PTHGRN: unraveling post-translational hierarchical gene regulatory networks using PPI, ChIP-seq and gene expression data

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ABSTRACT

Interactions among transcriptional factors (TFs), cofactors and other proteins or enzymes can affect transcriptional regulatory capabilities of eukaryotic organisms. Post-translational modifications (PTMs) cooperate with TFs and epigenetic alterations to constitute a hierarchical complexity in transcriptional gene regulation. While clearly implicated in biological processes, our understanding of these complex regulatory mechanisms is still limited and incomplete. Various online software have been proposed for uncovering transcriptional and epigenetic regulatory networks, however, there is a lack of effective web-based software capable of constructing underlying interactive organizations between post-translational and transcriptional regulatory components. Here, we present an open web server, post-translational hierarchical gene regulatory network (PTHGRN) to unravel relationships among PTMs, TFs, epigenetic modifications and gene expression. PTHGRN utilizes a graphical Gaussian model with partial least squares regression-based methodology, and is able to integrate protein–protein interactions, ChIP-seq and gene expression data and to capture essential regulation features behind high-throughput data. The server provides an integrative platform for users to analyze ready-to-use public high-throughput Omics resources or upload their own data for systems biology study. Users can choose various parameters in the method, build network topologies of interests and dissect their associations with biological functions. Application of the software to stem cell and breast cancer demonstrates that it is an effective tool for understanding regulatory mechanisms in biological complex systems. PTHGRN web server is publicly available at web site http://www.byannbioinfo.org/pthgrn.

INTRODUCTION

Gene regulation of eukaryotic organisms is a very complex process that is mainly carried out by tight interactions between transcription factors (TFs) and DNA sequences in specific ways (activation or inhibition). The ability of TFs to regulate target genes is modified by post-translational protein–protein interactions (PPIs) among TFs, cofactors and other proteins or enzymes upstream of transcriptional gene regulation. A variety of post-translational modifications (PTMs), including protein phosphorylation, acetylation and ubiquitination have been implicated in transcriptional gene regulation (1–3). Emerging evidences have indicated that PTMs of proteins are involved in many biological processes or human diseases through controlling DNA-binding ability to modulate downstream gene expression (4,5). In addition, epigenetic modifications, such as histone methylation, have been documented as crucial elements for regulation of genome function through changing chromatin architecture without altering DNA sequences. The combinatorial action of PTMs, TFs, as well as epigenetic alterations constitutes the hierarchical complexity in gene regu-
lution. Despite its importance, we have incomplete knowl-
edge of these processes, which limit our understanding of
complex regulatory mechanisms.

The advent of high-throughput technologies allows bi-
ologists to examine molecule interactions at different lev-
els, and generates multi-dimensional Omics datasets. There
are several technologies used to detect PPIs, like co-
immunoprecipitation with display technology (6), tandem
affinity purification (7) and yeast two hybrid (8). Mass
spectrometry-based methods can identify numerous PTMs
(9), providing more insights into the associations between
PTMs and gene expression. Coupling chromatin immuno-
precipitation with next-generation sequencing (ChIP-seq)
offers high resolution mapping of TFs or epigenetic modifi-
cations’s interaction sites to genomic locations (10). ChIP-
based studies detect the binding regions of these regulators
on DNA sequences and show a genome-wide binding affin-
ity between protein–DNA sequences. Therefore, ChIP-seq
is informative for transcriptional or epigenetic regulatory
relationships and reconstructing gene regulatory networks
(GRNs).

