Pathway mapping and development of disease-specific biomarkers: protein-based network biomarkers

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Abstract
It is known that a disease is rarely a consequence of an abnormality of a single gene, but reflects the interactions of various processes in a complex network. Annotated molecular networks offer new opportunities to understand diseases within a systems biology framework and provide an excellent substrate for network-based identification of biomarkers. The network biomarkers and dynamic network biomarkers (DNBs) represent new types of biomarkers with protein-protein or gene-gene interactions that can be monitored and evaluated at different stages and time-points during development of disease. Clinical bioinformatics as a new way to combine clinical measurements and signs with human tissue-generated bioinformatics is crucial to translate biomarkers into clinical application, validate the disease specificity, and understand the role of biomarkers in clinical settings. In this article, the recent advances and developments on network biomarkers and DNBs are comprehensively reviewed. How network biomarkers help a better understanding of molecular mechanism of diseases, the advantages and constraints of network biomarkers for clinical application, clinical bioinformatics as a bridge to the development of diseases-specific, stage-specific, severity-specific and therapy predictive biomarkers, and the potentials of network biomarkers are also discussed.

Keywords: network biomarkers • dynamic network biomarkers • protein-protein interactions • disease • diagnostics

Introduction
The disease consists of multiple dysfunctional proteins, cells, organs and systems of the body within the complexity and molecular mechanisms by which diseases occur remain unclear, although biotechnologies and knowledge on diseases have been improved tremendously.

Given the functional interdependencies between the molecular components, a disease is rarely a consequence of an abnormality of a gene, protein or cell, but reflects the interactions of genes, proteins or cells in a complex network [1]. Protein-protein interactions (PPIs)
are important in the interaction, communication and functional process in a living cell, between cells, or between organs. PPI networks are of central importance to modulate cell behaviour by interactively link genome, epigenome, transcriptome, proteome and metabolome. In response to the dynamically varying intrinsic (genetic) and extrinsic (environmental) perturbations, the interplay of these interconnected cellular signalling networks can converge towards disease states and ultimately can initiate and drive complex diseases (Fig. 1). Annotated data sets of PPIs offer new opportunities to understand diseases within a systems biology-based framework and provide a useful substrate for network-based identification and validation of multiple interacting markers [2, 3]. The PPIs are curated from the literature into databases [4], to improve the understanding of diseases and provide the basis for new therapeutic approaches.

Biomarkers are typically thought of as individual genes, proteins and metabolites (molecular biomarkers). However, with the recent innovation and progress of new biotechnologies, a new type of biomarkers, protein-based network biomarkers [5, 6], composed of a panel of proteins and their interactions or interactions with DNA, RNA or other molecules was initiated and investigated with the integration of knowledge on protein annotations, interactions and signalling pathways. But both the molecular biomarkers and the protein-based network biomarkers have limitations because of their static nature. To increase the ability to make early diagnosis, identify disease-specific biomarkers and therapeutic targets, and predict patient outcome, dynamic network biomarkers (DNBs) were created, monitored and evaluated at different stages and time-points of the disease, based on non-linear dynamical theory and complex network theory. DNBs differ from molecular biomarkers and network biomarkers to describe and identify disease progress situations and interactions rather than the static nature and approach. Thus, DNBs can demonstrate the expression density of genes or proteins and their time-dependent interactions to draw a three-dimensional imaging of altered proteins, interactions or regulations in the network and to discover and develop disease-specific biomarkers to predict and foresee pre-disease situations, monitor and regulate therapeutics in clinic, and indicate and guide of patient prognosis and life quality [7, 8].

Furthermore, the human biomarker discovery mandates the association of biological measurements with clinical information, ideally both statistically and mechanistically. Unlike biological data, clinical data, such as patient complaints, history, therapies, clinical symptoms and signs, physical examinations, biochemical analyses, imaging profiles, pathologies and other measurements, are descriptive and far less structured. The lack of integrative results in the loss or neglecting of valuable information, necessitates novel strategies to successfully combine these large collections of heterogeneous data sets, and identifies new disease-specific biomarkers. To address this issue, clinical bioinformatics was proposed as a new emerging science to combine clinical phenotypes with human tissue-generated bioinformatics and define relationships between collectively directs global function with clinical measurements [9, 10]. Comparing dynamic alterations of network biomarkers with clinical informatics may allow discovering disease-specific, stage-specific, severity-specific and therapy-sensitive biomarkers.

The present review aims to introduce the need and significance of protein-based network biomarkers, highlight the development of network biomarkers of human diseases, and discuss the clinical relevance and correlation between DNBs and clinical informatics. We will explore how network biomarkers help a better understanding of molecular mechanism of diseases, the advantages and constraints of network biomarkers for clinical application, and the potential values of network biomarkers in the future.

Biomarkers play an important role in the diagnosis of diseases, and in assessing prognosis and directing treatment of the diseases. Its value
is correlated with the disease-associated specificity, sensitivity, trace-ability, stability, repeatability and reliability. Conventional molecular biomarkers consist of single or a group of several biological mole-cules such as genes, RNAs, proteins and metabolites that can be measured to distinguish disease from health. Advances in high-throughput technologies such as genomics and proteomics make it possible to measure thousands of different variables in pathobiologi-cal conditions simultaneously, providing comprehensive and substan-tial information of a disease state. An increasing number of biomarkers identified through analysis of expression profiles have been seen. Unfortunately, this type of biomarker lists that obtained from omics data for similarly diagnosed patients by different research groups differ widely and share few in common [11]. As biomarkers are required to be reproducible to causally link to the disease pheno-type to discover potential targets for diagnosis and therapy, this lack of agreement imposed doubts about the reliability and robustness of the reported biomarker lists.

