Abstract. Osteoporosis is a common skeletal disorder characterized by a decrease in bone mass and density. The peak bone mass (PBM) is a significant determinant of osteoporosis. To gain insights into the indicating effect of PBM to osteoporosis, this study focused on characterizing the PBM networks and identifying key genes. One biological data set with 12 monocyte low PBM samples and 11 high PBM samples was derived to construct protein-protein interaction networks (PPINs). Based on clique-merging, module-identification algorithm was used to identify modules from PPINs. The systematic calculation and comparison were performed to test whether the network entropy can discriminate the low PBM network from high PBM network. We constructed 32 destination networks with 66 modules divided from monocyte low and high PBM networks. Among them, network 11 was the only significantly differential one (P<0.05) with 8 nodes and 28 edges. All genes belonged to precursors of osteoclasts, which were related to calcium transport as well as blood monocytes. In conclusion, based on the entropy in PBM PPINs, the differential network appears to be a novel therapeutic indicator for osteoporosis during the bone monocyte progression; these findings are helpful in disclosing the pathogenetic mechanisms of osteoporosis.

Introduction

Osteoporosis is a common skeletal disorder characterized by a decrease in bone mass and density, which results in an increased risk of fractures among the elderly (1). Women, especially postmenopausal women, are more susceptible to osteoporosis compared to other groups. Bone mass and strength achieved at the end of the growth period is defined as peak bone mass (PBM), which is a significant determinant of osteoporosis (2,3). It is regarded that an increase of PBM by one standard deviation would reduce the fracture risk by 50% (4).

Peripheral blood monocytes can serve as early precursors of osteoclasts (5-7). A growing body of literature has explored that blood monocytes deliver many kinds of factors for bone metabolism, such as interleukin-1 and tumor necrosis factor-α (8). Osteoclasts in peripheral skeleton (9) and the central skeleton come from circulating monocytes (10). Substantial research has focused on the effect of circulating monocytes on pathogenesis of osteoporosis in young and middle aged adults.

Research in systems biology has shown that variability in the activity of gene network and frame structure play an important role in the disease progression (11,12). Network-based systems biology approaches have emerged as powerful tools for analysis of molecular mechanisms of diseases (13-16). An integrated network method was applied to predict conserved regulators related to high and low viral pathogenicity, leading to therapeutic targets for intervention (17). In addition, recent study confirmed that signaling entropy computable from integrating a gene expression profile with a protein-protein interaction network (PPIN), correlate with phenotypic plasticity and is increased in disease compared to normal controls (18). Based on information theory, entropy opens new perspectives for gene inference methods and increases the accuracy of PPINs (19). Jin et al (20) compared the IAV-induced inflammatory regulatory networks and normal cellular networks by integrating the data from the highly pathogenic avian H5N1 and the pandemic H1N1 with PPINs, and these findings provide significant hypotheses for further exploring the molecular mechanisms of infectious diseases and developing control strategies.

To gain insights into the indicating effect of PBM to osteoporosis, this study focused on characterizing the PBM networks and identifying key genes by combining high-throughput data and computational techniques. In PPINs, we searched for genes functionally significant for PBM variation, which might be related to circulating monocytes in human premenopausal subjects. The findings are expected to provide effective insights for further exploring the pathology mechanisms of osteoporosis and obtaining better treatment strategies.

Materials and methods

Datasets and construction of PPINs. One biological data set E-GEOD-7158 of osteoporosis was derived from the Gene
Expression Omnibus (GEO) database (http://www.ncbi.nlm.nih.gov/geo/) (3). There were 12 monocyte low PBM samples and 11 high PBM samples in the total 23 samples. The platform is A-AFFY-44-Affymetrix GeneChip Human Genome U133 Plus 2.0 (HG-U133 Plus 2.0). The Linear Models for Microarray Data (LIMMA) was then used to preprocess data. After quantile data normalization performed by robust multi-array average (RMA) (21), 20,545 genes were obtained.