A major goal of mining high-throughput data is to find
underlying structures in the data that provide a basis for
identification of regulatory modules and reconstruction
of complex protein/gene networks. Computational algo-
rithms have proven efficient for addressing this issue. For
example, MINDy facilitates genome-wide identification of
PTMs of TFs, and was successfully applied to determine
the regulation of MYC activity in human B lymphocytes
(11). Other methodologies have been developed to assemble
both gene expression profiling and ChIP-seq or TF binding
data for identifying TF–gene interaction modules, such as
Bayesian multivariate modeling (12), matrix decomposition
(13) and regression model (14). Graphical Gaussian model
(GGM) was widely used for inferring GRNs by exploiting
high dimensional throughput data (15–17). Recently, an
integrative model Active Protein-Gene (APG) based on linear
GGM with matrix decomposition was designed to connect
upstream protein–TF networks with downstream TF–gene
networks (18). Partial least squares (PLS) is a well-known
regression tool suitable for statistical analysis of genomic
and proteomic data, and modeling of gene networks and
TF activity (19). Some studies have applied the PLS method
to examine connections between TF or microRNAs and
gene expression (20,21). Therefore, a strategy combining the
GGM with PLS regression could be powerful way to de-
velop network-related web software.

Online tools or servers have been reported that build
transcriptional or epigenetic regulatory networks, for in-
stance, ChIP-Array (22), ChEA (23) and RENATO (24).
Our web-based framework CMGRN was devised for inte-
grative analysis of causal relationships between regulators
and complex gene regulation mediated by TFs, epigenetic
factors and microRNAs (25). MAGNET server can gen-
erate and score PPI networks and coexpression gene–gene
networks but did not connect both together computa-
tionally (26). Although these methods show improved results in
uncovering transcriptional regulatory programs, there is a
lack of effective web-based software capable of constructing
underlying interactive networks linking post-translational
and transcriptional regulatory components.

Figure 1. A workflow of integrated data analysis and GRN construc-
tion. PPI, TF binding and gene expression data are used to infer protein–
TF interactions. The ChIP-seq binding peaks of TFs/epigenetic modifi-
cations are integrated with gene expression profile to infer TF/epigenetic
modification–gene interactions. Finally, the tabulated results are organized
to hierarchical GRNs.

A large number of PPI (such as BioGRID, STRING, HPRD and Reactome), ChIP-seq and gene expression
(such as ENCODE, modENCODE, GEO and ArrayExpress) resources are publically available. To provide an easy
bioinformatics tool to use and interpret the high through-
put Omics data, we present an integrative web server, post-
translational hierarchical gene regulatory network (PTH-
GRN) (http://www.byanbioinfo.org/ptghrn) to unravel re-
lationships among PTMs, TFs, epigenetic modifications
and gene expression. The newly developed server performs
a GGM with PLS regression-based methodology to gen-
erate and score potential interactions of both protein–TF
and TF/epigenetic modification–gene. It is system-wide and
enables biologists to process the mixture information from
PPI, ChIP-seq and gene expression, using standard data
formatted with minimal need of bioinformatics skills. At
the end, PTHGRN server robustly constructs hierarchical
GRN for further evaluating the effect of PTMs on tran-
scriptional regulatory complexity. Application of the soft-
ware in mouse embryonic stem (ES) cell and human breast
cancer demonstrates that it can explore biologically mean-
ful regulatory networks. Proof of principle analyses vali-
dates the use of PTHGRN for systems biology applications.

MATERIALS AND METHODS

Workflow of PTHGRN

As outlined in Figure 1, PTHGRN mainly conducts two
tasks simultaneously. To examine how proteins (including
cofactors, TFs, enzymes, etc.) affect TF performance post-
translationally, input datasets, comprised of PPI, TF bind-
ing and gene expression, are submitted to the server. In this
step, a GGM with PLS regression-based method will pro-
file protein effects on TF activity and evaluate all interac-
tions between TFs and proteins. The result is a list of the in-
ferred protein–TF interactions that satisfy cutoff P-values
(Table 1 of Figure 1), which can be used for constructing

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cations are integrated with gene expression profile to infer TF/epigenetic
modification–gene interactions. Finally, the tabulated results are organized
to hierarchical GRNs.
upstream protein–TF networks. Another task is inference of regulator (TF/epigenetic modification)–gene interaction networks. Two types of input data are required: ChIP-seq binding peaks of the regulators or TF binding and gene expression. By performing the similar methodology, PTHGRN generates and scores all possible interactions between the regulators and genes. The resulting table is a list of the inferred regulator–gene interactions that satisfy cutoff P-values (Table 2 of Figure 1). Depending on availability of input data, users can carry out transcriptional regulatory network task with only gene expression and binding data of regulators. If with gene expression and PPI data only, the server will construct PPI networks, where the proteins are present in expression data. With all the three input data, the server organizes the identified protein–TF and regulator–gene interactions together and constructs a post-translational hierarchical GRN.

