Identification and Validation of Immune-Related Biomarker Gene and Construction of ceRNA Networks in Septic Cardiomyopathy

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Septic cardiomyopathy (SCM) is a cardiac dysfunction caused by severe sepsis, which greatly increases the risk of heart failure and death, and its molecular mechanism is unclear. The immune response has been reported to be an important process in septic cardiomyopathy and is present in the cardiac tissue of patients with sepsis, suggesting that the immune response may be an underlying mechanism of myocardial injury in SCM. Therefore, we explored the role of immune-related genes (IRGs) in SCM and aimed to identify pivotal immune-related targets with the aim of identifying key immune-related targets in SCM and potential therapeutic mechanisms involved in the pathological process of SCM. To explore the regulatory mechanisms of immune responses in SCM, we identified differentially expressed genes (DEGs) shared in the SCM datasets GSE179554 and GSE40180 by bioinformatics analysis and then obtained hub genes from the DEGs. Then, we obtained the immune-related hub genes (IRHGs) by intersecting the hub genes with IRGs and performed quantitative reverse transcription polymerase chain reaction to confirm the abnormal expression of IRHGs. Finally, we further constructed an immune-related lncRNA–miRNA–IRHG ceRNA regulatory network. In this study, we identified an IRHG that may be involved in the pathogenesis of SCM, which helps us to further elucidate the role of immune response in SCM and gain insights into the molecular mechanisms and potential therapeutic targets of SCM.

Keywords: septic cardiomyopathy, non-coding RNA, immune-related gene, biomarker, ceRNA network

INTRODUCTION

Septic cardiomyopathy (SCM) is a systemic, multi-organ–linked malignant disease induced by sepsis, which often leads to host immune system disorders, organ dysfunction, and even death (Beesley et al., 2018). Not only that, but when patients with sepsis develop myocardial dysfunction, the mortality rate increases from 10% to 40% (Singer et al., 2016). At present, the diagnosis of SCM is still mainly based on echocardiography (Annane et al., 2005) and myocardial injury markers (Charpentier et al., 2004; Romero-Bermejo et al., 2011), and early intervention for SCM cannot be performed. Therefore, the exploration of
novel biomarkers for early diagnosis and intervention of SCM may be crucial to reducing the mortality of patients with SCM.

Non-coding RNAs (ncRNAs) [including long ncRNAs (lncRNAs) and microRNAs (miRNAs)] are transcripts with a variety of biological regulatory functions and are key regulators of many biological processes (BPs) such as proliferation, differentiation, apoptosis, and metabolism. Furthermore, competing endogenous RNA (ceRNA) networks, composed of lncRNAs that share miRNA response elements (MREs) with miRNAs, play regulatory roles in a variety of diseases, including sepsis (An et al., 2021) and SCM (Pei et al., 2021). Pei et al. revealed that the lncRNA TTN-AS1/miR-29a/E2F2 ceRNA network could successfully reduce SCM-induced myocardial injury (Pei et al., 2021). Therefore, the ceRNA network established by lncRNAs and miRNAs is likely to be a key regulatory mechanism in the progression of SCM.

The immune response is the body’s defense against external damage. In septic cardiomyopathy caused by sepsis, immune response is an important pathological process. Current research on immune responses in SCM has focused on the role of pro-inflammatory mediators at the molecular and cellular levels (Hoesel et al., 2007). For example, Zhang et al. found that inhibition of TLR4/NF-κB/TNF-a/IL-18 immune signaling in cardiomyocytes could prevent LPS-induced cardiac dysfunction in mice (Zhang et al., 2020). It is suggested that immune regulation may be an important part of myocardial injury in SCM. However, the exact mechanism of immune response in sepsis-induced myocardial injury remains unclear.

Here, to determine the role of immune responses in SCM, we obtained differentially expressed genes (DEGs) shared in datasets GSE179554 and GSE40180 by bioinformatics analysis. Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analysis of shared DEGs were performed subsequently. On the basis of the STRING online database, we established a protein–protein interaction (PPI) network of shared DEGs and used the cytoHubba plugin to screen out hub genes from the PPI network. We then analyzed shared DEGs and immune-related genes (IRGs) to obtain IRGs in SCM. The IRHG in SCM were finally obtained after the intersection of Hub gene and IRGs and verified by quantitative reverse transcription polymerase chain reaction (qRT-PCR) and dataset to confirm the abnormal expression of IRHG. Not only that, after screening the target IRHG, we obtained the upstream miRNAs and lncRNAs of IRHG from the relevant database. Then, key immune-related lncRNA–miRNA–IRHG ceRNA networks were constructed after dataset validation. We summarize the overall flow chart of this study and present it in Figure 1.

MATERIALS AND METHODS

Expression Profile of mRNA and IncRNA in Microarray Data From Mice and Rat With Septic Cardiomyopathy

Five septic cardiomyopathy mouse, rat, and human datasets (GSE179554, GSE40180, GSE171546, GSE159309 and GSE12624) were obtained from the National Institutes of Health (NIH)—National Center for Biotechnology Information (NCBI)–Gene Expression Omnibus (GEO) database (https://www.ncbi.nlm.nih.gov/gds/). Details of these GEO datasets are presented in Table 1. Myocardial tissues from septic cardiomyopathy mice were used in all five databases, and expression profiles were analyzed by high-throughput sequencing (GSE179554 and GSE171546) and array sequencing (GSE40180, GSE159309, and GSE12624), respectively. Therefore, we used these five datasets for the acquisition of SCM immune-related biomarker genes and the construction of ceRNA networks. It is important to note that our technique is achievable and accepted, and gene expression under pathophysiological conditions has been studied by scholars (Xu et al., 2020; Cheng et al., 2021).