The human PPIN was collected from the Retrieval of Interacting Genes (STRING; version 9.0) (22). Only edges whose correlations are greater than the threshold 0.6 were chosen, and set at 0.8. A total of 8,590 nodes and 53,975 protein-protein interaction (PPI) sets remained. Then we constructed an osteoporosis PPI subnet after getting intersection with expression profiles and the PPIN, which contained 7,953 genes and 48,778 PPI sets.

Identification of modules and paired comparison. Using the human subnet as a backbone, we infer a re-weighted PPIN with expression and mutation profiles of monocyte low PBM and high PBM in osteoporosis. Every side of the constructed PPI was assigned with absolute value of Spearman correlation coefficient of every interaction according to gene expression data.

Based on clique-merging, module-identification algorithm was used to identify modules from PPINs (23-25). We found all maximal cliques from low and high PBM PPIN, and the nodes <4 or >20 were filtered out. Then 8,002 maximal cliques in osteoporosis were obtained. We identified the set C of all maximal cliques of size at least k in the two groups using a fast depth-first search with pruning-based algorithm (CLIQUEs). Next, we calculated its weighted interaction density score (C) as (equation 1):

\[
\text{score (C)} = \sum_{u \in C, v \in C} w(u, v) / |C|(|C|-1)
\]

We ranked these cliques in descending order of their score (C) (23). A predefined overlap-threshold t=0.5 was set to go through the list repeatedly. The modules were gathered by merging highly overlapping cliques. Accordingly, we captured the effect of differences in interaction weights between monocyte low and high PBM group via the weighted density-based ranking of cliques.

The set of disrupted or altered module pairs were identified by modeling it as a maximum weight bipartite matching. The module correlation density was calculated between low and high PBM group as follows (equation 2):

\[
d_{cc}(S_i) = \frac{\sum_{p,q \in S_i} \text{PCC}((p, q), N)}{|S_i| \cdot |S_i| / 2}
\]

Characterizing networks from network entropy. Destination networks were constructed with modules of common genes of monocyte low and high PBM groups.

Network entropy, one of metrics of the inflammatory network, was detected in osteoporosis (20). The local network entropy of a node i, denoted S_o, is defined as (equation 3):

\[
S_i = -\frac{1}{k_i} \sum_{j \in N(i)} p_{ij} \log p_{ij}
\]

In which, k_i is the degree of node j, N(i) is the set of neighbor nodes of node i and p_{ij} defines a stochastic probability matrix on the PBM network, which is defined by (equation 4):

\[
p_{ij} = \frac{|c_{ij}|}{\sum_{k \in N(i)} c_{ik}}
\]

c_{ij} is the PCC between protein i and j.

The global entropy of destination networks, denoted S, is defined as follows (equation 5):

\[
S = \sum_{i=1}^{n} C_i S_i
\]

n is the total number of nodes in the destination networks, and C_i is the degree centrality of node i, which is defined by (equation 6):

\[
C_i = \frac{k_i}{n-1}
\]

The differential network entropy, denoted \(\Delta S_i\), is defined as follows (equation 7):

\[
\Delta S_i = S_i^l - S_i^H
\]

S_i^l, S_i^H is the local network entropy of node i in low and high PBM networks.

Significance test. The non-parametric one-tailed Wilcoxon rank sum test was explored to judge whether the distributions of the global entropy of low and high PBM networks were significantly different (20). We first permuted sample labels and recalculated the global entropy of the low PBM network, which was repeated L times. The significance level (P-value) of the tests was calculated by (#\(S_i^l\) \leq S_{obs}^l, \#\(S_i^H\) \leq S_{obs}^H) for l = 1,...,L)/L. \(S_i^l\) and S_{obs}^l are the global entropy of the low PBM network at the 1st time-point and of the high PBM network before this test, respectively. A P-value <0.05 was used to define significantly differential network in this study.

