clinical clinicopathological category. The larger the circumference of the arc, the more active the canonical pathway or the more significant the influence of this clinicopathological category on the signaling network. The area of each colored ribbon delineates the proportion of the signaling pathway that contributes to a particular clinicopathological category.

**Fig. 7** Circos plot: Among all eight clinicopathological categories, the gender occupied the most significant proportion of the distribution, suggesting that it is the clinical factor that has the most impact on the signaling network. Among 20 canonical pathways altered in the disease, the HER-2 and p53 are affected most, suggested that they play essential roles in the pathogenesis of the disease.

**Single cell proteomics**

Single cell proteomics is an emerging technique focused on single cells. It will compete and complement single cell transcriptomics for understanding single-cell biology in the near future. Single cell proteomics recently became a reality when advanced technology
showed that peptides in a single cell could be efficiently delivered to the MS instruments\textsuperscript{10,11}. These single-cell MS methodologies can be broadly divided into cell-free and multiplex methods, the latter of which allows proteomic analyses of multiple cells at the same time. The SCope2 and Scp are the R-packages for analyzing single-cell proteomic data\textsuperscript{112,113}, while the SCeptre is their counterpart implemented in Python\textsuperscript{114}. Some general proteomic pipelines may also be used to process single-cell proteomic data. They include computational quality control tools\textsuperscript{115} and a single pipeline (MSNbase) for data processing and visualization\textsuperscript{116,117}.

In conclusion, during the last decade, proteomic technology and research has advanced tremendously. The increasing ability of high-throughput proteomics methods have generated real-time and in-depth datasets. The effective data mining technologies also significantly helped with the pursuit of novel and useful biomarkers, which are essential for disease early-detection and treatment. With the breakthrough of computing power and the rise of artificial intelligence, the role of proteomics has been further expanded. The highly advanced statistic/computational models enable proteomics to be integrated into multi-omics. Under this new trend, proteomics data analysis will be revolutionized for a bigger blueprint with a large amount of clinical and health-related data. It is an exciting time for proteomics developing into an essential new discipline and integrated with other disciplines. Although proteomics has to face emerging challenges during this process, it will move toward more in-depth single-cell biology and individualized precision medicine to boost both basic research and clinical practice to another level.