The small overlap of biomarkers for similar phenotypes may have various technique reasons, such as platforms differences, samples differences, protocols differences, and statistical reasons, leading to unstable selection of the biomarkers. Besides, identification of a phe-notype-associated pathway solely on the basis of differentially expressed molecules is frequently difficult because of the high inter-dependency of the omics data. But more importantly, these biomar-kers are not identified from the systems perspective. With the rapid growing knowledge of the cellular molecular network in diseases, most of the diseases are not considered to be caused by a single effector gene product but the interrelated malfunction of genes and proteins [1, 12–14]. This system-level understanding of the diseases has brought about novel strategies of biomarker discovery that inte-grate systemic information of the molecular networks (i.e. PPI net-works, RNA networks, metabolic networks and regulatory networks) to contextualize the differential expression patterns observed in a phe-notype. The system-based ‘network biomarkers’ [5, 6] was proposed to consider not only differentially expressed molecules but also the molecules association network structure that even allows an accurate identification of biomarkers with low discriminative potentials provided such molecules were associated with other significant mole-cules. The past few years have witnessed systematic efforts to inte-grate gene network knowledge in the gene expression analysis [15–19]. However, a clearly limitation of gene network analysis is that genes are not the proper end-point context of a phenotype. Moreover, the majority of human genes have not yet been assigned to a definitive pathway. Proteomic profiling provides information at the post-translational level, therefore can be used to bridge the genotype-phenotype gap, to help providing a global picture of cellular mecha-nisms. With the recent significant improvements of mapping human protein networks, networks approach have been studied in proteo-mic researches to understand disease-related pathobiological pro cesses and to identify candidate disease biomarkers [6, 7, 20–24].

Studies have also integrated multiple data types to generate more accurate molecular networks of diseases [5, 25–46] and revealed the dynamic modular structure of the protein interaction networks [47–49]. Integration of condition-specific co-expression information can provide clues to the dynamic features of these networks, when PPI data constitute static network maps – such knowledge-integrated interaction is relatively defined and fixed. The ‘DNBs’ is an innovative concept to integrate network biomarkers and dynamic biomarkers by monitoring and evaluating the alterations of network biomarkers at different stages and time-points. Unlike static molecular biomarkers and network biomarkers with consistent values, DNB is a group of molecules, which are highly fluctuating but strongly correlated without consistent values during disease progression [7]. DNB is a pow-erful way to detect the bifurcation of gene or protein interactions to unravel the dynamic aspects of cellular networks and answer when, where and why proteins interact, and to indicate the early change in biomarkers and to predict the occurrence of diseases [50, 51].

Chen et al. [7] proposed to consider a disease progress as three stages, including the normal stage, the pre-disease stage and the dis-ease stage. The normal stage is a relatively healthy state, as well as the chronic inflammation period or the period that the disease is under control. The pre-disease stage [52–55] is a state just before the presence of disease symptoms, defined as the limit of the normal state immediately before the critical point is reached. This stage is usually reversible to the normal state if appropriately treated, or becomes irreversible if the system passes over the critical point to the disease stage. Therefore, detection of the pre-disease stage is crucial to achieve early diagnose and treatment. Rather as a wide, general definition been discussed before, the authors developed DNB in particular as an early warning indicator of pre-disease state and to fulfill three criteria [7]: firstly, DNB is an observable subnetwork of the system, composed of a group of molecules that are dynamically cor-related when the system is in a pre-disease state; secondly, DNB is an isolated subnetwork or functional module, behaves independent of other non-DNB molecules that are in the same system or network; and thirdly the expressions of DNB molecules increasingly fluctuate as the system is approaching the critical point. Based on these condi-tions, a composite index I was constructed to computationally identify DNB from high-throughput omic data:

$$I = \frac{SD_d \cdot PCC_d}{PCC_o}$$

where $PCC_o$ is the numerical measurement of the average Pearson’s correlation coefficient among the molecules of DNB in absolute value; $PCC_d$ is the numerical measurement of the average Pearson’s correlation coefficient of the molecules of DNB with the other molecules in absolute value; $SD_d$ is the numerical measurement of the average standard deviation of the molecules of DNB. The composite index I was shown to be effective to provide reliable and significant early warning signal, despite the stochastically fluctuation in the expression of each molecule, in complex diseases such as acute lung injury, liver cancer and lymphoma [7]. Furthermore, DNB was demonstrated as the leading/driving network causally related to disease initia-tion and progression [8].

Figure 2 summarizes the evolution of these three types of biomar-ker concepts, namely, molecular biomarkers that provide static, one-dimensional information (Fig. 2A), network biomarkers that provide static, two-dimensional profiles (Fig. 2B), and DNBs that provide
dynamic, three-dimensional images of biomarker–biomarker interactions (Fig. 2C).

The development of protein-based network biomarkers

Protein–protein interactions play major role in a living system, provide information at the post-translational level, and bridge the genotype–phenotype gap. Annotated data sets of PPIs provide key substrate for network-based identification of biomarkers. The systematic approach of protein-based network biomarker discovery typically involves three pivotal processes: (i) obtaining global expression profiles of a disease phenotype; (ii) integrating such information into protein network frameworks or literature-curated pathways that contain key pathobiological events of phenotypes and (iii) interpreting, identifying and validating phenotype-associated candidate key network molecules or modules (Fig. 3). This section mainly introduces some foundations of protein-based network biomarker discovery and demonstrates several representative studies in human diseases and how such network biomarkers shed light onto the molecular mechanism of diseases.