Scoring interaction networks

PTHGRN used a GGM (15–17) with PLS regression (19) to perform an integrated analysis of gene expression profiles with PPI and binding data of a set of regulators, and to discover protein–TF and regulator–gene interactions. The basic algorithm is to generate and then score every possible interaction. Based on the input data, the algorithm sets several matrices: $E$ represents expression level of genes, $T$ refers to concentrations of TF targeting genes, $P$ stands for concentrations of both TFs and proteins, and $R$, $E$, $T$, $P$, $E$, $T$, $F$, $A$ represents TFs activities under different conditions. Except the four matrices, the algorithm will also generate two temporary connectivity matrices (consisting of 1 and 0), $A$ referring to one with TFs targeting genes and $B$ referring to one with proteins modifying TFs. In the matrix $A$, ‘1’ indicates that a TF target gene in the ChIP-seq or TF binding data is also found in matrix $E$, and ‘0’ otherwise. Similarly, ‘1’ in the matrix $B$ indicates that a protein in input PPI is also present in the ChIP-seq or TF binding data, and ‘0’ otherwise. To estimate initial observations of the matrix $T$, the median expression level of TF target genes from $E$ and $A$ are used. Similarly, initial observations of the matrix $P$ is calculated from $E$ and $B$ based on the median strategy. According to the dependent and conditional independent features of Bayesian network, the algorithm can establish the joint distribution $Pr(T, P, E, T, F, A)$ corresponding to the four matrices. A natural choice for representing continuous variables is the use of Gaussian distribution. According to the Gaussian model, if $S$ is a node in graphical network, we have the conditional density of $S$ given its parents:

$$Pr(S | R_1, \ldots, R_n) \propto N \left( \sum_{ij} \delta_{ij} r_{1\ldots n}, \sigma^2 \right),$$

where $R_1, \ldots, R_n$ are the parents of $S$; $N(\mu, \sigma^2)$ is the density function of the normal distribution with mean $\mu$ and standard variations $\sigma$, and $r_1, \ldots, r_n$ are the observations of $R_1, \ldots, R_n$ respectively. $\delta_{ij}$ is the effect strength of the $i$th variable on the $j$th variable. Since the activities of TFs depend on concentrations of both TFs and proteins, as described in APG (18), we can use $\delta_{ij}$ representing the interaction scores of TFs targeting genes or proteins modifying TFs to update the joint distribution $Pr(T, P, E, T, F, A)$ above. The TFA is a hidden variable depending on concentrations of both TFs and proteins. To set the joint distribution parameters, we employed PLS regression (19) to find true TFAs and the associated interactions. PLS-based network component analysis offers computationally highly efficient and statistically robust strategies to identify likely true TFAs for any given connectivity matrix. In addition, it allows statistical assessment of the available connectivity information, and also the discovery of interactions and natural groupings among regulatory components. During the whole scoring process, we used the Maximum Likelihood Estimation method to obtain optimal parameters or network structure of the graphical model and maximize the probability of observed interaction scores (17,18).