REFERENCES

1. Collins, FS, McKusick, VA. Implications of the Human Genome Project for medical science. \textit{JAMA} \textbf{285}, 540-544 (2001).
2. Wang, K, Huang, C, Nice, E. Recent advances in proteomics: towards the human proteome. \textit{Biomed Chromatogr} \textbf{28}, 848-857 (2014).
3. Mathivanan, S. Integrated Bioinformatics Analysis of the Publicly Available Protein Data Shows Evidence for 96% of the Human Proteome. J Proteomics \textbf{Bioinform 7}, 41-49 (2014).
4. Deutsch, EW, Lane, L, Overall, CM, Bandeira, N, Baker, MS, Pineau, C, et al. Human Proteome Project Mass Spectrometry Data Interpretation Guidelines 3.0. J Proteome \textbf{Res 18}, 4108-4116 (2019).
5. Wilhelm, M, Schlegel, J, Hahne, H, Gholami, AM, Lieberenz, M, Savitski, MM, et al. Mass-spectrometry-based draft of the human proteome. \textit{Nature} \textbf{509}, 582-587 (2014).
6. Ren, AH, Diamandis, EP, Kulasingam, V. Uncovering the Depths of the Human Proteome: Antibody-based Technologies for Ultra-sensitive Multiplexed Protein Detection and Quantification. \textit{Mol Cell Proteomics} \textbf{20}, 100155 (2021).
7. Aslan, B, Baist, M, Nisar, MA, Khorshed, M, Rasool, MH. Proteomics: Technologies and Their Applications. J Chromatogr Sci \textbf{55}, 182-196 (2017).
8. Kulasingam, V, Diamandis, EP. Strategies for discovering novel cancer biomarkers through utilization of emerging technologies. \textit{Nat Clin Pract Oncol} \textbf{5}, 588-599 (2008).
9. Huang, W, Zhan, D, Li, Y, Zheng, N, Wei, X, Bai, B, et al. Proteomics provides individualized options of precision medicine for patients with gastric cancer. \textit{Sci China Life Sci} \textbf{64}, 1199-1211 (2021).
10. Forler, S, Klein, O, Kloele, J. Individualized proteomics. \textit{J Proteomics} \textbf{107}, 56-61 (2014).
11. Uzozie, AC, Aebersold, R. Advancing translational research and precision medicine with targeted proteomics. \textit{J Proteomics} \textbf{189}, 1-10 (2018).
12. Parker CE, Borchers CH. The Special Issue: Clinical Proteomics for Precision Medicine. \textit{Proteomics Clin Appl} \textbf{12} (2018).
13. Macklin, A, Khan, S, Kisliger, T. Recent advances in mass spectrometry based clinical proteomics: applications to cancer research. \textit{Clin Proteomics} \textbf{17}, 17 (2020).
14. Li, X, Wang, W, Chen, J. Recent progress in mass spectrometry proteomics for biomedical research. \textit{Sci China Life Sci} \textbf{60}, 1093-1113 (2017).
15. Nilsson, T, Mann, M, Aebersold, R, Yates, JR, 3rd, Bairach, A, Meredith, J, Mass spectrometry in high-throughput proteomics: ready for the big time. \textit{Nat Methods} \textbf{7}, 681-685 (2010).
16. Peng, L, Cantor, DI, Huang, C, Wang, K, Baker, MS, Nice, EC. Tissue and plasma proteomics for early stage cancer detection. \textit{Mol Omics} \textbf{14}, 405-423 (2018).
markers: Platform performance in post-traumatic stress disorder and Parkinson's disease. Cytokine X 2, 100027 (2020).

44. Lim, SY, Lee, JH, Welsh, SJ, Ahn, SB, Breen, E, Khan, A, et al. Evaluation of two high-throughput proteomic technologies for plasma biomarker discovery in immunotherapy-treated melanoma patients. Biomark Res 5, 32 (2017).

45. Chowdhury, F, Williams, A, Johnson, P. Validation and comparison of two multiplex technologies, Luminox and Mesoscale Discovery, for human cytokine profiling. J Immunol Methods 340, 55-64 (2009).

46. Pan, J, Zheng, QZ, Li, Y, Yu, LL, Wu, QW, Zheng, JY, et al. Discovery and Validation of a Serologic Autoantibody Panel for Early Diagnosis of Esophageal Squamous Cell Carcinoma. Cancer Epidemiol Biomarkers Prev 28, 1454-1460 (2019).

47. Cui, L, Shu, C, Liu, Z, Tong, W, Cui, M, Wei, C, et al. The expression of serum sFLR, sFLR, and sFLR in preeclampsia. Pregnancy Hypertens 13, 127-132 (2018).

48. Cui, L, Shu, C, Liu, Z, Tong, W, Cui, M, Wei, C, et al. Serum protein marker panel for predicting preeclampsia. Pregnancy Hypertens 14, 279-285 (2018).

49. Tong, W, Ye, F, He, L, Cui, L, Cui, M, Hu, Y, et al. Serum biomarker panels for diagnosis of gastric cancer. Onco Targets Ther 9, 2455-2463 (2016).

50. Tanouchi, M, Verwey, JJ, Noor, Z, Sobuz, SU, Liezhout, L, Petri, WA, Jr., et al. High-throughput multiplex PCR and probe-based detection with Luminox beads for seven intestinal parasites. Am J Trop Med Hyg 84, 332-337 (2011).