Human protein–protein interaction network

Human PPI network mapping is a crucial component of systematic approach for protein-based network biomarker discovery. Extensive efforts have been seen recently to increase the coverage of human PPI maps by high-throughput yeast two-hybrid (Y2H) interaction mating [2, 3, 56–58], affinity purification mass spectrometry (AP-MS) [59], literature curation of published experiments [60–70] or computational integrating approaches [71–76]. Some of the major public PPI databases are summarized in Table 1. The global efforts to map protein interactions with curated interactions from the literature also has resulted in the formation of the International Molecular Exchange (IMEx) consortium (http://www.imex consortium.org/) which can facilitate literature curation standards, data exchange and comparison. However, considering the full human protein interaction network has been estimated to be between 154,000 and 369,000 [77] or approximately 650,000 [78], current human PPI maps are still incomplete, noisy and prone to biases [74, 79], caution should be exercised when using them to research diseases. Given the magnitude of this challenge, a number of strategies have been proposed to maximize the efficiency and correct biases for PPI network mapping [56–58, 80].

Methodologies for integrating and identifying network biomarkers

A disease can be linked to a well-defined neighbourhood of PPI network, which refers to as ‘disease module’, representing a panel of network components responsible for cellular function and disruption of which results in a specific disease phenotype [1]. Disease-related profiles can be integrated within a network framework by a number of technologies and algorithms [81, 82]. Computational programs were developed to integrate selected genes or proteins into the knowledge-based networks via the combination of genomics, proteomics and bioinformatics, such as GRNInfer [83], MDCinfer [84], TRNInfer [85], Samo [86], MNAAligner [87], PTG [88], PRNA [89], NOA [90], differential dependency network (DDN) [91], WGCNA [92], SurvNet [93] or DiME [94], each of them has its own advantages and strength on basis of scientific needs and investigative goals, as summarized in Table 2.

Methodologies for expression data integration could be categorized as ‘univariate’ or ‘multivariate’ on basis of statistics [95], to interpret the dysregulation (differential expression) at the system
Defining the subnetworks dysregulation as the aggregate significance of the dysregulation of each gene, the univariate approaches measure the dysregulation of subnetworks by combining the results of differential expression of each gene that are assessed separately [47, 49, 96–99]. However, multivariate studies consider the dysregulation of the subnetworks as the mutual information between phenotype and subnetworks activity, and access the coordination of multiple gene differential expressions in discriminating normal and disease samples [99]. The coordinate dysregulation was shown to be effective integrating protein and mRNA expression data for identification of important subnetworks in colorectal cancer [30, 33]. The coordination of subnetwork dysregulation could be captured by ‘additive’ or ‘combinatorial’ formulations. The additive dysregulation formulates the coordination between genes through the additive representation of their expression profiles and utilizes fast heuristics to identify dysregulated subnetwork; while the combinatorial formulation assesses the combining degrees of gene expressions in the subnetwork that can discriminate control and phenotype samples. The combinatorial approach was shown to be powerful in predicting liver metastasis in human colorectal cancer [100].
is not explained by smaller parts of the subnetwork, the synergistic dysregulation was formulated to delineate the complementarity of genes in the subnetwork [101]. Synergy corrects for the coordinate dysregulation of the subsets of the subnetwork, capturing the pattern of dysregulation that emerges only when all genes in the subnetwork are considered. Although identification of multiple genes with synergistic dysregulation is intractable, this methodology provides important insight through pair-wise assessment of synergy, which jointly analyses two sets of expression data, one in the presence and one in the absence of a disease, identifying gene pairs whose correlation with disease is because of co-operative, rather than independent, contributions of genes [102]. Besides coordinate dysregulation, differential co-expression approaches are also shown to be effective in finding co-expressed genes in the disease samples, rather than controls, and vice versa [103, 104].

Briefly, disease-related molecules identified from omic profiling studies or other sources, which serve as the ‘seeds’ of the disease module, are placed on their corresponding proteins in the properly constructed PPI network, and by exploiting both the functional and topological modularity of the network through quantitative systematic approaches, subnetworks and/or pathways with the disease-related components could then be identified as disease module, and as potential network biomarker. Numerous computational methods and algorithms have been proposed for network biomarker identification. For instance, the DDN analysis [91, 105], which detects topological changes in biological networks by comparing the topological differences between networks, is a straightforward way in distinguishing disease samples when the topology of disease networks is significantly different from the topology of normal samples. However, the network structure learning is inconsistent because of the limitation of the data samples, making this approach not convenient in real applications. The active subnetwork identification approach [5, 106] identifies disease module as active subnetworks that show significant changes in particular conditions by using existing PPI networks. While this approach can identify disease-related subnetworks that are not differentially expressed, it is limited by the availability of the PPI networks. The disease-specific pathway identification method [37, 73, 107] is another systemic approach to extract disease-specific subnetworks or pathways by using regression models or scoring modules. This approach is effective in identifying network biomarkers based on the integration of PPIs and pathway knowledge. However, it is limited by the exhaustive search procedure. Besides, caution should be excised that the regression model-based method is not suitable for small sample cases, in which the parameters are biased. The information flow modelling approach [108, 109] identifies dysfunctional modules in complex disease by modelling the information flow from source disease genes to targets of differentially expressed genes via a context-specific PPI network. This approach is effective in

