RESEARCH ARTICLE

Investigating multiple dys-regulated pathways in rheumatoid arthritis based on a pathway interaction network

Running title: Dys-regulated pathways in rheumatoid arthritis

Author: Xian-Dong Song¹; Xian-Xu Song²; Gui-Bo Liu³; Chun-Hui Ren⁴; Yuan-Bo Sun⁵; Ke-Xin Liu¹; Bo Liu¹; Shuang Liang⁴; Min Zhu⁴*

¹ Department of Orthopaedics, Hongqi Hospital of Mudanjiang Medical University, Mudanjiang 157000, Heilongjiang, People’s Republic of China

² Department of General Surgery, Second Affiliated Hospital of Mudanjiang Medical University, Mudanjiang 157000, Heilongjiang, People’s Republic of China

³ Department of Anatomy, Mudanjiang Medical University, Mudanjiang 157000, Heilongjiang, People’s Republic of China

⁴ Department of MRI, Hongqi Hospital of Mudanjiang Medical University, Mudanjiang 157000, Heilongjiang, People’s Republic of China

⁵ Department of Kidney Internal Medicine, Hongqi Hospital of Mudanjiang Medical University, Mudanjiang 157000, Heilongjiang, People’s Republic of China

*Corresponding author:
Min Zhu, Email: zhuminbio@126.com, Address: Department of MRI, Hongqi Hospital of Mudanjiang Medical University, No.5 Tongxiang Street, Aimin District, Mudanjiang 157000, Heilongjiang, People’s Republic of China

Keywords: rheumatoid arthritis, dys-regulated pathways, pathway interaction network
Abstract:

**Objective:** Traditional methods of identifying biomarkers in rheumatoid arthritis (RA) have focused on the differentially expressed pathways or individual pathways, which however neglected the interactions between pathways. To better understand the pathogenesis of RA, we aimed to identify dys-regulated pathway set using a pathway interaction network (PIN) which considered the interactions among pathways.

**Methods:** Firstly, RA-related gene expression profile data, protein-protein interactions (PPI) data and pathway data were recruited from the corresponding databases. Secondly, principal component analysis (PCA) method was used to calculate the pathway activity for each pathway, and then a seed pathway was identified using data gleaned from the pathway activity. A PIN was then constructed based on gene expression profile, pathway data, and PPI information. Finally, the dys-regulated pathways were extracted from the PIN based on the seed pathway using the method of support vector machines (SVMs) and an Area Under the Curve (AUC) index. **Results:** A PIN was constructed, which included 854 pathways and 1064 pathway interactions. The greatest change of the activity score between RA and control samples was observed in the pathway of epigenetic regulation of gene expression which was extracted and regarded as the seed pathway. Starting with this seed pathway, one maximum pathway set (MPS) containing 10 dys-regulated pathways was extracted from the PIN, having an AUC of 0.8249, and this result indicated that this pathway set could distinguish RA from the controls. **Conclusion:** These 10 dys-regulated pathways might be the potential biomarkers for RA diagnosis and treatment in the future.

**Keywords:** rheumatoid arthritis, dys-regulated pathways, pathway interaction network
1. Introduction

Rheumatoid arthritis (RA) is a chronic inflammatory joint disease, characterized by the burden of swollen joints, pain, and decreased physical functions (Michaud and Wolfe, 2007). Usually, patients with RA are at an increasing risk of mortality, with the survival rate decreasing by 3-10 years (Alamanos and Drosos, 2005). Although many biological agents have been developed to treat this disease, the curative treatment is still undiscovered yet (Andersen et al., 2016). Moreover, immune-modulatory properties of drugs also increase the risk of potential adverse events (such as nosocomial infections, congestive heart failure, and malignancy) which put a significant burden on the health care systems (Yamada et al., 2016). The risk of adverse effects and expensive treatment for RA patients have driven the seek for predictive signatures that can be used to detect and treat RA early.