51. Simpson, RJ & Greening, DW. Serum/plasma proteomics: methods and protocols. (Humana Press/New York, NY, USA, 2017).

52. Lantoine, J, Proces, A, Villers, A, Halliez, S, Buee, L, Ris, L, et al. Inflammatory Molecules Released by Mechanically Injured Astrocytes Trigger Presynaptic Loss in Cortico-Neuronal Networks. ACS Chem Neurosci 12, 3885-3887 (2021).

53. Jiang, Z, Kamerud, J, You, Z, Basak, S, Seletskaia, E, Steeno, GS, et al. Feasibility of plasma proteome. Proteomics 11, 1671-1772 (2021).

54. Rohloff, JC, Gelinas, AD, Jarvis, TC, Ochsner, UA, Schneider, DJ, Gold, L, et al. Nucleic Acid Ligands With Protein-like Side Chains: Modified Aptamers and Their Use as Diagnostic and Therapeutic Agents. Mol Ther Nucleic Acids 3, e201 (2020).

55. Sivakumaran, D, Ritz, C, Gjoen, JE, Vaz, M, Selvam, S, Ottenhoff, THM, et al. Host proteome. Proteomics 10, 3626-3635 (2010).

56. Fraser, DD, Chen, M, Ren, A, Miller, MR, Martin, C, Daley, M, et al. Novel Outcome Biomarkers Identified With Targeted Proteomic Analyses of Plasma From Critically Ill Coronavirus Disease 2019 Patients. Crit Care Explor 2, e0189 (2020).

57. Patel, H, Ashton, NJ, Dobson, RJU, Andersen, LM, Yilmaz, A, Blenow, K, et al. Proteomic blood profiling in mild, severe and critical COVID-19 patients. Sci Rep 11, 6357 (2021).

58. Carlson, AC, Ingelsson, E, Sundstrom, J, Carrero, JJ, Gustafsson, S, Feldtreich, D, et al. Use of Proteomics To Investigate Kidney Function Decline over 5 Years. Clin J Am Soc Nephrol 12, 1226-1235 (2017).

59. Mayer, SF, Cao, C, Dal, Peraro, M. Biological nanopores for single-molecule sensing. iScience 25, 104145 (2022).

60. Oppenheim, S, Cao, X, Rueppel, O, Krongard, S, Phokasem, P, DeSalle, R, et al. Whole Genome Sequencing and Assembly of the Asian Honey Bee Apis dorsata. Genome Biol Evol 12, 3607 (2020).

61. Zhou, A, Lin, X, Jing, J. Evaluating nanopore sequencing data processing pipelines for structural variation identification. Genome Biol 20, 237 (2019).

62. Yan, S, Zhang, J, Wang, Y, Guo, W, Zhang, S, Liu, Y, et al. Single Molecule Ratcheting Motion of Peptides in a Mycobacterium smegmatis Porin A (MspA) Nanopore. Nano Lett 21, 6703-6710 (2021).

63. Boskovic, F, Keyser, UF. Toward single-molecule proteomics. Science 374, 1443-1444 (2021).

64. Brinkerhoff, H, Kang, ASW, Liu, J, Aksimentiev, A, Dekker, C. Multiple reiterations of single proteins at single-amino acid resolution using nanopores. Science 374, 1509-1513 (2021).

65. Xu, J, Lin, Y, Yang, M, Zhang, L. Statistics and pitfalls of trend analysis in cancer research: a review focused on statistical packages. J Cancer 11, 2957-2961 (2020).

66. Krempel, R, Kulkarni, P, Yim, A, Lang, U, Habermann, B, Frommolt, P. Integrative analysis and machine learning on cancer genomics data using the Cancer Systems Biology Database (CancerSysDB). BMC Bioinformatics 19, 156 (2018).

67. Rashidi, HH, Tran, NK, Betts, EY, Howell, LP, Green, R. Artificial Intelligence and Machine Learning in Pathology: The Present Landscape of Supervised Methods. Acad Pathol 6, 2374289519873088 (2019).

