Identification of Candidate Genes and MicroRNAs for Acute Myocardial Infarction by Weighted Gene Coexpression Network Analysis

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Background. Identification of potential molecular targets of acute myocardial infarction is crucial to our comprehensive understanding of the disease mechanism. However, studies of gene coexpression analysis via jointing multiple microarray data of acute myocardial infarction still remain restricted. Methods. Microarray data of acute myocardial infarction (GSE48060, GSE66360, GSE97320, and GSE19339) were downloaded from Gene Expression Omnibus database. Three data sets without heterogeneity (GSE48060, GSE66360, and GSE97320) were subjected to differential expression analysis using MetaDE package. Differentially expressed genes having upper 25% variation across samples were imported in weighted gene coexpression network analysis. Functional and pathway enrichment analyses were conducted for genes in the most significant module using DAVID. The predicted microRNAs to regulate target genes in the most significant module were identified using TargetScan. Moreover, subpathway analyses using iSubpathwayMiner package and GenCLiP2.0 were performed on hub genes with high connective weight in the most significant module. Results. A total of 1027 differentially expressed genes and 33 specific modules were screened out between acute myocardial infarction patients and control samples. Ficolin (collagen/fibrinogen domain containing) 1 (FCN), CD14 molecule (CD/one.fitted/four.fitted), S100 calcium binding protein A9 (S/one.fitted/zero.fitted/zero.fittedA/nine.fitted), and mitochondrial aldehyde dehydrogenase 2 (ALDH/two.fitted) were identified as critical target molecules; hsa-let-/seven.fittedd, hsa-let-/seven.fittedb, hsa-miR-/one.fitted/two.fitted/four.fitted-/three.fitted, and hsa-miR-/nine.fitted-/one.fitted were identified as potential regulators of the expression of the key genes in the two biggest modules. Conclusions. FCN, CD14, S100A9, ALDH2, hsa-let-7d, hsa-let-7b, hsa-miR-124-3, and hsa-miR-9-1 were identified as potential candidate regulators in acute myocardial infarction. These findings might provide new comprehension into the underlying molecular mechanism of disease.

1. Introduction

Acute myocardial infarction (AMI) is characterized by definite evidence of myocardial necrosis in a clinical background of acute myocardial ischemia [1]. According to the report of global burden of cardiovascular disease in 2015, there were an estimated 7.29 million AMI in the world [2], which contributed to a high morbidity and mortality of global health. AMI has a high risk of death due to congestive heart failure and malignant arrhythmia, although many endeavors and monies have been spent on developing new therapies, it remains a major challenge for clinicians to prevent the adverse cardiac events and cure this disease. Established risk factors, such as hypertension, diabetes, hypercholesterolemia, smoking, and obesity, cannot explain all the risk on morbidity and mortality of AMI, and, moreover, substantial numbers of patients have an inexplicit etiology in myocardial damage [3]. Therefore, urgent research to discover potential pathogenesis
of AMI and exploit novel medication targets and therapeutic strategies is needed, thereby reducing the threat of this disease to human life.

The clinical manifestations and characteristics of AMI are heritable features, and transcriptomics-based screening of genetic biomarker is helpful in early recognition of risk carriers and improving diagnosis and treatment of AMI. It is a general recognition that expression microarrays have advantages of rapid unbiased screening and extensive coverage of nearly all transcripomes to reveal the most promising targets. IL-1RL-1, Interleukin 1 receptor-like 1 (ST2) should be a representation of cardiac biomarker which was identified as a target at initial microarray analyses and then resulted in the development of suitable assay [4]. Recently, microRNAs (miRNAs) as noncoding small RNAs are involved in a broad range of regulation for biological processes and disease development, and also several evidences have shown that circulating miRNAs are stable and can be used as novel diagnostic markers for AMI [5–10].

Coexpression networks as transcriptomic technologies have grown in popularity since they allow for the integration of large transcriptional data sets, as well as cointaneous identification, clustering and analysis of thousands of genes with similar expression patterns across a wide range of conditions [11]. Weighted Gene Coexpression Network Analysis (WGCNA) has been established by means of introducing several adjacency functions that convert the coexpression measure to a connection weight, and furthermore the parameters of the adjacency function have been determined by the scale-free topology criterion [12]. WGCNA provides a powerful network-based strategy to expedite the clarification of molecular mechanisms underlying important biological processes and applies to a diverse range of human disease researches. Accordingly, WGCNA could be used to analyze AMI microarray data sets in this study.

