Combinatorial Regulatory Networks Provide Novel Biomarkers for Diagnosis and Therapeutics

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Introduction

One of the most important discoveries of the post-genomic era is that a large fraction of the genome transcribes a heterogeneous population of non-coding RNAs (ncRNA). NcRNAs shorter than 200 nucleotides are usually identified as short/small ncRNAs - examples include PIWI-interacting RNAs, small interfering RNAs, and microRNAs (miRNAs)- whereas those longer than 200 nucleotides are classified as long ncRNAs (lncRNAs). They play a crucial role in RNA silencing and post-transcriptional regulation of gene expression [1]. These molecules are emerging as important regulators of cellular process, such as development, differentiation, and metabolism [2]. Long and short ncRNAs not only regulate developmental processes and cell physiology, but also involved in disease states, host-pathogen interactions, and a variety of stress responses [3]. Transcriptional regulation of gene expression is often the prior process due to the response of system changes, wherein the information contained in a genome is converted and then ultimately used to produce the proteins required for a given response.

It is arranged by transcriptional regulatory proteins and their target genes working in assemblage through a variety of mechanisms [4]. These regulators, known as transcription factors (TFs), are proteins bind to either enhancer or promoter regions of DNA adjacent to the genes that they regulate [5]. MicroRNAs (miRNAs) have arisen as another eminent class of transcription regulator component which have huge effects on protein formation at pre-transcriptional and post-transcriptional levels. Furthermore, it was found that IncRNAs get involved in transcriptional regulation; as well as [6]. And they are responsible for activation or inhibition of TFs [7]. Though determining of the disease related gene and/or ncRNA is crucial to improve knowledge about disease pathogenesis, these components remain meager in most situations to understand the complex screen underlying disease mechanisms rev [8].

Systems biology approaches may be provide integrated analysis of different experimental data sets and understanding of how alterations in regulatory networks in disease progression by using interactions of molecules (i.e: gene, protein, metabolite) at different scale (genome, RNA, proteome, metabolome level) [9-11]. On the contrary one component (i.e: TF, miRNA, lncRNA and gene) based approaches, network biology perspective is a versatile tool to evaluate interactions of all transcriptional regulatory components (i.e: mRNA and ncRNA) with comprehensive aspect. On the other hand, many studies revealed that miRNA-TF interaction play a significant role in gene expression mechanism. Arora et al. [12] summarized that a combinatorial action of miRNAs and TFs which share similar regulatory logics and bring about a cooperative action in the gene regulatory network were determined in biological processes.

Several transcriptional regulatory network studies in which include notably miRNA-TF- target gene interaction network have been carried out. Tsang et al. [13] investigated the expression pattern between miRNAs and their target genes, and suggested that miRNAs and their targets can share common regulator(s). Gennarino et al. [14]. Were performed an analysis of human miRNA regulatory networks by focusing on the expression relationships among miRNA targets. Yu et al. [15] reported that the most important tumor-regulating miRNAs and TFs were highlighted from a topological point of combinatorial regulatory network of transcriptional regulation in human cancer. Recently, reconstruction of human generic transcriptional regulatory network by using experimentally verified data and demonstrated new insights on the reciprocal interplay between miRNAs, TFs and their target genes [16].

Data-driven reconstruction of networks included lncRNAs have previously performed [21-23]. However, disease specific ncRNA enriched transcriptional regulatory network will be investigated
to identify RNA biomarkers for diagnosis and treatment by using systems biology perspective. Publicly available databases of non-coding RNA such as miRBase [24], MiRDB [25] and ChIPBase [26] for miRNAs, LNCipedia.org [27] IncRNAdb [28] and NONCODE [29], for lncRNAs can be used for construction of integrative transcriptional regulatory network.

Conclusion

The crucial aim for the further studies is to develop tools and databases to identify putative functional and pathological interactions between ncRNAs, TFs and their target genes associated with various diseases. Integration of multiple data (i.e: gene expression, ncRNA and interactions data) allows us to create more holistic aspect of the disease active transcriptional regulatory networks, namely ncRNA-enriched disease networks. This can lead us to understand properties of TF-miRNA-lncRNA- their targets regulatory mechanisms and role of ncRNAs in large gene expression regulatory network. Integrative and systematic analysis of ncRNA enriched transcriptional regulation is key element to detect more reliable RNA biomarkers for diagnosis and treatment strategies.

Moreover, the discovery of novel ncRNAs, as well as their prominent role in disease specific transcriptional regulation has accentuated with respect to identify diagnostic and therapeutic targets, but to fully understand how these RNAs contribute to gene expression need to be experimental studies. Identification of the proteins and pathways regulated by these miRNA and IncRNA is likely to provide novel insights into disease pathogenesis provide novel biomarker for disease diagnosis and determine novel drug targets for therapeutic applications. However, complicated gene expression mechanisms can be solved with RNA molecules which open new exciting avenues, as our understanding of RNAs’ role improves, RNA world represent us amazing knowledge.

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References