Estimating probability of interaction networks

In order to select potential interaction networks, PTHGRN will evaluate $P$-values through a randomization test. First, the server will generate all possible interactions between protein–TF and regulator–gene based on the original data without data randomization, so called signal interactions. Next, to conduct the randomization test, the input data will be randomly permuted a given number of times. The number, or ‘iteration’, can be chosen in the input web interface, for example, iteration = 100, indicating that the server is going to run 100-time data permutations, i.e. 100-time tests. For every permutation, PTHGRN randomly picks up the same number of proteins as true protein–TF from the PPI database, and the same number of genes as true regulator–gene from the ChIP-seq or TF binding data. The server then randomly sets the parameters for the matrices with the same structure as signal interaction generation for every test. Following the same procedure of scoring interaction networks described above, the server will score each of the randomized interaction networks, so called background interactions, to form background scores. PTHGRN then compares the obtained score with the background scores, and calculates the $P$-values of every signal interaction. Furthermore, we corrected the $P$-value based on False Discovery Rate using $R$ function $P$-adjust. The networks with $P$ or corrected-$P$-value satisfying a certain cutoff (for example, $P < 0.05$ or corrected-$P < 0.05$) will be chosen for the lists in output tables and visualization. In general, the score of the likely true signal is higher than its corresponding backgrounds due to data randomization, indicating lower $P$-values. In the web interface, users can set cutoff of $P$-value or corrected-$P$-value for selecting the output interactions. The randomization-based method has been widely used in many computational biology studies (13,27–29).

Web interface

PTHGRN provides an easy web interface platform for incorporating three types of input data. The first one is a tab-delimited list of PPI derived from public databases Biogrid, STRING, Dip, HPRD, Intact, Mint and Reactome, and consists of two columns containing pairs of interacting proteins. Currently, PTHGRN contains PPI data
that covers human, mouse, rat, *Drosophila melanogaster* and *Caenorhabditis elegans*. The server system also supports uploading of user datasets with a similar format. For example, results from proteomics or metabolomics experiments could be treated as PPI input. The second one is tab-delimited regulatory signal, and consists of two columns including the name of regulators (TFs or epigenetic modifications) and their target genes. A regular method to prepare the signal data is to analyze significantly enriched ChIP-seq binding peaks of the regulators along genomic sequences by using MACS (30) or other ChIP-seq analysis software. ENCODE project stores available ChIP-seq binding peaks of the regulators. We mapped these binding loci to the genomic regions from transcriptional start site 2000 bp to transcriptional end site 1000 bp on human or mouse genome. We extracted these binding data, including 171 and 47 TFs, as well as 45 and 13 epigenetic modifications in human and mouse, respectively. Overall, our server recruited a total 135 cell types of ENCODE, 108 from human and 27 from mouse. We also collected ChIP-seq binding peaks of *D. melanogaster* (including 42 TFs and 28 epigenetic modifications) and *C. elegans* (including 91 TFs and 91 epigenetic modifications) from public database modENCODE. Currently, PTHGRN provides ready-to-use binding peaks of the regulators from the two projects. Moreover, users can upload their own binding data in the required format. For example, conserved binding motif of TFs on the promoter can be treated as regulatory signals. In the server, we provide TF target genes containing conserved binding sites on promoters of human, mouse and rat (retrieved from ECRbase, http://ecrbase.dcode.org/), as well as *D. melanogaster* (22). The last input is tab-delimited numeric matrix and gene expression data, where the columns are sample ID and each row represents gene names. The expression data should consider the same biological materials as ChIP-seq experiments if using ChIP-seq binding data. We suggest that expression data contain differentially expressed genes, and are separated to up-regulated and down-regulated subgroups, respectively. In all the three input files, the gene and protein ID format should be consistent for data integration. The current databases of PTHGRN support official gene/protein symbols.

The web interface provides example data from mouse ES cell line V6.5 and human breast cancer cell line MCF-7. The ES cell data contains mouse PPI, ChIP-seq binding peaks of five histone methylations (H3K4me3, H3K27me3, H3K79me2, H3K36me3 and H4K20me3), four TFs (Oct4, Sox2, Nanog and Tcf3) and PRC2 component Suz12 from GSE12241 and GSE11724 of NCBI GEO database. The gene expression data was extracted from GSE32321 of GEO and characterized with a time-course profiling of genes differentially expressed from embryonic status to embryoid bodies. In the breast cancer example study, we processed human PPI, ChIP-seq binding peaks of five histone modifications (H3K4me3, H3K27me3, H3K36me3, H3K9me3 and H3K27ac) from GSE31755 of ENCODE project, and six TFs (ERα, CEBPB, FOXM1, GATA3, JUND and MAX), as well as cofactors p530 and CTCF from GSE32465 of ENCODE and GSE25684 of GEO. The gene expression data is based on time-course profiling of genes differentially over-expressed in the estrogen-stimulated breast cancer cell from accession number E-TABM-742 of the EBI database.