So far gene expression studies of AMI have been limited in sample size and lack of myocardial tissue samples from patients. On the other hand, development and utility of circulating biomarkers would be optimum to noninvasive diagnosis and early identification of AMI. We combined several blood-based microarray expression data to screen out mutual differentially expressed genes (DEGs) among data sets. WGCNA was used to construct gene coexpression network based on DEGs profiling, and significant modules and hub genes were detected by the WGCNA as well. Furthermore, miRNAs that could be predicted to regulate DEGs in the most significant module were identified and miRNAs-DEGs regulatory relationships were analyzed. This study aimed to detect out more candidate genes and miRNAs that were involved in the pathogenesis and progression of AMI, among which some genetic biomarkers might be converted into the promising targets for the diagnosis and treatment of AMI.

2. Materials and Methods

2.1. Source of Microarray Data. Four microarray data, including GSE48060 (USA; 31 patients with AMI and 21 controls), GSE66360 (USA; 49 patients with AMI and 50 controls), GSE97320 (China; 3 patients with AMI and 3 controls), and GSE19339 (Switzerland; 4 patients with AMI and 4 controls), were downloaded from Gene Expression Omnibus (GEO http://www.ncbi.nlm.nih.gov/geo/) database, which were based on the platform of GPL570 Affymetrix Human Genome U133 Plus 2.0 Array (Affymetrix Inc., Santa Clara, California, USA). This microarray meta-analysis made use of four data sets based on 156 blood samples derived from 87 AMI patients and 78 control subjects.

2.2. Data Preprocessing. The raw array data (CEL files) were imported into Expression Console software and subjected to background adjustment, quantile normalization, and log₂ transformation by Robust Multiarray Average (RMA) [13]. After that, probe identifiers (IDs) were transformed into gene symbols according to the annotation files, and the average expression value of multiple probes corresponding to one same gene was calculated as the single expression value of this gene.

2.3. Quality Control of Microarray Data and Differential Expression (DE) Analysis. Six quality control (QC) indices provided in MetaQC package of R software were used to assess quality of selected studies, which included internal homogeneity of coexpression structure among studies (IQC), external consistency of coexpression pattern with pathway database (EQC), accuracy and consistency of differentially expressed gene detection (AQCg and CQCg), and enriched pathway identification (AQCp and CQCp) [14]. By applying this procedure, we could filter out the low-quality studies and eliminate the biases among datasets. Moreover, principal component analysis (PCA) biplots and standardized mean ranks were provided assisting in the detection of deflected studies. MetaDE package in R software provided 12 major statistical methods for differential expression (DE) analysis [14]. We chose DEGs between disease group and control group using the MetaDE.pvalue algorithm of MetaDE package. A false discovery rate (FDR) of < 0.05 was considered as the cut-off for the detection of DEGs.

2.4. Gene Ontology (GO) and Pathway Enrichment Analysis. Gene ontology (GO) is a tool for gene annotation using a dynamic, controlled vocabulary that classifies genes into three categories, including biological process, molecular function, and cellular component [15]. Kyoto Encyclopedia of Genes and Genomes (KEGG) database is used to assign gene sets to specific pathway maps of molecular interactions, reactions, and relation networks [16]. We performed GO functional annotation and KEGG pathway enrichment analyses of DEGs by the Database for Annotation, Visualization and Integrated Discovery (DAVID) [17].

2.5. Weighted Gene Coexpression Network Construction. We conducted gene coexpression analysis of DEGs in order to check out more potential genes associated with AMI. For reducing the deviation, we chose DEGs having upper 25% variation across samples to implement subsequent weighted gene coexpression network analysis (WGCNA).
WGCNA was used to construct weighted adjacency matrix that reported the connection strength between gene pairs. For weighted networks, the concordance of gene expression was measured with the Pearson correlation matrix, and the absolute value of the correlation coefficient $s_{ij} = |\text{cor}(x_i, x_j)|$, where $x_i$ and $x_j$ were the vectors of the expression value of gene $i$ and gene $j$. Then, the Pearson correlation matrix was transformed continuously with the power adjacency function into weighted gene network, and the adjacency matrix $a_{ij}$ was calculated as $a_{ij} = s_{ij}^\beta$. Here, the exponential $\beta = 20$ were chosen by scale-free topology criterion (Supplementary Figure 1). For detecting gene coexpression modules, the adjacency matrix was converted into the topological overlap matrix (TOM). The following formula was used to compute the weighting coefficient $W_{ij}$ (referred to TOM):