Animals and modeling. A total of 60 female SD rats were purchased from Hebei Medical University Animal Center. All rats were housed at room temperature with 12 h light-dark cycle, and fed with normal chow diet and free access to water. The rat ovariectomy experiment was approved by the Ethics Committee of Hebei Cangzhou Central Hospital, (Cangzhou, China).

After a week of feeding, all rats were randomly and on average separated into 2 groups in which rats were ovariectomized and sham operated at the age of 6 months. Three months after surgery, bone mineral density examination was used to verify if the model was successfully established. All rats in ovariectomized and sham groups were euthanized via exposure to gradually increasing concentrations of isoflurane and carbon dioxide gas (30% gradual-fill chamber vol/min), blood was drawn from the abdominal aorta of rats in each group after the blood pressure and the right femur and then the right femur and third lumbar spine were quickly removed. The number of animals used and their suffering was minimized. Surface-attached soft tissue were removed, eardrums, vertebral bows and other accessories were retained. The obtained bone tissue
was packaged by gauze soaked with saline, and kept at -80℃ in a refrigerator. Dual-energy X-ray absorptiometry (Lunar-DPX-IQ; GE Medical Systems, Madison, WI, USA) was used to detect bone density of right femur and third lumbar vertebra.

Reverse transcription-quantitative PCR (RT-qPCR). Venous blood sample of models were collected and total RNA was extract by TRIzol RNA kit (Invitrogen; Thermo Fisher Scientific, Inc., Waltham, MA, USA) according to the manufacturer's instructions. The concentration and purity of total RNA were measured by ultraviolet spectrophotometer. Reverse transcription into cDNA was then conducted by a reverse cDNA transcription kit (Takara Bio, Inc., Otsu, Japan). SYBR Ex Script RT-PCR kit (Takara Bio, Inc.) was used to detect the expression of target genes and the primers used in this study is shown in Table I.

Statistical analysis. GraphPad Prism version 5 software (GraphPad Software, Inc., La Jolla, CA, USA) was used for statistical analyses. All data were shown as mean ± standard deviation (SD). For experiments with two groups, t-test was performed. P<0.05 was considered to indicate a statistically significant difference. Experiments were repeated with a minimum of 3 times for each condition.

Results

Identification of modules. We used the expression profile data to construct PPINs for the low and high PBM group, respectively. Based on merging highly overlapping cliques, we captured the effect of differences in interaction weights between monocyte low and high PBM group via the weighted density-based ranking of cliques. After the simplification of the modules, 19 modules in low PBM network and 38 in high PBM network were obtained. Then module correlation density was calculated to identify sets of disrupting or altering module pairs between the two networks. A total of 66 modules were identified by modeling it as a maximum weight bipartite matching.

Characterizing differential networks from network entropy. We constructed 32 destination networks with modules divided from monocyte low and high PBM networks. The characteristics of 32 networks are shown in Table II. The network entropy was ranged from -3.09 to 0.87 and the max value of network entropy of PBM was 0.87 in network 11. Among them, network 11 was the only significantly differential one with P-value of 0.047 (P<0.05). Fig. 1 shows the differential PBM network with 8 nodes and 28 edges. All genes belonged to precursors of macropain, which were related to calcium transport as well as blood monocytes. PBM, peak bone mass; PPI, protein-protein interaction.

Bone density of osteoporosis models. As shown in Fig. 2, 3 months after operation, bone density of right femur and third lumbar vertebra in the osteoporosis group was significant.
Expression of target genes in osteoporosis models. The results of RT-qPCR showed that compared with sham controls, the expression of target genes in osteoporosis was obviously increased (Fig. 3). The results indicated that these precursors of osteoclasts, such as PSMD7, PSMC1, PSMC2, PSMD5, PSMD7, PSMD8 and PSMD11, were identified in both monocyte low and high PBM networks. They belonged to precursors of macropain, which were related to calcium transport as well as blood monocytes. In a previous study, the surplus/PSMD5 was found to inhibit the assembly and activity of 26S proteasome in human disease (32). Importantly, neuronal activity and calcium/calmodulin-dependent protein kinase II could regulate the expression of proteasome (33). Intracellular calcium mobilization regulates the activity of 26S proteasome during the metaphase-anaphase transition in meiotic cell cycle (34). At the same time, proteasome inhibitor lactacystin hinders the calcium homeostasis of dopamine neurons (35). Similar results were also obtained in this study, the results of RT-qPCR in this study showed that compared with sham controls, the expression of target genes in osteoporosis were obviously increased. In other words, the method of network entropy could be used to detect differential networks as indicator of disease. These 8 genes may play important roles in controlling inflammation of the osteoporosis. It suggested that the network is a novel therapeutic indicator for osteoporosis during the bone monocyte progression. These findings are helpful in disclosing the pathogenetic mechanisms of osteoporosis.