Therefore, several scholars focused on the differentially expressed genes (DEGs) (Diogo et al., 2013) in order to reveal the etiology and explore the effective treatment modalities for RA. However, facing with the multi-factorial disease, RA, identifying biomarkers can not just rely on a single DEG, because a set of genes interact with each other (Glazier et al., 2002; Merikangas et al., 2006). Furthermore, for the same disorder, many of the gene biomarkers extracted in one dataset have not been found to work efficiently in another dataset (Braga-Neto and Dougherty, 2004). Based on the poor performance of gene biomarkers, many scholars focused on the pathways related to this disease which would improve the accuracy when the pathways are considered as biomarkers, compared with these gene biomarkers. For example, Cheng et al. (CG et al., 2013) found that Wnt signaling pathway played an important role in the pathogenesis of RA. Traditional pathway analysis mainly focused on the single dys-regulated pathways, but the most important point neglected the interactions
between pathways (Khatri et al., 2012). Generally, the functions related pathways are interconnected with each other, and any of the dys-regulated pathways could affect the activities of the other. Remarkably, detecting and understanding the interactions between pathways is beneficial for exploring the molecular mechanisms of disease (Li and Agarwal, 2009). Notably, network-based biology is broadly used to analyze interactions, in turn further shedding lights on the action mechanism of how cellular systems operate (Barabasi and Oltvai, 2004; Xia et al., 2004). As a result, a pathway interaction network (PIN) was built to identify dys-regulated pathways, which considered the interactions between pathways (Liu et al., 2012).

Herein, in our study, a computational method was used to detect significant pathways by constructing a PIN. The findings demonstrate that this method is beneficial in predicting biomarkers and even drug targets for RA in a more robust fashion.

2 Materials and methods

2.1 Retrieving datasets and preprocessing

2.1.1 Gene expression profile data

In this paper, the gene expression profile data (E-GEOD-57405) (Rosenberg et al., 2014) were obtained from the ArrayExpress database, which were deposited in the A-MEXP-1171-Illumina HumanHT-12v3.0 Expression BeadChip platform. In detail, E-GEOD-57405 was comprised of 46 samples, including 19 normal samples and 27 RA samples. In order to make the quality of the gene expression profile data more accurate, the standard pre-processing steps (including scale transformations, management of missing values, replicated handing, flat pattern filtering and pattern standardization) were conducted (Herrero et al., 2003). Then, the probes were mapped
to the human genomics to further identify gene symbols. Ultimately, a total of 7352 genes were obtained for further study. Next, the expression levels of all genes were standardized.

2.1.2 Protein-protein interaction (PPI) data

PPIs provide a valuable framework for a better understanding of the functional organization of proteomes and are crucial for all biological processes (Stelzl et al., 2005). All human PPI data was obtained from the Search Tool for the Retrieval of Interacting Genes/Proteins (STRING, http://string-db.org/) which has been designed to assemble, evaluate and disseminate protein-protein interaction information, in a user-friendly and comprehensive manner (Franceschini et al., 2013). Based on the STRING database, 16,730 genes and 787,896 interactions were retrieved. To increase the reliability of these PPIs, only the interactions with confidence score > 0.5 were selected to establish the informative PPI network. Then, taking the intersections of informative PPI network and gene expression profile, a total of 26,855 protein interactions among 4890 genes were obtained to construct the background PPI network for further analysis.

2.1.3 Pathway data

The biological pathways for human beings were obtained from the Reactome (http://www.reactome.org) pathway database. This database actually is an expert-authored, peer-reviewed knowledgebase of human reactions and pathways, that functions as a data mining resource and electronic textbook (Matthews et al., 2009). A total of 1675 pathways were collected from this database. Because different pathways consist of different number of genes, some having too few genes might not have enough biological information, and some having too many genes might be too generic (Ahn et al., 2014). Hence, only pathways with a gene set size ranging from 5 to 100
were reserved as the study objectives. Finally, a total of 854 pathways were selected.

2.2 Pathway activity calculation and PIN construction

2.2.1 Pathway activity calculation

After filtering procedures for pathways, only those genes that were mapped to the 854 pathways were reserved for further analysis. After the genes were aligned to the 854 pathways, an activity score for each pathway was defined as the sum of the expression levels of all genes enriched in this given pathway using the PCA method. PCA method is a mathematical algorithm that reduces the dimensionality of data while retaining most of the variation in the data sets (Hotelling, 2010; Ringn and Eacute, 2008). In particular, the first principal component was determined as the activity score for the corresponding pathway. Thus, there were different activity scores for each pathway in the disease samples and controls. That meant that the activity score for a corresponding pathway between RA and controls was different, and the difference implied the correlation to RA development. A bigger difference in activity score may lead to closer correlation of the pathway to the disease. Therefore, the pathway with the biggest change in activity score between the two groups was regarded as the most important pathway, and was denoted as the seed pathway.