$$W_{ij} = \frac{l_{ij} + a_{ij}}{\min \{k_i, k_j\} + 1 - a_{ij}}$$

where $k_i$ indicated the total connectivity of gene $i$ with all other genes in the weighted network, $W_{ij}$ was considered the topology overlap between neighbor genes of $i$ and $j$. The topological overlap dissimilarity $(1 - \text{TOM})$ was used as input of hierarchical clustering. Modules were defined as branches of a hierarchical clustering dendrogram using the average linkage hierarchical clustering coupled with the topological overlap dissimilarity measure. For each module, the module eigengene (ME) was summarized by the first principal component of the module expression levels, and the correlation between each gene expression values and module eigengene was defined as the module membership (MM).

A gene significance (GS) was measured as minus log of a $p$-value with the $T$-test. The module significance (MS) was determined as the mean absolute GS for all genes in a given module.

2.6. Module Preservation Statistics. Module preservation statistics are used to verify whether an identified module in the reference network can be found in the test network. We investigated the preservation of coexpression network between the reference data set (meta-analysis) and the test data set (GSE123487) using the network-based Zsummary calculated by the module preservation function from WGCNA [18]. The statistics are calculated twice: once to evaluate whether modules are reproducible in the reference data set consisting only of genes in common with the test data set (called Zsummary.qual as “quality” statistics) and the second time to assess the conservation of the modules in the test data set (called Zsummary.pres as “preservation” statistics). $Z\text{summary.pres} < 2$ implies no evidence for module preservation, $2 < Z\text{summary.pres} < 10$ implies weak to moderate evidence of preservation, and $Z\text{summary.pres} > 10$ implies strong evidence that the module is preserved. $Z\text{summary.qual}$ was a complementary statistic to evaluate the robustness of the identified modules.

2.7. Screening of Hub Genes and Subpathway Analysis. In general, the genes with the largest number of connections are the most important genes in a module. The hub gene is termed as an abbreviation of “highly connected gene” that tends to have high connectivity in a coexpression module. Additionally, the hub genes were described as the genes most closely associated with disease. Intramodular connectivity (IC) corresponds to the connection degree of a gene with other genes in a given module. Here, the hub genes were considered as the highly connected genes according to their characterization with high IC, high MM, and high GS. After screening of the hub genes, a web-based text-mining server of GenCLiP 2.0 (http://ci.smu.edu.cn/GenCLiP2/analysis.php) was used to identify biological functions and molecular interactions in the hub genes list. The iSubpathwayMiner provides the k-clique method for identification of metabolic subpathways associated with studying disease based on the interested gene sets [19]. We used iSubpathwayMiner in mining the most relevant metabolic subpathways of the hub genes by merging information from genes and metabolites. The significantly enriched subpathways were detected by hypergeometric test ($p$-value < 0.05).

2.8. MicroRNAs (miRNAs) Target Prediction. To identify putative miRNA binding sites at the 3′-untranslated regions (3′-UTR) of mRNAs, we conducted miRNAs target prediction using the TargetScanHuman 7.2 algorithm. The outputs of prediction are ranked based on either the predicted efficacy of targeting (context++ scores) [20] or the probability of conserved targeting ($P_{CT}$) [21]. Here, the combination of two measures on context score percentile $\geq 90$ and aggregate $P_{CT}$ score $\geq 0.8$ was used to determine effective miRNAs. The
Table 1: Microarray data information and quality control measure summary.