| Database      | Method                  | Proteins number | Interactions number | Ref.      | Website                                      |
|---------------|-------------------------|-----------------|---------------------|-----------|----------------------------------------------|
| CCSB          | Y2H                     | 1549            | 2754                | [2]       | http://interactome.dfci.harvard.edu/         |
| MDC           | Y2H                     | 1705            | 3186                | [3]       | http://www.mdc-berlin.de/neuroprot/database.htm |
| BIND          | Literature              | 6089            | 14955               | [60]      | http://bind.ca                               |
| DIP           | Literature              | 3877            | 6103                | [62]      | http://dip.doe-mbi.ucla.edu/dip              |
| MINT          | Literature              | 8762            | 26830               | [63]      | http://mint.bio.uniroma2.it/mint             |
| HomoMINT      | Prediction & Literature | 8634            | 323595              | [70]      | http://mint.bio.uniroma2.it/HomoMINT         |
| InteAct       | Literature              | 60932           | 197974              | [64]      | http://www.ebi.ac.uk/intact/                |
| BioGRID       | Literature              | 18208           | 220390              | [65]      | http://thebiogrid.org/                      |
| HPRD          | Literature              | 30047           | 41327               | [66]      | http://hprd.org/                            |
| Reactome      | Literature              | 7085            | 6744                | [68]      | http://www.reactome.org/                    |
| UniHI         | Database integration    | 36023*          | 374833*             | [74]      | http://www.unihi.org/                       |
| HAPPI         | Database integration    | 70829           | 601757              | [75]      | http://bioinformatics.iupui.edu/HAPPI       |

The table displays the number of proteins and the number of interactions derived from each database. The column termed Methods shows the general approaches how PPIs were compiled in the different resources. In addition, the literature reference for the resource and the websites of the databases are given.

CCSB: Center for Cancer Systems Biology; MDC: Max Delbrück Center; BIND: Biomolecular Interaction Network Database; DIP: Database of Interacting Proteins; MINT: Molecular INTeraction database; HomoMINT: inferred human network of MINT; InteAct: the protein interaction database; BioGRID: Biological General Repository for Interaction Datasets; HPRD: Human Protein Reference Database; Reactome: a curated pathway database; UniHI: Unified Human Interactome; HAPPI: Human Annotated and Predicted Protein Interaction.

*All species.
classifying the functional dependency or crosstalk between pathways provided differentially expressed genes are detected. Unlike the conventional clustering approaches based on differential expressions, classification of differential interactions [110, 111] investigates the differential interactions between disease and normal samples and network rewiring between molecules related to pathogenesis. Applied to

Table 2 Examples of software packages for mapping phenotype-related subnetworks

| Name      | Full name                                      | Website                                      | Description                                                                 | Ref. |
|-----------|------------------------------------------------|----------------------------------------------|-----------------------------------------------------------------------------|------|
| GRNInfer | Gene Regulatory Network Inference Tool         | http://digbio.missouri.edu/grninfer/         | A gene regulatory network inference tool from multiple microarray data sets  | [83] |
| MDCinfer | Inferring protein–protein interactions based on multi-domain Co-operation | http://intelligent.eic.osaka-sandai.ac.jp/chenen/MDCinfer.htm | PPI prediction tool based on multiple domain co-operation analysis         | [85] |
| TRNInfer | Inferring transcriptional regulatory networks from high-throughput data | http://intelligent.eic.osaka-sandai.ac.jp/chenen/TRNinfer.htm | Infer direct relationships between transcription factors and target genes    | [85] |
| Samo      | Protein Structure Alignment tool based on Multiple Objective optimization | http://doc.aporc.org/wiki/Samo | A protein structure alignment tool based on multiple objective optimization | [86] |
| MNAAligner | Molecular Network Aligner                      | http://doc.aporc.org/wiki/MNAAligner        | Alignment of molecular networks by quadratic programming                   | [87] |
| PTG       | Parsimonious Tree-Grow method for haplotype inference | http://doc.aporc.org/wiki/PTG | Parsimonious tree-grow method for haplotype inference                      | [88] |
| PRNA      | Protein–RNA Binding-Site Prediction            | http://doc.aporc.org/wiki/PRNA              | Prediction of protein–RNA binding sites by a random forest method with combined features | [89] |
| NOA       | Network Ontology Analysis                      | http://www.aporc.org/noa/                   | Collection of gene ontology tools aiming to analyse functions of gene network instead of gene list | [90] |
| DDN       | Differential dependency network analysis       | http://www.cbil.ece.vt.edu/software.htm      | Detect statistically significant topological changes in the transcriptional networks between two biological conditions | [91] |
| WGCNA     | Weighted correlation network analysis          | http://www.genetics.ucla.edu/labs/horvath/CoexpressionNetwork/Rpackages/WGCNA | A comprehensive collection of R functions for performing various aspects of weighted correlation network analysis patterns among genes across microarray samples | [92] |
| SurvNet   | N/A                                             | http://bioinformatics.mdanderson.org/SurvNet | A bioinformatics web app for identifying network-based biomarkers that most correlate with patient survival data | [93] |
| DiME      | Disease Module Extraction                      | www.cs.bham.ac.uk/~szh/DiME                | A novel algorithm based on the Community Extraction criterion, to extract topological core modules from biological networks as putative disease modules | [94] |

The table displays the abbreviated name and full name of the computational programs with respective description and website. In addition, the literature reference for the resource is given.
gastric cancer, this method demonstrated that the differential interactions are effective on identifying dysfunctional modules from the molecular interaction network and can be applied as network biomarkers [111]. However, this approach is time consuming because of the large-scale interaction networks. The supporting vector machine (SVM) approach [35, 112, 113] identifies a comprehensive key interaction map and integrating different types of interaction information of heterogeneous data sources within the SVM scheme. By using various biological knowledge and data sources such as gene co-expression, regulatory networks, evolutionary relationship and functional similarity, the effectiveness and efficiency are significantly improved. The major shortcoming of SVM is its high computational cost for real applications.