2.2.2 PIN construction

The interactions between the correlative pathways constituted a network which was called PIN. In the PIN, each node represented a pathway, and one edge was laid between two pathways if they shared at least one gene or there were interactions between genes from the two pathways on the basis of PPIs. The PIN is important not only to understand the drug response, but also for the development of novel drugs and therapy in human disease (Song et al., 2014). For the sake of constructing the PIN, two conditions should be met.
Firstly, based on the gene expression data and student’s t-test, DEGs were determined using p-value < 0.05 between RA and controls. In the process of establishing the PIN, it was necessary that there was at least one common DEG between two pathways.

Secondly, in order to measure whether a pair of interacting proteins used to lay an edge between two pathways was highly co-expressed, a Pearson correlation coefficient (PCC) for all PPIs between RA and control samples was calculated. The distribution of PCC was obtained after the correlation strength between the two pathways was evaluated. The PCC determines the strength of linear association between two variables, and it is measured on a scale with no units and can take a value from -1 through 0 to +1 (Sedgwick, 1996). The absolute difference in PCC values for the PPIs in RA and control groups was then calculated. In this study, the weight score for a pathway interaction was determined as the total |PCC| values of all genes. The other condition was that the two genes coding a pair of interacting proteins employed to lay an edge between two pathways were highly co-expressed (the absolute value of PCC > 0.8) between genes in the two pathways. If not, the edges between the two pathways were removed.

Based on the above two conditions, an original PIN was built. In order to simplify the pathway network, the score values of each pathway interactions in PIN, eg. the sum of the absolute values of PCC for the PPIs in every two pathways, were computed, and then the pathway interactions were ranked in a descending order based on the score values. Next, the top 5% of pathway interactions were selected to construct an informative PIN for RA to further identify dys-regulated pathways.

2.2.3 Identifying dys-regulated pathway from the informative PIN

The informative PIN that was extracted above increased the difficulty in
distinguishing the diseases from the controls, which had a large number of genes and pathways. Hence, in order to make the differentiation more accurately, detecting a minimum set of pathways (MSP) which were considered to be more possibly dys-regulated pathways from the PIN might be the best way to discriminate RA samples from controls.

As mentioned above, the seed pathway that had the biggest change activity score was the first pathway biomarker used to discriminate between diseases and controls. Subsequently, the second pathway was selected from the available pathways and was added to the first pathway to get a better classification ability, which interacted with the first pathway. This process was repeated to add new pathways to extract pathway biomarkers, until no more pathways could be added to enhance the accuracy of the classification. The final selected pathway sets were regarded as potential MSP in RA. During the process, support vector machines (SVMs), a widely used kernel based method especially useful for smaller number of samples with high dimensional variables (Liu et al., 2012), was utilized to select the dys-regulated pathways. Utilizing a five-fold cross validation, the classification performance was evaluated, and an Area Under the Curve (AUC) value was adopted as the classification performance index. With the goal of obtaining robust results, the five-fold cross-validation was repeated for 100 times and the mean value of classification accuracy was defined as the final result.

3. Results

3.1 PIN establishment

Utilizing student’s t-test, a total of 807 DEGs were obtained with a p-value of <
0.05. In order to establish the PIN, the edges between any two pathways were randomly selected, provided that at least one of the common genes in both pathways was differentially expressed in the two groups, or the two genes coding a pair of interacting proteins used to lay an edge between the two pathways were highly co-expressed (PCC absolute value > 0.8). Ultimately, an original PIN covering a total of 21,281 pathway-pathway interactions was established. However, due to the complicated scale of the network, the sum of the absolute values of PCC for the PPIs in every pair of pathways was calculated, and only the top 5% of pathway interactions were extracted to construct the informative PIN to identify dys-regulated pathways. Finally, a total of 1064 pathway interactions were collected to build the informative PIN.

3.2 Seed pathway

After analyzing 854 pathways using the PCA, it was discovered that different activity scores existed in pathways between the disease and normal controls. In order to identify the most important pathway, referred to as the pathway related to the disease, the activity scores of 854 pathways between disease and normal controls were analyzed. Meanwhile, to make the changes in scores more intuitively, the regularities in distribution of the activity score was illustrated in Figure 1. This figure showed that the greatest absolute change of activity score in the pathways was located on the top point, which was the pathway of the epigenetic regulation of gene expression (absolute change of activity score = 4.8836), and was defined as the seed pathway.