<table>
<thead>
<tr>
<th>GEO accession</th>
<th>Platform</th>
<th>Probe number</th>
<th>Country</th>
<th>AMI sample</th>
<th>Control sample</th>
<th>IQC</th>
<th>EQC</th>
<th>CQCg</th>
<th>CQCp</th>
<th>AQCg</th>
<th>AQCp</th>
<th>SMR</th>
</tr>
</thead>
<tbody>
<tr>
<td>GSE97320</td>
<td>GPL570 / Affymetrix HU133 Plus 2.0</td>
<td>54,676</td>
<td>China</td>
<td>3</td>
<td>3</td>
<td>5.62</td>
<td>2.92</td>
<td>41.88</td>
<td>65.55</td>
<td>27.32</td>
<td>22.53</td>
<td>1.67</td>
</tr>
<tr>
<td>GSE66360</td>
<td>GPL570 / Affymetrix HU133 Plus 2.0</td>
<td>54,676</td>
<td>USA</td>
<td>49</td>
<td>50</td>
<td>3.30</td>
<td>4.00</td>
<td>17.76</td>
<td>59.30</td>
<td>16.73</td>
<td>34.49</td>
<td>2.08</td>
</tr>
<tr>
<td>GSE48060</td>
<td>GPL570 / Affymetrix HU133 Plus 2.0</td>
<td>54,676</td>
<td>USA</td>
<td>31</td>
<td>21</td>
<td>5.62</td>
<td>2.92</td>
<td>22.33</td>
<td>12.49</td>
<td>16.70</td>
<td>3.55</td>
<td>3.00</td>
</tr>
<tr>
<td>GSE19339</td>
<td>GPL570 / Affymetrix HU133 Plus 2.0</td>
<td>54,676</td>
<td>Switzerland</td>
<td>4</td>
<td>4</td>
<td>1.30*</td>
<td>4.00</td>
<td>4.92</td>
<td>39.73</td>
<td>4.49</td>
<td>13.15</td>
<td>3.25</td>
</tr>
</tbody>
</table>

GEO: Gene Expression Omnibus; * represents nonstatistical significance of quality control measures; AMI: acute myocardial infarction; IQC: internal quality index; EQC: external quality index; CQCg and CQCp: consistency quality control indexes; AQCg and AQCp: accuracy quality control indexes; SMR: standardized mean rank.

3. Results

3.1. Screening of DEGs. In order to eliminate problematic study and obtain reliable data, we performed MetaQC analysis of four data sets (GSE48060, GSE66360, GSE97320, and GSE19339) downloaded from GEO database. Six QC indexes and the standardized mean rank summary (SMR) scores were created by R software. The QC measures indicated that a study (GSE19339) had relatively low correlation with other three studies (Table 1). Coupled with the visualization of PCA biplots describing that a study on the opposite side of arrows had large SMR scores (Figure 1), the data set of GSE19339 was finally excluded from meta-analysis. DEGs were selected by MetaDE analysis. A total of 1027 DEGs were identified between AMI group and control group under the threshold of FDR < 0.05 (Supplementary Figure 2).

3.2. WGCNA Network Construction and Key Modules Detection. One hundred and fifty-seven samples with disease phenotypes were involved in gene coexpression analysis. For reducing the deviation, we selected DEGs having upper 25% variation across samples to implement WGCNA. To construct a weighted network, the lowest power of $\beta = 20$ ($R^2 \geq 0.8$) was chosen for a criterion of scale-free topology. By the means of the average linkage hierarchical clustering, two hundred and fifty-seven DEGs were grouped into a total of 33 modules (Figure 2). We selected the first largest module (56 DEGs clustered in turquoise module) and the second largest module (45 DEGs clustered in blue module) as the key (interesting) modules for further analysis. For each key module, the MS value was used to test their association with predicted results of miRNAs were validated with data set of GSE123487.
the disease. That the turquoise module had a higher MS value (MS_{turquoise} = 5.71) than the blue module (MS_{blue} = 2.64) suggested that the turquoise module had stronger correlation to AMI. In addition, the relevance between ME per key module and the disease status was also measured via calculation of Pearson's correlation coefficient and the significant p-value. The turquoise module (r = 0.40, p-value = 1.73E-07) was still considered to be highly relevant to AMI than the blue module (r = -0.27, p-value = 7.35E-04) (Supplementary Figure 3). We conducted statistical comparisons of correlations between the turquoise module and the blue module with AMI. The calculations relied on the tests implemented in the package cocor for the R programming language. The results of a comparison between two correlations showed that there was statistically significant difference (z=4.71, p-value <0.001). Subsequently, we conducted functional and pathway enrichment analysis for all DEGs clustered in the turquoise module. The mainly enriched results of significant functions and pathways were described in Table 2.

### 3.3. Identification of Hub Genes Associated with AMI

To identify intramodular hub genes, we computed the IC, MM, and GS for each gene in the interesting module. Through comparing with these measures, we found a positive correlation between MM and IC, but not GS. So, we adopted the IC and MM values as selective measures of the hub
Table 2: Top significantly GO function and KEGG pathway enrichment analysis in the turquoise module.