Network biomarker studies in humans

Variants of aforementioned approaches have been applied to a wide range of disease phenotypes for identifying protein-based network biomarkers, such as breast cancer [5, 35, 36, 49], colorectal cancer [30, 33, 100], prostate cancer [97, 114], gastric cancer [115], lung cancer [37], ovarian cancer [116], acute myeloid leukaemia [39], glioma [117], ageing [23, 98], Alzheimer’s disease [38], inherited ataxias [118], cardiovascular diseases [6, 32, 34, 109, 119, 120], chronic obstructive pulmonary disease [22], diabetes [43], asthma [121], osteoarthritis [42, 122], multiple sclerosis [29], primary immunodeficiency [31], systemic inflammation [25], or acute aortic dissection [123], and demonstrated promising results. For example, Jin et al. [6] developed a cardiovascular-related network based on protein information databases, and discovered network biomarkers of major adverse cardiac events (MACE) in the MS data on basis of protein knowledge. Candidate network biomarkers could classify patients with MACE more accurately than current single ones without network information. Similarly, Lim et al. [118] developed a PPI network of 54 proteins involved in 23 inherited ataxias, and expanded the network by incorporating literature-curated and evolutionarily conserved interactions. This phenotype-based PPI network reveals several previously unsuspected ataxia-causing proteins interactions and provides candidate genes for target-based therapies.

Integrating multiple types of data sources could enhance the accuracy of the network and improve the quality of identified disease-specific biomarkers [38, 108]. Chuang et al. [5] integrated the gene expression and PPI network data sets to identify biomarkers associated with breast cancer metastasis. The gene expression profiles of two cohorts of breast cancer patients were obtained from literatures, assigned as either ‘metastatic’ or ‘non-metastatic’, and a corresponding human PPI network was constructed by integrating data sets from Y2H, orthology and literature curation experiments. The expression values of each gene were then mapped onto their corresponding proteins in the network to combine the gene and protein data sets. The discriminative potential of candidate subnetwork was computed based on the mutual information between its activity score and the metastatic/non-metastatic disease status over all patients and the significantly discriminate subnetworks were identified by comparing their discriminative potentials to those of random networks. The results showed that known breast cancer genes such as P53, KRAS, HRAS, HER-2/neu and PIK3CA that do not change their expression profile might still play a central role interconnecting genes in the protein network. The identified subnetworks may be more reproducible than individual gene marker selected by traditional approaches, and be better to define metastatic tumours. Systems-based approaches were also used to identify novel biomarkers and understand related mechanisms in a more comprehensive way by integrating protein network with data types or sources, such as phenome [28, 40], microRNA [42] or mRNA [33].

The network biomarkers have also been investigated dynamically. Taylor et al. [49] examined the dynamic structure of human protein interaction network by analysing ‘intermodular’ or ‘intramodular’ hub proteins that are co-expressed with their interacting partners in a tissue-restricted manner or in all or most tissues. Substantial differences in biochemical structure were observed between the two types of hubs. Hub proteins that displayed dynamic modularity were useful indicators for predicting the outcome of patients with breast cancer. Similar observations were noticed by other groups in the yeast [47, 48]. Lin et al. [119] proposed a network-based approach that integrates PPIs with gene expression profiles and biological function annotations to analyse the interaction networks in different biological states. They found that hub proteins in condition-specific co-expressed PPI networks tended to be differentially expressed between biological states. Applying this method to a cohort of heart failure patients, they identified two functional modules that significantly emerged from the interaction networks that can provide new insights into the cause of dilated cardiomyopathy and might be used as potential drug targets. The dynamic features of network biomarker were also investigated in ageing [98], liver cancer [124], breast cancer [107], glioma [105] and influenza [125].

Table 3 summarized some representative network biomarker studies in humans. While the results seemed promising, these studies are method and algorithms oriented; apparently, there is a great need to validate these approaches according to clinical application.