Discussion

Recently genome-wide gene expression profiles and PPIs hold the major promise to uncover the biological progression, which are subsequently pivotal in getting across molecular mechanisms of diseases (26). Based on the landscape, functional network entropy can be readily detected to facilitate the valuable application of PPIs on disease research. Previous studies have revealed that increased entropy appears to be a hallmark of cancer systems (27,28). Thus we utilized the previous hypothesis that the network entropy could discriminate the inflammatory network from the normal network (20). The systematic calculation and comparison were performed to test whether the network entropy can also discriminate the low PBM network from high PBM network. By applying a significance test for the difference in global network entropy of low and high PBM networks, we found the global network entropy of PBM networks is significantly differential (P<0.05). The result claimed that the network entropy provided good discrimination between low and high PBM networks. Moreover, this method can serve as a significant foundation for further exploring the molecular mechanisms of other diseases and developing control strategies.

One unique advantage of the present method compared with custom networks identifying analysis is that the entropy performs better than other topological metrics of the network in characterizing the inflammatory disease, which was detected by the Network Analyzer plug-in in Cytoscape (20,29). Thereby, the significantly different network we captured was more valuable and promising in understanding molecular mechanisms of diseases and reliable in therapeutic options.

Identifying key genes and modules from PPINs is of great help for uncovering the biological functions of genes in networks (30,31). In the present study, a total of 8 genes, PSMA2, PSMB1, PSMC1, PSMC4, PSMD5, PSMD7, PSMD8 and PSMD11, were identified in both monocyte low and high PBM networks. They belonged to precursors of macropain, which were related to calcium transport as well as blood monocytes. In a previous study, the surplus/PSMD5 was found to inhibit the assembly and activity of 26S proteasome in human disease (32). Importantly, neuronal activity and calcium/calmodulin-dependent protein kinase II could regulate the expression of proteasome (33). Intracellular calcium mobilization regulates the activity of 26S proteasome during the metaphase-anaphase transition in meiotic cell cycle (34). At the same time, proteasome inhibitor lactacystin hinders the calcium homeostasis of dopamine neurons (35). Similar results were also obtained in this study, the results of RT-qPCR in this study showed that compared with sham controls, the expression of target genes in osteoporosis were obviously increased. In other words, the method of network entropy could be used to detect differential networks as indicator of disease. These 8 genes may play important roles in controlling inflammation of the osteoporosis. It suggested that the network is a novel therapeutic indicator for osteoporosis during the bone monocyte progression. These findings are helpful in disclosing the pathogenetic mechanisms of osteoporosis.

Acknowledgements

Not applicable.

Funding

This study did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

Availability of data and materials

The datasets used and/or analyzed during the present study are available from the corresponding author on reasonable request.

Authors' contributions

GC and LM contributed to the conception and design of the study. LM and HD contributed significantly to the acquisition,
analysis and interpretation of data. LM, HD and GC performed the writing, review, and/or revision of this manuscript. All authors have read and approved the final manuscript.

Ethics approval and consent to participate

The rat ovariectomy experiment was approved by the Ethics Committee of Hebei Cangzhou Central Hospital, (Cangzhou, China).

Consent for publication

Not applicable.

Competing interests

The authors have no financial or personal relationships with other people or organizations that could inappropriately influence their work.

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