<table>
<thead>
<tr>
<th>Category</th>
<th>ID</th>
<th>Term</th>
<th>Count</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>GO_BP</td>
<td>GO:0006954</td>
<td>inflammatory response</td>
<td>11</td>
<td>7.50E-08</td>
</tr>
<tr>
<td>GO_BP</td>
<td>GO:0038124</td>
<td>toll-like receptor TLR6:TLR2 signaling pathway</td>
<td>3</td>
<td>4.58E-05</td>
</tr>
<tr>
<td>GO_BP</td>
<td>GO:0071726</td>
<td>cellular response to diacyl bacterial lipopeptide</td>
<td>3</td>
<td>4.58E-05</td>
</tr>
<tr>
<td>GO_BP</td>
<td>GO:0031663</td>
<td>lipopolysaccharide-mediated signaling pathway</td>
<td>4</td>
<td>9.63E-05</td>
</tr>
<tr>
<td>GO_BP</td>
<td>GO:0002755</td>
<td>MyD88-dependent toll-like receptor signaling pathway</td>
<td>4</td>
<td>1.06E-04</td>
</tr>
<tr>
<td>GO_BP</td>
<td>GO:0006955</td>
<td>immune response</td>
<td>8</td>
<td>1.57E-04</td>
</tr>
<tr>
<td>GO_BP</td>
<td>GO:0045087</td>
<td>innate immune response</td>
<td>8</td>
<td>1.78E-04</td>
</tr>
<tr>
<td>GO_BP</td>
<td>GO:0071223</td>
<td>cellular response to lipoteichoic acid</td>
<td>3</td>
<td>2.73E-04</td>
</tr>
<tr>
<td>KEGG Pathway</td>
<td>hsa05134</td>
<td>Legionellosis</td>
<td>3</td>
<td>1.60E-02</td>
</tr>
<tr>
<td>KEGG Pathway</td>
<td>hsa04145</td>
<td>Phagosome</td>
<td>4</td>
<td>1.71E-02</td>
</tr>
<tr>
<td>KEGG Pathway</td>
<td>hsa04640</td>
<td>Hematopoietic cell lineage</td>
<td>3</td>
<td>3.74E-02</td>
</tr>
<tr>
<td>KEGG Pathway</td>
<td>hsa04915</td>
<td>Estrogen signaling pathway</td>
<td>3</td>
<td>4.92E-02</td>
</tr>
</tbody>
</table>

Table 3: The hub genes in the turquoise and blue module.

<table>
<thead>
<tr>
<th>Module</th>
<th>Gene Symbol</th>
<th>Gene Title</th>
<th>p-value</th>
<th>IC</th>
<th>MM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Turquoise</td>
<td>FCN1</td>
<td>ficolin (collagen/fibrinogen domain containing) 1</td>
<td>6.49E-07</td>
<td>25</td>
<td>0.96</td>
</tr>
<tr>
<td>Turquoise</td>
<td>S100A9</td>
<td>S100 calcium binding protein A9</td>
<td>4.26E-07</td>
<td>25</td>
<td>0.96</td>
</tr>
<tr>
<td>Turquoise</td>
<td>IGSF6</td>
<td>immunoglobulin superfamily, member 6</td>
<td>3.20E-04</td>
<td>21</td>
<td>0.95</td>
</tr>
<tr>
<td>Turquoise</td>
<td>HCK</td>
<td>HCK proto-oncogene, Src family tyrosine kinase</td>
<td>6.80E-05</td>
<td>18</td>
<td>0.96</td>
</tr>
<tr>
<td>Turquoise</td>
<td>CD14</td>
<td>CD14 molecule</td>
<td>2.10E-05</td>
<td>17</td>
<td>0.95</td>
</tr>
<tr>
<td>Turquoise</td>
<td>TLR2</td>
<td>toll-like receptor 2</td>
<td>4.54E-08</td>
<td>17</td>
<td>0.95</td>
</tr>
<tr>
<td>Turquoise</td>
<td>VCAN</td>
<td>versican</td>
<td>6.21E-08</td>
<td>17</td>
<td>0.96</td>
</tr>
<tr>
<td>Turquoise</td>
<td>PTAFR</td>
<td>platelet-activating factor receptor</td>
<td>1.40E-05</td>
<td>16</td>
<td>0.95</td>
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<tr>
<td>Turquoise</td>
<td>GLT1D1</td>
<td>glycosyltransferase 1 domain containing 1</td>
<td>6.57E-07</td>
<td>15</td>
<td>0.95</td>
</tr>
<tr>
<td>Turquoise</td>
<td>MS4A6A</td>
<td>membrane-spanning 4-domains, subfamily A, member 6</td>
<td>4.00E-06</td>
<td>15</td>
<td>0.95</td>
</tr>
<tr>
<td>Blue</td>
<td>ABCA5</td>
<td>ATP-binding cassette, sub-family A (ABC1), member 5</td>
<td>2.24E-02</td>
<td>24</td>
<td>0.96</td>
</tr>
<tr>
<td>Blue</td>
<td>LUC7L</td>
<td>LUC7-like (S. cerevisiae)</td>
<td>1.78E-02</td>
<td>23</td>
<td>0.97</td>
</tr>
<tr>
<td>Blue</td>
<td>RBM6</td>
<td>RNA binding motif protein 6</td>
<td>4.01E-03</td>
<td>12</td>
<td>0.95</td>
</tr>
<tr>
<td>Blue</td>
<td>FAM134B</td>
<td>family with sequence similarity 134, member B</td>
<td>6.32E-03</td>
<td>11</td>
<td>0.96</td>
</tr>
<tr>
<td>Blue</td>
<td>CLUAP1</td>
<td>clusterin associated protein 1</td>
<td>8.74E-04</td>
<td>9</td>
<td>0.96</td>
</tr>
<tr>
<td>Blue</td>
<td>GABPB2</td>
<td>GA binding protein transcription factor, beta subunit 2</td>
<td>8.30E-03</td>
<td>8</td>
<td>0.94</td>
</tr>
</tbody>
</table>