Better understanding of molecular mechanism of diseases

The integration of disease-specific molecules into the knowledge-based protein networks and subnetworks is a new and better way to understand mechanistic hypotheses about the causes of disease. The interactions within such subnetworks are often suggestive of functional signalling cascades, metabolic pathways or molecular complexes responsible for or/and contributing to the phenotypes and dysfunction of the disease. Thus, the network approach offers a potent means of mapping the molecular mechanisms underlying complex pathobiological processes. While the networks of genes and proteins present the links and association between them, such knowledge-integrated interaction is relatively defined and fixed. Rather than only the expression, the strength of interactions between genes or proteins varies during the development of diseases. Moreover, DNB is proved to be the leading network to initiate the critical transition during disease progression, and is highly related to causal factors of
| Disease       | Seeds                                                                 | PPI data sources                                                                 | Ref. | Key findings                                                                                                                                 |
|--------------|----------------------------------------------------------------------|----------------------------------------------------------------------------------|------|----------------------------------------------------------------------------------------------------------------------------------------------|
| Breast cancer| Microarray data of two cohorts of breast cancer patients from literature | Database integration of Y2H, prediction and literature curation, includes 11203 proteins and 57235 interactions | [5]  | The identified subnetwork biomarkers increased the reproducibility and accuracy in differentiating metastatic from non-metastatic breast tumours, compared to traditional molecular biomarkers. |
| Colorectal cancer | Two sets of proteomic targets of colorectal cancer obtained from tissue biopsies | HPRD database includes 9299 proteins and 35023 interactions | [33] | Integration of complementary data sources can enhance the discovery of candidate subnetworks in cancer that are well-suited for mechanistic validation in disease. |
| Prostate cancer | A prostate gene data set from literature | Database integration of DIP and HPRD, includes 6509 proteins and 23157 interactions | [97] | The proposed approach can discover condition relevant functional modules efficiently. |
| Breast cancer | Microarray data of two cohorts of breast cancer patients from literature | Database integration of HPRD database and IPA, includes 584 genes and 2280 interactions | [35] | The identified network biomarkers are highly enriched in biological pathways associated with cancer progression and the prediction performance is much improved when tested across different data sets. |
| Prostate cancer | 67 proteins identified from tumour tissue of a cohort of colorectal cancer patients | MetaCore from GeneGo Inc. (version 4.6 build 12332) | [30] | The identified protein subnetwork biomarkers can discriminate late stage cancer versus control. Proteins in the subnetworks were associated with the tumour progression or identification. |
| Disease               | Seeds                                      | PPI data sources                                      | Ref.   | Key findings                                                                 |
|----------------------|--------------------------------------------|-------------------------------------------------------|--------|------------------------------------------------------------------------------|
| Gastric cancer       | 272 differentially expressed genes in the metastatic gastric cancer | UniHI database                                       | [115]  | The identified subnetwork biomarkers are promising diagnostic markers for liver metastasis of gastric cancer |
|                      | Gene expression data from GEO database (ID: GSE27342) | HPRD database includes 9465 proteins and 37039 interactions | [111]  | Identified network biomarkers include 34 genes shown to be directly connected by the gastric cancer-related genes to all phases, and a functional transition from normal phenotypes to cancer phases was demonstrated |
| Lung cancer          | Microarray data from GEO database (ID: GSE4115) | Database integration of BioGRID and HPRD              | [37]   | Identified 40 proteins related to lung carcinogenesis. The network-based biomarker was effective in diagnosing smokers with signs of lung cancer |
| Cardiovascular disease | Mass spectrometry data from major adverse cardiac events patients | HPRD includes 18796 proteins and 37056 interactions | [6]    | The identified network biomarkers can classify the patients with major adverse cardiac events more accurately than traditional molecular biomarkers |
|                      | 105 heart failure associated proteins from the literature | Database integration of HPRD, BioGRID and MINT        | [32]   | The identified network biomarkers support accurate prediction of heart failure and provide novel clue to the underlying mechanisms |
|                      | Known inflammation biomarkers from clinical practice and literature | Database integration of DIP, IntAct and MINT         | [34]   | Identified a panel of gene biomarkers with high discriminatory capability predicts clinical outcome after myocardial infarction |
|                      | CHD microarray data from GEO database (ID: GSE26125 & GSE14790) | Database integration of HPRD, BIND, BioGrid, IntAct and MINT, includes 4761 proteins and 18084 interactions | [109]  | Identified 12 dysfunctional modules from the constructed CHD subnetwork, which provide clue to molecular mechanisms of CHD |
the disease [8]. In this regard, DNBs, by integrating of condition-specific information of network biomarkers at different stages and time-points, promise an improved understanding of the causes of human disease [8, 49, 119].

Xue et al. [98] examined the modular structure of the protein interaction networks during the ageing of fruitfly and human brains and found two modules associated with the cellular proliferation to differentiation temporal switch that display opposite ageing-related changes in expression. This particular study provides a modularized network view of the ageing process and found the dynamic network stability might be associated with the ageing. Such a dynamic network view provides a molecular explanation for the stochastic nature of ageing, that is, isogenic population age at vastly different paces, for the states of the network can be differentially affected by developmental and environmental factors. Li et al. [126] constructed dynamic physical and functional protein interactions network regulating the production of type I interferon (IFN) and identified 22 unique genes that regulated NF-kB and ISRE reporter activity, viral replication or virus-induced IFN production. Among them, mind bomb (MIB) E3 ligases played a role in K63-linked ubiquitination of TBK1 kinase that phosphorylates IRF transcription factors controlling IFN production. MIB genes were found selectively controlled responses to cytosolic RNA viruses, and MIB deficiency reduced antiviral activity. This study established the role of MIB proteins as positive regulators of antiviral

The table displays biomarkers studies in humans with respective network approach description. In addition, the literature reference for the resource is given.

GEO: Gene Expression Omnibus; OMIM: Online Mendelian Inheritance in Man database; IPA: Ingenuity Pathway Analysis; MetaCore: Data-mining and pathway analysis (http://thomsonreuters.com/metacore/); CHD: Congenital heart disease; OPHID: Online Predicted Human Interaction Database; I2D: Interologous Interaction Database; MYC: myelocytomatosis oncogene.
responses and demonstrated that mapping a dynamic physical and regulatory network of type I IFN can be a valuable source for understanding the connections between innate immunity and other cell processes.