After uploading the input files and selecting certain cut-off parameters, users can start the PTHGRN procedure by click ‘Submit’ button. The interface displays three cut-off choices, *P*-value, corrected-*P*-value and iteration. A low *P*-value indicates a high degree of confidence that a protein/TF can be modified by another one or that a gene can be targeted by a regulator. Following the submission action, the server generates an output interface showing result networks exploring post-translational, transcriptional and epigenetic regulatory hierarchies in graphical and tabular formats. Based on the two main output tables (see Figure 1), the server automatically generates a third table with a list of interacting protein pairs that can modify the TFs in the second table. This process can be carried out by directly searching the PPI database in PTHGRN server. The graphical interface would show a view of post-translational GRN including the three output tables above. Users can choose the interaction links of TF–protein or regulator–gene under post-translational, transcriptional and epigenetic levels. For example, users can click a TF of interest, and see an overview of all nodes (its target genes and proteins that modify it) and edges linking to this TF. The graphical topology generated enables users to manipulate the pan/zoom choice, search and retrieve text information showing score and *P*-values of all interactive TF–protein or regulator–gene for a selected node and its associated subgraph. In addition, the server can retrieve NCBI protein or gene information for the selected node.

**Implementation**

PTHGRN platform is modularly designed to allow other combination of regulatory networks, metabolic networks, pathway networks and drug-target networks. The web server is written in PHP with supporting scripts written in Pascal. Network display was created using Cytoscape web. PTHGRN does not have excessive computing requirements. In its current deployment and under typical server loading conditions, PTHGRN can finish processing a typical input dataset with ~5–15 min. Optionally, users provide an Email address to receive a link for finished results if large datasets or large numbers of iteration are requested. The results will be stored for a month by default, and storage times can be longer or shorter depending on user’s requirement.

**EXAMPLE STUDY**

**Mouse ES cell differentiation**

Orchestrated interplays among three pluripotency TFs Oct4, Sox2 and Nanog with other TFs, cofactors and epigenetic modifications are critical for controlling ES cell self-renewal and differentiation (31,32). We applied PTHGRN to investigate the complex regulatory associations in mouse ES cell. Figure 2 displays an output hierarchical GRN coordinated by PTMs, TFs and histone methylations. This result implies that a multi-level regulatory complex is involved in down-regulating genes that are related to transcriptional regulation, DNA
metabolic process and RNA processing, and thereby promoting ES cell differentiation. Under protein networks, we identified 44, 14 and 58 proteins that interact with the Oct4, Sox2 and Nanog respectively (Supplementary Table S1). In particular, several important modification interactions among TFs, epigenetic enzymes and cofactors were discovered (Figure 2), such as Sox2-Oct4-Nanog, Jarid2-Ezh2-Suz12-Eed-Mtf2-Rbbp4, Ezh2-Rbbp4-Sall4-Sox2-Klf5-Oct4, Phc1-Oct4-Nanog-Sox2-Eed, Klf4-Oct4-Suz12 and Dnmt3b-Dnmt3l-Nanog-Oct4, some of which are consistent with known reports (33–40). This finding validates that three core TFs Oct4, Sox2 and Nanog interact with each other and also with other TFs or proteins for the regulation of their downstream genes in stem cells (33). Three PRC2 components Ezh2 (H3K27 methyltransferase), Suz12 and Eed interact with each other to contribute to H3K27 methylation for repressing gene expression of mouse ES cells (37). Their cofactor Jarid2 can recruit the PRC2 proteins to common promoters for jointly regulating gene expression (36). Interestingly, ChIP-based experiments indicated that PRC2 binding is highly correlated with binding of Oct4, Sox2 and Nanog (41). Our analysis supports the roles of these key regulators during ES cell differentiation. At transcriptional level, Oct4, Sox2, Nanog can regulate 31–36% of genes differentially under-expressed respectively, where 102 are common targets. Their target genes include key TF or epigenetic enzyme genes, such as Mybl2, Mycn, Sall4, Trp53, Esrrb, Eed, Ezh2, Phc1, Jmjd1a, Jmjd2c and Jarid2 (Supplementary Table S1). At epigenetic layer, we identified 41–43% of differentially under-expressed genes modified by three histone modifications, H3K4me3, H3K79me2 and H3K36me3 respectively. Epigenetic factors Suz12, H3K27me3 and H4K20me3 only regulate a few numbers of genes (Supplementary Table S1). Similarly, we tested differentially over-expressed genes using the same method. The result shows that repressive Suz12 and H3K27me3 are able to target 35 and 15% of up-regulated genes respectively, which are much greater than down-regulated ones (Supplementary Tables S1 and S2). This observation supports previous perspectives regarding role of PRC2 and the repressive epigenetic factors in leading to over-expression of developmental genes in the differentiated stem cells (42,43).