genes for each of the key modules (Table 3). The top hub genes in the turquoise module included FCN1, S100A9, IGSF6, HCK, CD14, TLR2, VCAN, PTAFR, GLT1D1, and MS4A6A, and the top hub genes in the blue module included ABCA5, LUC7L, RBM6, FAM134B, CLUAP1, and GABPB2. The iSubpathwayMiner can detect the local structure of an entire pathway (subpathway) that will help us to understand the pathogenic mechanism caused by dysfunction of the subpathways. For the turquoise module, the subpathway analysis displayed that ALDH2 gene was enriched in the subpathway of glycolysis/gluconeogenesis (Figure 3). Furthermore, all highly connective genes (adjacency weight greater than 0.1) of the turquoise module were submitted to GenCLiP 2.0 for data mining. The results showed that the hub genes in the network enriched in several functional terms of biological processes, including regulation of response to wounding and regulation of phosphate metabolic process.

3.4. Replication of Module Structure and Hub Genes. We tested the module preservation of the reference data set (meta-analysis) in the validation data set (GSE123487). Among 33 identified modules, we found there was low-to-moderate evidence of preservation (Zsummary.pres = 3.5) for the turquoise module; however, other modules were not preserved (Zsummary.pres < 2). To further confirm the findings of intramodular hub genes, we performed differentially expressed mRNAs analysis in the test data set. The hub genes FCN1 (p-value = 0.002), HCK (p-value = 0.012), CD14 (p-value = 0.024), TLR2 (p-value = 0.009), VCAN (p-value = 0.009), and GLT1D1 (p-value = 0.022) in the turquoise module identified by the reference data set, as well as the hub genes ABCA5 (p-value = 0.0004), LUC7L (p-value = 0.013), and GABPB2 (p-value = 0.039) in the blue module, were also found to be differentially expressed in the test data set.
3.5. **MiRNAs Target Prediction and Validation.** TargetScan was applied to predicate the miRNAs regulated to the target DEGs in the network of key module. Only a context score percentile ≥ 90 and an aggregate PCT score ≥ 0.8 were identified as the putative miRNA of the target gene (Figures 4 and 5). The main results represented that the target genes PTAFR, BCL6, DUSP6, and KCTD12 in the turquoise module were regulated by hsa-let-7-5p, hsa-miR-124-3p, hsa-miR-145-5p, and hsa-miR-9-5p, respectively. Moreover, hsa-miR-124-3p and hsa-miR-9-5p also regulated the target genes FAM134B and SFXN2 in the blue module. The predicted results of miRNAs were tested with data set of GSE123487 using microarray assays. There were hsa-miR-9-1 (p-value = 0.026), hsa-miR-124-3 (p-value = 0.046), hsa-miR-5195 (p-value = 0.014), hsa-miR-124-3 (p-value = 0.028), hsa-miR-5195 (p-value = 0.024), hsa-let-7d (p-value = 0.037), hsa-miR-133b (p-value = 0.040), hsa-miR-526b (p-value = 0.036), and hsa-miR-5195 (p-value = 0.014) that were confirmed to be differentially expressed between AMI patients and normal controls.