Based on the computational algorithm (composite index J) described above, researchers found two dynamical network biomarkers which can be separately used to predict the peri-insultis of the early stage of disease and the onset of type 1 diabetes with overt hyperglycaemia [127]. These two DNBs were adopted to analyse and revealed that mitochondrion electron transport induces the apoptosis function of the second DNB and pushes the peri-insultis to diabetes. Li et al. [128] identified tissue-specific DNBs corresponding to the critical transitions occurring in liver, adipose and muscle during type 2 diabetes mellitus (T2DM) progression, and found two different critical states during T2DM development, characterized as responses to insulin resistance and serious inflammation respectively. The identified DNB genes are significantly associated with T2DM, either to be the disease genes or participate in important biological processes related to the T2DM development, such as response to insulin stimuli, abnormal lipid metabolism and immune system response. DNB genes were also found tend to be located at the upstream of pathway rather than the consequence so that DNB genes act as the causal factors to drive the downstream molecules to change their transcriptional activities. These studies demonstrated that DNB approaches can detect the early warning signals for detecting the normal and pre-disease states, and provide insights to the molecular mechanism of disease phenotype or complex physiological processes.

Fang et al. investigated the relationship of cigarette smoking and lung cancer development [44]. The disease states (tumour or normal), smoking states (current smokers or non-smokers or former smokers), and the disease stage (stages I–IV) were pair-wise compared using a novel strategy that incorporates network-based approach with gene set enrichment analysis. They identified panels of gene candidates that involve in cell proliferation and drug metabolism, such as cytochrome P450 and WW domain containing transcription regulator 1, in smoking or lung cancer development. Pathways of cell cycle, DNA replication, RNA transport, protein processing in endoplasmic reticulum, vascular smooth muscle contraction and endocytosis were found commonly involved in smoking and lung cancer. Furthermore, semaphorin 5A and protein phosphatase 1F were identified as the common genes represented in major hubs in both the smoking and cancer-related network. This study provides an excellent example not only to understand the complex and dynamic relationships between cigarette smoking and lung cancer but also to reveal molecular mechanisms of cancer initiation and progression at a network level.

Correlation between network biomarkers and clinical informatics

Network approaches allow an accurate identification of biomarkers. DNBs have the advantage to demonstrate pathophysiological changes at different stages and periods. One of the most challenges is to translate network biomarkers into clinical application and validate the disease specificity [9, 10]. The biological measurements need to be statistically and mechanistically correlated with clinical information. But conventional clinical measurements, such as complaints, history, symptoms and signs, physical examinations, laboratory tests, medical imaging and therapies, are descriptive and rarely integrated, comparing with biological data. As such, clinical bioinformatics was proposed to combine clinical phenotypes with human tissue-generated bioinformatics, to understand molecular mechanisms of the disease, and to define relationships between collectively direct global function with clinical measurements [9]. Defined as 'the clinical application of bioinformatics-associated sciences and technologies to understand molecular mechanisms and potential therapies for human diseases', clinical bioinformatics emphasizes the association and specificity complex biomedical data sets with the disease complexity of patient information. It suggests that the integration of biology data with clinical informatics can be a new way to validate and optimize disease-specific network biomarkers [10]. It would be even more values if clinical bioinformatics can integrate network-based approaches to prioritize disease-specific interaction subnetworks between gene–gene, gene–protein, or protein–protein with disease signature and clinical phenotypes, to improve the accuracy of clinical diagnostics and prediction.

The strategies to integrate biological and clinical data have been proposed and are still under the rapid development [129, 130]. Using a clinical bioinformatics approach, Schwarz et al. [131] quantified relationships between specific variables of patients with schizophrenia (i.e. cerebrospinal fluid and serum samples, obtained from two different profiling platforms and standard laboratory tests) as networks, and detected a subgroup of patients featuring remarkable abnormalities in a network of serum primary fatty acid amides. The identified disease-associated patterns of biomarkers were suggested to be able to describe the complicity of psychiatric diseases. This particular study demonstrated that simultaneous evaluation of clinical data and molecular biology data via a clinical bioinformatics approach could improve the understanding of complex diseases and lead to better diagnosis, prediction and therapy.

Because of the large and independent nature of the clinical data, the application of controlled vocabulary and ontology for the standard nomenclature of clinical trial data is critical and important for clinical data integration [129]. In a preliminary study, Chen et al. [22] utilized chemokine multiplex antibody array to detect inflammatory mediators in the circulation of patients with acute exacerbation (AECOPD) or stable condition (sCOPD) of chronic obstructive pulmonary disease to correlate DNBs with clinical informatics. Clinical informatics, which translates clinical descriptive information into the digital data, was achieved by a digital evaluation score system (DESS) for assessing severity of the patients. DESS is a score index established by senior chest physicians that take into account patient symptoms, signs, doctor examination, clinical imaging and biochemical analyses in patients with AECOPD or sCOPD. For the assessment of the severity, each component was assigned with different scores as 0, 1, 2 and 4. The score of 4 (maximal value) indicates far more above normal range or more severe condition, while 0 (minimal value) means the variable is within physiological range. The value of 3 was missed in the scoring.
system for exponential values to better define the severity stages. The points of each variable were added after compiling patients’ data and DESS values ranged from 0 to 264, of which higher scores indicate severer conditions. By integrating proteomics-based bioinformatics with clinical informatics, disease-specific biomarkers in the circulation were scanned and a multi-scale predictive model was established. The authors identified 13 mediators (BTC, IL-9, IL-18Bpa, CCL22, CCL23, CCL25, CCL28, CTACK, LIGHT, MSPa, MCP-3, MCP-4 and OPN) that could discriminate AECOPD patients from both healthy and sCOPD patients.