68. Baxi, V, Edwards, R, Montalto, M, Saha, S. Digital pathology and artificial intelligence in translational medicine and clinical practice. Mod Pathol 35, 23-32 (2022).

69. Bera, K, Schalper, KA, Renner, DL, Velcheti, V, Madabhushi, A. Artificial intelligence in digital pathology - new tools for diagnosis and precision oncology. Nat Rev Clin Oncol 16, 703-715 (2019).

70. La Porta, CAM, Zappieri, S. Explaining the dynamics of tumor aggressiveness: At the crossroads between biology, artificial intelligence and complex systems. Semin Cancer Biol 53, 42-47 (2018).

71. Small-Tabbone, M, Rance, B, Section Editors for the IYSoS, Translation I. Contributions from the 2019 Literature on Bioinformatics and Translational Informatics. Yearb Med Inform 29, 188-192 (2020).

72. Stechow, von, L. Cancer Systems Biology: Methods and Protocols. (Humana Press: New York, NY, USA, 2016.).

73. Korenberg MJ. Microarray Data Analysis: Methods and Applications: Springer Science & Business Media, 2007.

74. Ward, JH. Hierarchical grouping to optimize an objective function. J. Am. Stat. Assoc. 51, 236-244 (1956).

75. MacArthur, BD, Lachmann, A, Lemischka, IR, Ma, X, Jones, G, Eissenberg, J, et al. Human homology and high-throughput proteomic technologies for plasma biomarker discovery in patients with end-stage renal disease. Diagnostics (Basel) 8, 2 (2018).

76. Raffield, LM, Dang, H, Pratte, KA, Jacobson, S, Gillenwater, LA, Ampleford, E, et al. Comparison of Proportion of Proteins Identified Between Four Methods in Multiple Cohort Studies. Proteomics 20, e1900278 (2020).

77. Fredriksson, S, Dixon, W, Ji, H, Koong, AC, Mindrinos, M, Davis, RW. Multiplexed protein detection by proximity ligation for cancer biomarker validation. Nat Methods 4, 327-329 (2007).

78. Gullberg, M, Gustafsdottir, SM, Schallmeiner, E, Jarvis, J, Bjarnegard, M, Betsholtz, C, et al. Cytochrome detection by antibody-proximity based ligation. Proc Natl Acad Sci U S A 101, 8420-8424 (2004).

79. Fredriksson, S, Gullberg, M, Jarvis, J, Olsson, C, Pietras, K, Gustafsdottir, SM, et al. Protein detection using proximity-dependent DNA ligation assays. Nat Biotechnol 20, 673-677 (2002).

80. Assarsson, E, Lundberg, M, Holmquist, G, Bjorksten, J, Thorsen, SB, Ekmann, D, et al. Homogenous 96-plex PEA immunoassay exhibiting high sensitivity, specificity, and excellent scalability. PLoS One 9, e95192 (2014).

81. Fraser, DD, Chen, M, Ren, A, Miller, MR, Martin, C, Daley, M, et al. Novel severe traumatic brain injury blood biomarkers identified with proximity extension assay. Clin Chem Lab Med 59, 1662-1669 (2021).

82. Fraser, DD, Cepinskas, G, Patterson, KD, Slessarev, M, Martin, C, Daley, M, et al. Novel Outcome Biomarkers Identified With Targeted Proteomic Analyses of Plasma From Critically Ill Coronavirus Disease 2019 Patients. Crit Care Explor 2, e0189 (2020).
100. Swinscow TDV, Campbell MJ. 

101. Roher, AE, Maarouf, CL, Sue, LI, Hu, Y, Wilson, J, Beach, TG. Proteomics-derived markers of autopsy-confirmed Alzheimer’s disease. Bio-markers 14, 493-501 (2009).

102. Li, J, Zhang, Z, Rimer, T, Wang, YY, Chan, DW. Proteomics and bioinformatics approaches for identification of serum biomarkers to detect breast cancer. Clin Chem 48, 1296-1304 (2002).