4. Discussion

This study aimed to identify candidate genes and miRNAs involved in the occurrence and development of AMI. In this study, we screened out total of 1027 DEGs in blood samples from AMI patients compared with normal controls. The average linkage hierarchical clustering analysis was carried out to group coexpressed DEGs into modules, and 33 modules were identified. Among them, the turquoise and blue modules were detected as the two biggest modules to be associated with AMI. From the functional enrichment and GO analysis, we found that DEGs were mainly correlated to functional annotations of inflammatory and immune responses. The subpathway enrichment analysis revealed that ALDH2 gene in the turquoise module was enriched in glycolysis/glucogenesis subpathway. Furthermore, hsa-let-7d, hsa-let-7b, hsa-miR-124-3, and hsa-miR-9-1 were identified to regulate the key genes in the turquoise and blue modules.

Myocardial ischemia causes a reduction of oxygen supply to the heart, leading to cardiac myocytes in the hypoxia status dependent of the glycolysis metabolism, the major source of energy supply in the hypoxic circumference. In spite of the fact that cardiovascular drugs and surgical interventions can improve the oxygen supply in myocardium for increased cardiac work, these interventions were shown to be unsatisfactory to decrease cardiac events or increase patient survival [22]. Aldehydes have been reported to be highly associated with myocardial ischemia and cardiac reperfusion damage [23, 24]. Here, subpathway analysis found that ALDH2 might exert its role by influencing glycolysis/glucogenesis metabolic pathway. Mitochondrial aldehyde dehydrogenase 2 (ALDH2) is an important enzyme that catalyses the removal of reactive aldehydes, whose activation was indicated to
be correlated with reduced ischemic myocardium damage in rodent models [25]. Acetaldehyde metabolites, such as 4-hydroxynonenal (4-HNE), can affect glycolysis by modifying key glycolytic enzymes, including glyceraldehyde-3-phosphate dehydrogenase and glucose-6-phosphate dehydrogenase [26], and also they can inhibit mitochondrial respiratory chain function, promote mitochondrial membrane permeability transformation channel open, and directly lead to mitochondrial dysfunction. Therefore, ALDH2 confers profound treatment-aided value in individualized cardioprotective strategies.

The genes in turquoise module, including HMOX1, TLR2, VCAN, SI00A9, CD14, and CD36, were significantly correlated to wound healing function; in addition, HCK, HMOX1, BCL6, SI00A9, SI00A12, TLR2, and CD36 were significantly enriched in response to wounding associated biological process. It is well established that myocardial infarction is accompanied by local and systemic inflammation and leads to rapid necrosis of myocardia in the ischemic heart [27]. For postmyocardial infarction, healing the myocardium wound is essential for tissue integrity and function of the heart. There is widespread concern and increased interest in the damage associated molecular pattern molecules SI00A8 and SI00A9 in human cardiovascular disease. SI00A8 and its binding partner SI00A9 are members of the SI00 calcium-binding family of proteins, the circulating levels of which are elevated by activated inflammatory cytokines and autoimmune state [28]. The growing evidence indicated that SI00A9 played an important role in leukocyte trafficking and arachidonic acid metabolism [29, 30]. In human, several SI00 proteins, including SI00A7, SI00A8, SI00A9, and SI00A12, are linked with the severity of coronary and carotid atherosclerosis [31–34]. Thus, SI00A8 and SI00A9 might serve as a useful biomarker and therapeutic target in human cardiovascular disease; besides, SI00A8 and SI00A9 blockers have been developed and are approved for clinical testing.

Some DEGs clustered in the turquoise module have been reported to be associated with AMI, including SI00A9 [35], CD14 [36], TLR2 [37], and HMOX1 [38]. However, human studies on the association between FCN1 and AMI have not been reported, although there was a study in mice.
that MBL/ficolin-associated protein-1 (MAP-1) can attenuate myocardial injury and arterial thrombogenesis [39]. We analyzed the interactional associations of proteins encoded by DEGs in the turquoise module using the Search Tool for the Retrieval of Interacting Genes/Proteins (STRING; https://string-db.org/). The results showed that there were complicated interactions among TLR2, HMOX1, MMP9, S100A9, CD14, and FCN1. Therefore, the modules identified in the present study associated with AMI could not be considered to be independent of already established genes.