Using similar approach, the authors further investigated the potential correlation of proteomic profile with clinical informatics in COPD patients. Plasma samples from 18 patients including healthy individuals or patients with sCOPD or AECOPD were collected to measure 507 inflammatory mediators using antibody microarray [24]. Clinical descriptive information was translated into digital data by DESS for severity assessment. Twenty mediators were significantly different between three groups, of which Cerberus 1, inhibit B, osteoactivin and thrompoietin were firstly reported in COPD and AECOPD. Ten cytokines such as Cerberus 1, Growth Hormone R, IL-1F6, IL-17B R, IL-17D, IL-19, Lymphotoxin beta, MMP-10, Thrombopoietin and TLR4 were found inversely correlated with DESS scores. A down-regulation of systemic inflammatory responses was indicated in the occurrence of AECOPD. These studies demonstrated that the integration of omic profiles with clinical informatics as part of clinical bioinformatics is important to discover, validate and optimize disease-specific and disease-staged biomarkers. The proposed protocol for disease-specific biomarker discovery by integrating bioinformatics and clinical informatics is illustrated in Figure 4. Firstly, the expression profiles and the clinical data of a disease phenotype during disease progression are obtained. Second, the disease-associated networks are measured by bioinformatics, and clinical informatics is generated through a digital evaluation score system. Thirdly, the molecular-phenotype networks are then measured and ranked by integration of bioinformatics and clinical informatics to identify candidate biomarkers. And lastly, the identified disease-specific biomarkers are validated for clinical applications.

The advantages and constraints for clinical application

Protein-based network biomarkers with systems information can provide a more precise and complete profile of cellular changes in human diseases. As proteins perform the major cellular functions essential to signal transduction that role cell growth, differentiation, proliferation and death, protein-based network biomarkers are critical in providing valuable information at the post-translational level that can be used to establish diagnosis or prognosis of a disease and to develop personalized therapeutics, with favourable clinical feasibilities. The effectiveness of protein-based network biomarkers has been demonstrated in the context of various diseases.

On the other hand, a number of proteomics-associated challenges should be bear in mind. For instance, proteomics experiments typically screen only a limited fraction of proteins, in particular, gel-based expression experiments are most likely to detect high abundance proteins. The human PPI data are still incomplete and variable because of different curated collections. Therefore, the advantages of the network biomarker discovery include not only easily avoiding data noises by knowledge-based network but also deriving high confident network biomarkers.

A number of challenges still exist in every step of the network biomarker development pipelines, despite of technological advances. Protein post-translational modification and alterations in protein stability may influence network modularity on a global scale during disease progression. Most high-throughput methods can suffer from high false-positive or -negative rates that may lead to misclassifications. Large-scale networks are not specific to diseases or clinical phenotypes. The level of certainty is constrained by the issues of data collection, interpretation of large size of the proteome, and the diversity of cells and tissues. On the other hand, noise generated during different network methodologies integration remains a major constraint to correctly interpret complex networks and needs to be critically evaluated and managed. Thus, the principles used in network discovery, validation and development remains to be further defined and quantified, and the development of novel and reliable statistical tools for the network environment is urgently needed. Also, the lack of standardized vocabularies for a definitive translation of networks into the clinical arena represents a main challenge in the integration and interpretation of clinical bioinformatics. Another important challenge is to translate DNBs into the understanding of clinical phenotypes, molecular mechanisms of disease development and progress, and development of therapeutic strategy [10].

Despite these limitations related to knowledge incompleteness and uncertainty in the network inference process, the characterization of complex biological phenomena on the basis of functional modular architectures and topological parameters present us with new opportunities to improve our understanding of the aetiology, evolution and therapeutics of the diseases. To reach clinical application, the advantages and disadvantages of protein-based network biomarkers should be furthermore investigated to evaluate the potential values of network biomarkers in the development. We believe that clinical bioinformatics can play an important role in identification and validation of disease-specific DNBs.

Prospective and conclusions

Better biomarkers are urgently needed to disease detection, diagnosis and prognosis. Network approaches have revolutionized the traditional ways of biomarkers discovery and offered a powerful way for pathway mapping and development of disease-specific biomarkers. Although challenges exist in steps of the network biomarker development, network biomarkers are proving to play importance roles in disease-causing genes prediction, disease-related subnetworks identification, disease classification, disease in vitro/in silico modelling, drug discovery and target screening, and ultimately improving the outcome and life quality of patients.
In the era of network medicine, new biomarker discovery depends on a comprehensive view of transcriptome, genome, proteome, and metabolome, or diseaseome. The dynamic nature of human protein interaction network because of the diversity and regulative structure of post-translational modifications, gives in-depth insight into disease mechanism. Scientists and physicians are facing more challenges to keep the pace with the growing availability of a variety of high-throughput data. Global efforts are been done to improve our understanding of diseases through integrative approaches to translate science into the clinical practice [132–134]. With the development of clinical bioinformatics, biomarker discovery should not only integrate different types of omics data, but also consolidate of such molecular biological measurements with clinical phenotypes. The development of profiling technologies, biological databases, data mining, biostatistics and clinical bioinformatics will tremendously speed up the identification, validation and development of disease-specific network biomarkers and DNBs.

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Conflicts of interest

The authors confirm that there are no conflicts of interest.

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