3.3 Identification of dys-regulated pathways

Taking the seed pathway of epigenetic regulation of gene expression as start, dys-regulated pathways were detected based on the increased classification accuracy. Ultimately, an MSP (including 10 dys-regulated pathways) with AUC of 0.8249 was
obtained from the informative PIN, which was regarded as the most possibly dys-regulated pathways. Good performance indicated that the identified dys-regulated pathways might be acted as robust biomarkers of RA. The illustrative diagram of the network of these 10 pathways was shown in Figure 2.

Additionally, the genes annotated in the identified dys-regulated pathways were compared with the DEGs (Table 1). From this Table, it can be seen that only a small fraction (from 3.5% to 17.6%) of the genes enriched in the dys-regulated pathways overlapped with the DEGs. This phenomenon further suggested that the pathway as an entity might have a better propensity to diagnose complex diseases rather than individual genes, even if the genes enriched in the pathway were not differentially expressed. From this table, we found that the seed pathway of epigenetic regulation of gene expression owned the largest number of DEGs (N = 9), and the pathway of DNA replication had the second largest number of DEGs (N = 7). Another dys-regulated pathway, Toll-like Receptor 2 (TLR2) Cascade was enriched by relatively much more DEGs (N = 6), and served as an immunological pathway. Significantly, TLRs, particularly TLR2, play a key action in RA (Seibl et al., 2003), which is specific for the immune-related diseases. RA is a chronic autoimmune disease, characterized by cytokines-mediated inflammation of the synovial lining of joints.

4. Discussion

RA is a common systemic autoimmune disorders characterized by inflammation of the synovial tissue (de Hair et al., 2014). Although it has been reported that T cells, B cells, the orchestrated interaction of pro-inflammatory cytokines and Wnt signaling pathway significantly participate in the RA pathogenesis (CG et al., 2013; Choy, 2012), the etiology of this disease is still unknown (CG et al., 2013). In addition, this disease is associated with the risk of morbidity and mortality,
especially an increase in mortality rate due to the increased risk of cardiovascular events (Avinazubieta et al., 2012; Roubille et al., 2015). With the advent of the era of genes, understanding the diseases at the molecular level will help us pinpoint the root cause of these diseases. Hence, it is considered that biological functions do not just rely on a single gene or an individual pathway, but also the interactions between genes or pathways. Meanwhile, as the interactions between the function-related genes or pathways constitute a large network, it has led to confusion among scholars as to which pathways are closely related to the disease. Consequently, in order to determine the most closely related pathways about RA, the MSP was extracted from the PIN, which was denoted as the dys-regulated pathways. Compared with the traditional methods that focused on a single pathway (Begovich et al., 2004; Choy, 2012), our method considered the interactions between pathways, and extracted the minimum set of interactions in the pathways from the PIN. These would serve as better biomarkers for the disease and would help us to diagnose the disease (Okada et al., 2014).

In this paper, a total of 10 dys-regulated pathways were obtained from the PIN, among which epigenetic regulation of gene expression was designated as the seed pathway. Epigenetic gene regulation refers to different states of phenotypic expression caused by differential effects of chromosome or chromatin packaging (Hendrich and Willard, 1995). Emerging evidences have suggested that the epigenetics plays a key role in human pathologies, including in inflammatory disorders, and such epigenetic factors may be important in understanding the origins of interindividual variations in the inflammatory response (Wilson, 2008). RA is the most common inflammatory disease (Firestein, 2003). In addition, Emmanuel et al. (Karouzakis et al., 2009) have discovered that three main mechanisms of epigenetic control, including DNA
methylation, histone modifications and microRNA activity, interact in the development of the RA synovial fibroblasts (Karouzakis et al., 2009). Therefore, the results of this study suggested that the pathway of epigenetic regulation of gene expression might play an important part in the progression of RA.

Another dys-regulated pathway, Toll-like Receptor 2 (TLR2) Cascade, was also identified, which was enriched by relatively much more DEGs, and served as an immunological pathway. TLR2 is a member of TLRs which have been demonstrated to mediate the activation of NF-κB, thereby leading to the generation of mediators in the immune system such as TNF-α and IL-1β (Takeuchi et al., 2000; Wang et al., 2001). Further, proinflammatory cytokines TNF-α and IL-1β stimulates cultured synovial fibroblasts (SFs) to cause the significant increase of TLR2 (Seibl et al., 2003). The activation of SFs has been found to be an important characteristic in RA. An earlier study has implicated that TLR2-dependent mechanisms induce the activation of synovial cells, possibly resulting in the destruction of cartilage, accounting for the etiology of RA (Kyburz et al., 2003). In addition, Pierer et al. (Pierer et al., 2004) have indicated that chemokine secretion by activating SFs using TLR-2, contribute to the development of RA. Accordingly, our result support the strategy of targeting the pathway of Toll-like Receptor 2 (TLR2) Cascade to suppress joint inflammation in RA patients.