In this study, several miRNAs were predicted to regulate the key genes that contributed to the pathophysiological consequences of AMI. The let-7 is the second miRNA found in C.elegans, and it is highly expressed in the cardiovascular system [40], exerting important regulatory roles after myocardial infarction [41]. There were hsa-let-7d and hsa-let-7b confirmed to be differentially expressed between AMI patients and normal controls in the validation data set, suggesting that they might play a similar role in regulated process of AMI. Clinical data revealed that circulating miR-124 and miR-145 was significantly associated with AMI [42, 43], and also miR-145 was correlated with the severity of coronary artery [44] and played roles in regulating the evolution of atherosclerotic plaque toward instability and rupture [45]. Besides that, miR-9 was also found to be probably involved in myocardial regeneration [46].

However, some limitations should be reinforced in this study. Firstly, majority of data derives from two large studies in the USA. Therefore the results are not surely applicable to other ethnicities. Secondly, some findings need to be further studied. Despite these limitations, this study still provides some novel viewpoints in current understanding of AMI mechanism.

5. Conclusions

In summary, we screen out candidate genes for AMI, such as FCN1, CD14, S100A9, and ALDH2; in addition, we identified hsa-let-7d, hsa-let-7b, hsa-miR-124-3, and hsa-miR-9-1, as potential regulators on pathogenesis of AMI. Thus this study may offer potential therapeutic targets and new therapeutic strategies for AMI.

Abbreviations

AMI: Acute myocardial infarction
ST2: IL-1RL-1, Interleukin 1 receptor-like 1
MiRNAs: MicroRNAs
WGCNA: Weighted Gene Coexpression Network Analysis
DEGs: Differentially expressed genes
GEO: Gene Expression Omnibus
RMA: Robust Multichip Analysis
DE: Differential expression
QC: Quality control
IQC: Internal homogeneity of coexpression structure among studies

EQC: External consistency of coexpression pattern with pathway database
AQCg: Accuracy of differentially expressed gene detection
CQCg: Consistency of differentially expressed gene detection
AQCp: Accuracy of enriched pathway identification
CQCp: Consistency of enriched pathway identification
PCA: Principal component analysis
FDR: False discovery rate
GO: Gene Ontology
KEGG: Kyoto Encyclopedia of Genes and Genomes
DAVID: Database for Annotation, Visualization and Integrated Discovery
TOM: Topological overlap matrix
ME: Module eigengene
MM: Module membership
GS: Gene significance
MS: Module significance
IC: Intramodular connectivity
3’-UTR: 3’-untranslated region
Pcr: Probability of conserved targeting
SMR: Standardized mean rank
FCN1: Ficolin (collagen/fibrinogen domain containing) 1
S100A9: S100 calcium binding protein A9
IgSF6: Immunoglobulin superfamily, member 6
HCK: HCK protooncogene, Src family tyrosine kinase
CD14: CD14 molecule
TLR2: Toll like receptor 2
VCAN: Versican
PTAFR: Platelet-activating factor receptor
GLT1D1: Glycosyltransferase 1 domain containing 1
MS4A6A: Membrane-spanning 4-domains, subfamily A, member 6A
ABCA5: ATP-binding cassette, subfamily A (ABCI), member 5
LUC7L: LUC7-like (S. cerevisiae)
RBM6: RNA binding motif protein 6
FAM134B: Family with sequence similarity 134, member B
CLUAPI: Clusterin associated protein 1
GABPB2: GA binding protein transcription factor, beta subunit 2
ALDH2: Mitochondrial aldehyde dehydrogenase 2
BCL6: B-cell CLL/lymphoma 6
DUSP6: Dual specificity phosphatase 6
KCTD12: Potassium channel tetramerization domain containing 12
SFXN2: Sideroflexin 2
4-HNE: 4-hydroxynonenal
MDA: Malonaldehyde
HMOX1: Heme oxygenase (decycling) 1
S100A12: S100 calcium binding protein A12.
Data Availability


Conflicts of Interest

The authors declare that they have no conflicts of interest relevant to this work.

Authors’ Contributions

Yan Li conceived the study. Min Liu gave the analysis, provided the results, and wrote the manuscript. Xiao_nan He, Chao Li, and Ling Gong participated in the biological interpretation of the results. All authors read and approved the final manuscript.

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Supplementary Materials

Supplementary Figure 1. Picking up a soft-threshold approximating to a scale-free topology criterion based on R square and mean connectivity. Supplementary Figure 2. Number of detected DEGs under different q-value threshold. Supplementary Figure 3. Correlation coefficient and significant $p$-value between each module and AMI. (Supplementary Materials)

References