103. Keerthikumar, S & Mathivanan, S. Proteome Bioinformatics. (Humana Press:New York, NY, USA, 2017).

104. Kramer, A, Green, J, Pollard, J, Jr, Tugendreich, S. Causal analysis approaches in Ingenuity Pathway Analysis. Bioinformatics 30, 523-530 (2014).

105. Subramanian, A, Tamayo, P, Mootha, VK, Mukherjee, S, Ebert, BL, Gillette, MA, et al. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc Natl Acad Sci U S A 102, 15545-15550 (2005).

106. Zaim, SR, Li, Q, Schissler, AG, Lussier, YA. Emergence of pathway-level composite biomarkers from converging gene set signals of heterogeneous transcriptomic responses. Pac Symp Biocomput 23, 484-495 (2018).

107. Kuleshov, MV, Jones, MR, Rouillard, AD, Fernandez, NF, Duan, Q, Wang, Z, et al. Enrichr: a comprehensive gene set enrichment analysis web server 2016 update. Nucleic Acids Res 44, W90-97 (2016).

108. Kuleshov, MV, Diaz, JEL, Flamholz, ZN, Keenan, AB, Lachmann, A, Wojcikiewicz, ML, et al. modEnrichr: a suite of gene set enrichment analysis tools for model organisms. Nucleic Acids Res 47, W183-W190 (2019).

109. Krzywinski, M, Schein, J, Biau, I, Connors, J, Gascoyne, R, Horsman, D, et al. Circos: an information aesthetic for comparative genomics. Genome Res 19, 1639-1645 (2009).

110. Slavov, N. Scaling Up Single-Cell Proteomics. Mol Cell Proteomics 21, 100179 (2022).

111. Vistain, LF, Tay, S. Single-Cell Proteomics. Trends Biochem Sci 46, 661-672 (2021).

112. Petekski, AA, Emmott, E, Leduc, A, Huffman, RG, Specht, H, Perlman, DH, et al. Multiplexed single-cell proteomics using SCoPE2. Nat Protoc 16, 5398-5425 (2021).

113. Cheung, TK, Lee, CY, Bayer, FP, McCoy, A, Kuster, B, Rose, CM. Defining the carrier proteome limit for single-cell proteomics. Nat Methods 18, 76-83 (2021).

114. Schoof, EM, Furtwangler, B, Uresin, N, Rapin, N, Savickas, S, Gentil, C, et al. Quantitative single-cell proteomics as a tool to characterize cellular hierarchies. Nat Commun 12, 3341 (2021).

115. Bittremieux W, Valkenburg D, Martens L, Laukens K. Computational quality control tools for mass spectrometry proteomics. Proteomics 17, (2017).

116. Gatto, L, Gibb, S, Rainer, J. MSnbase, Efficient and Elegant R-Based Processing and Visualization of Raw Mass Spectrometry Data. J Proteome Res 20, 1063-1069 (2021).

117. Gatto, L, Breckels, LM, Naake, T, Gibb, S. Visualization of proteomics data using R and bioconductor. Proteomics 15, 1375-1389 (2015).

AUTHOR CONTRIBUTIONS
M.C. and L.Z. drafted the manuscript. All authors reviewed, discussed, and edited the manuscript.

FUNDING
This work is in part supported by National Science Foundation (IIS-2128307 to L.Z.) and Cancer Prevention & Research Institute of Texas (RR180061 to C.C.). The funders have no role in writing this work or the decision to submit it for publication.

COMPETING INTERESTS
The authors declare no competing interests.

ADDITIONAL INFORMATION
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41374-022-00830-7.

Correspondence and requests for materials should be addressed to Chao Cheng or Lanjing Zhang.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Springer Nature or its licensor holds exclusive rights to this article under a publishing agreement with the author(s) or other rightholder(s); author self-archiving of the accepted manuscript version of this article is solely governed by the terms of such publishing agreement and applicable law.