Consequently, unlike the previous method, this integration-based analysis had several merits. Firstly, we paid attention to the functional dependency between pathways via establishing a PIN, thereby indicating the robustness of the identified pathway biomarkers. Secondly, when pathways had marginal p-values, they still might bring a stronger signal if these pathways could form a cluster in the PIN. The results of our study suggested that dys-regulated pathways, especially epigenetic
regulation of gene expression and Toll-like Receptor 2 (TLR2) Cascade might be important in RA initiation, development and progression. However, there were still several limitations in this study. To begin with, the sample size was a little small. Secondly, this study was conducted based on existing data using only bioinformatics method, yet the findings lacked experimental validation. Although there were limitations, the findings of this study provided some preliminary evidence to uncover alternative candidate therapeutic strategies for RA. These 10 dys-regulated pathways might be the potential biomarkers for RA diagnosis and treatment in the future.

Reference


Diogo D., Kurreeman F., Stahl E., Liao K., Gupta N., Greenberg J. et al. 2013 Rare,
Low-Frequency, and Common Variants in the Protein-Coding Sequence of Biological Candidate Genes from GWASs Contribute to Risk of Rheumatoid Arthritis. American Journal of Human Genetics. 92, 15-27.


Rosenberg A., Fan H., Chiu Y.G., Bolce R., Tabechian D., Barrett R. et al. 2014 Divergent gene activation in peripheral blood and tissues of patients with rheumatoid arthritis, psoriatic
arthritis and psoriasis following infliximab therapy. Plos One. 9, e110657-e110657.


Table 1 Dys-regulated pathways extracted from the informative PIN, and the DEGs involved in dys-regulated pathways

<table>
<thead>
<tr>
<th>Index</th>
<th>Pathway</th>
<th>Number of genes</th>
<th>Number of DEG</th>
</tr>
</thead>
<tbody>
<tr>
<td>223</td>
<td>Epigenetic regulation of gene expression</td>
<td>51</td>
<td>9</td>
</tr>
<tr>
<td>597</td>
<td>Removal of licensing factors from origins</td>
<td>37</td>
<td>3</td>
</tr>
<tr>
<td>186</td>
<td>DNA replication</td>
<td>57</td>
<td>7</td>
</tr>
<tr>
<td>168</td>
<td>Degradation of beta-catenin by the destruction complex</td>
<td>48</td>
<td>4</td>
</tr>
<tr>
<td>76</td>
<td>Autodegradation of cdh1 by cdh1:APC/C</td>
<td>34</td>
<td>2</td>
</tr>
<tr>
<td>48</td>
<td>Antigen processing-cross presentation</td>
<td>48</td>
<td>4</td>
</tr>
<tr>
<td>386</td>
<td>M/G1 transition</td>
<td>46</td>
<td>4</td>
</tr>
<tr>
<td>97</td>
<td>Cdc20: phospo-APC/C mediated degradation of cyclin A</td>
<td>38</td>
<td>5</td>
</tr>
<tr>
<td>756</td>
<td>Toll like receptor 2 (TLR2) cascade</td>
<td>39</td>
<td>6</td>
</tr>
<tr>
<td>98</td>
<td>CDK-mediated phosphorylation and removal of Cdc6</td>
<td>29</td>
<td>1</td>
</tr>
</tbody>
</table>

Note: PIN: pathway interaction network; DEG: differentially expressed genes; “pathway IDs (Indexes)” are defined based on alphabetical order.

Figure legends

Figure 1: Distribution of the absolute changes in the activity score of 854 pathways.

The pathway with the greatest change in activity score between rheumatoid arthritis (RA) sample and normal control was considered to be the seed pathway.
Figure 2: Altered pathway interaction network in RA, involving 10 dys-regulated pathways which were assembled into a pathway network on the basis of the interactions. Each node stood for a pathway. Red node denoted seed pathway. Blue ones was the dys-regulated pathways interacted with the seed pathway. The number represented the pathway ID defined in this paper, in alphabetical order.