**Estrogen-induced human breast cancer cell**

Estrogen receptor alpha (ERα) is an estrogen-inducible TF that has been implicated in the development of human breast cancer. As a nuclear receptor, it can interact with many TFs, coregulators and growth factor-activated signalling to form a regulatory complex to modulate cancer-related biological processes (44). PTMs of ERα have been found through phosphorylation, ubiquitination, sumoylation and acetylation that affect its stability and activity (45). To dissect this complexity, we applied PTHGRN to analyze breast cancer data. The observation shows that ERα and cofactor p300 interacts with 73 and 33 proteins upon 24-h estrogen stimulation, respectively, where including TF proteins E2F1, FOS, MYC, FOXM1, IRS1 and IRS2 (Supplementary Table S3). ERα was found to link with other proteins to form protein networks, such as ERα-MYC-IRS2-CAV1, ERα-MYC-HSPA8-HSPH1, ERα-MYC-TUBA1-TUBB, ERα-MYC-FOS-NCL and ERα-MYC-CTSD. The previous study has revealed that ERα and MYC physically interact with each other to stabilize the ERα-coactivator complex and to facilitate estrogen-mediated signaling networks (46). ERα also forms a complex with AP-1 family members (including FOS) to modulate bone-specific genes in osteoblasts (47). Although p300 and ERα does not directly interact each other, both can link with MYC and FOS (Supplementary Table S3). This analysis highlights a considerable interaction of ERα as a master player with many coregulators upstream of transcription in breast cancer cell. Furthermore, we identified target genes of eight TFs or cofactors and five epigenetic modifications (Supplementary Table S3). It is apparent that 25–27% of over-expressed genes could be modified by three active epigenetic factors, H3K4me3, H3K36me3 and H3K27ac, whereas repressive H3K9me3 and H3K27me3 only target a small number (4–6%) of genes. The epigenetic features of histone modifications could contribute to up-regulate estrogen-mediated gene expression in breast cancer. Taken together, the example in breast cancer confirms that ERα is a key factor of regulatory complex networks, which modulate estrogen-induced gene expression programs and promote molecular pathogenesis of breast cancer.

**CONCLUSIONS**

PTHGRN is a freely available web server for an integrated analysis of PPIs, ChIP-seq binding data and gene expression profiling. Using a GGM with PLS regression-based method, it can generate and score all possible interactions of protein–TF and TF/epigenetic modification-gene.
Through statistical assessment, the server performs identification of highly potential interactions and reconstruction of hierarchical GRNs for evaluating roles of PTMs in transcriptional gene regulation. The web interface provides users ready-to-use input data derived from major high-throughput Omics resources, so that users have an option to submit these public or user-provided data. The workflow of PTHGRN produces output networks in graphical and tabular formats, which can be downloaded for further systems biology studies and visualization. The examples presented here using mouse ES and human breast cancer cells show the identified interaction networks to be biologically meaningful, in agreement with conclusions previously drawn from experiments and reports. It is expected that the newly developed server is able to provide a pilot framework for extensively unraveling regulatory associations among multilayered molecules in many biological complex systems, and thus would benefit not only biologists but also the bioinformatics community.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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