Functional Enrichment Analyses

The biological functions of recognized DE-mRNAs of pastime had been assessed through the usage of the Metascape bioinformatics sources (Metascape, a gene annotation and analysis resource, http://metascape.org, version 3.5) (Zhou et al., 2019). DE-mRNAs had been first screened from the dataset, and then, the shared DE-mRNAs had been imported into Metascape for GO and KEGG enrichment analysis. For GO analyses, enriched BPs, molecular functions (MFs), and cellular components (CCs) have been assessed.

Protein–Protein Interaction Analysis

PPI networks have been generated from the STRING database (https://string-db.org/) (Szklarczyk et al., 2021). A rating of 0.4 (medium confidence) used to be set as a threshold. Next, to obtain permission from the central gene, we imported the PPI evaluation consequences into the Cytoscape software program software (version 3.4.0, https://cytoscape.org/) (Shannon et al., 2003), using the cytoHubba plugin (Chin et al., 2014) perceived vital genes in this neighborhood as hub genes. We used five algorithms, specifically Degree, Maximal Clique Centrality (MCC), Maximum Neighborhood Component (MNC), Density of MNC (DMNC), and Clustering Coefficient, to calculate the top 10 hub genes.

Identification of Immune-Related Hub Genes

To also elucidate whether the shared DEGs have immune-related functions, we obtained 782 immune cell marker genes from a previously published article (Bindea et al., 2013) and intersected the shared DEGs with immune cell marker genes to obtain IRGs in SCMs. DEGs that overlap with immune cell-specific marker genes are defined as immune-associated DEGs. Then, we crossed the top 10 hub genes with IRGs to obtain immune-related hub genes (IRHGs) in SCM. For detailed research methods, refer to the published articles (Yu et al., 2021).
Emergency Medicine of the First Affiliated Hospital of Kunming Medical University, Yunnan Province, and medical information was collected in accordance with the Declaration of Helsinki after informed consent of the patients. This study was approved by the Ethics Committee of the First Affiliated Hospital of Kunming Medical University, Yunnan Province (2022) ethical review L, No.23). Patients with a clear history of sepsis and reduced left ventricular ejection fraction (LVEF) <0.55 were included.
in the study. Total RNA was isolated using TRIzol reagent (Invitrogen) followed by amplification grade (Invitrogen) DNase 1 for further processing. TaqMan Reverse Transcription Reagents (Applied Biosystems) were used for cDNA synthesis. qRT-PCR was performed in a My qQ Thermal Cycler (Bio-Rad) using 2× qQ SYBR Green Supermix (Bio-Rad). Each reaction contains 1 μl of cDNA for a total volume of 20 μl. Using glyceraldehyde-3-phosphate dehydrogenase (GAPDH) as the internal control of the target gene, the $2^{-ΔΔCt}$ method was used for data analysis. PCR primers were designed for human sequences and synthesized by Invitrogen.

**Construction of ceRNA Networks**

RNA22 (https://cm.jefferson.edu/rna22/Interactive/) (Loher and Rigoutsos, 2012) was used to predict IncRNAs that interacted with the selected miRNAs. The predicted IncRNAs of were intersected with the GSE159309-IncRNA database to obtain the interaction IncRNA. We selected the IncRNA with the largest fold change and entered NCBI as the target-IncRNA. Then, the ceRNA network based on the interaction between mRNA, miRNA, and IncRNA was constructed using Cytoscape.

**Statistical Analysis of Microarray Data**

The two-tailed Students’ t-test was once used to pick out the variations between groups. We used the R statistical evaluation bundle (version 3.5.3) to statistically analyze the data. An alpha price of $P < 0.05$ was once measured and regarded as a statistically tremendous match or evaluation in this study.

**RESULT**

**Identification of DE-mRNAs in Myocardial Tissue of SCM Mice and Rat**

Our research covers GEO datasets GSE179554 and GSE40180. GSE179554 has a total of 104 samples, from which we selected four normal myocardial tissue data as the control group and four cecal ligation and puncture (CLP) mouse model myocardial tissue data as the test group for analysis. GSE40180 has a total of 72 samples, from which we extracted five normal control myocardial tissue data and five CLP mouse model myocardial tissue data for subsequent analysis. The cluster heatmap and volcano plot in Figures 2A, B, 3A, B show the data distribution of the GSE179554 and GSE40180 datasets, respectively. In the GSE179554 dataset, a total of 2,198 differentially expressed mRNAs (DEGs, $P < 0.05$) were identified (1,665 upregulated and 533 downregulated). In the GSE40180 dataset, a total of 5,720 differentially expressed mRNAs (DEGs, $P < 0.05$) were identified (2,578 upregulated and 3,142 downregulated). Then, we crossed the differential genes of GSE179554 and GSE40180 to obtain 453 shared DEGs and used these differential genes for subsequent analysis.

**Pathway Enrichment Analyses**

The consequences of GO analysis indicated that these shared DEGs were associated with diverse BPs, MFs, and CCs. The top three BPs those shared DEGs were involved in were proteasomal protein catabolic process, proteolysis involved in cellular protein catabolic process, and ubiquitin-dependent protein catabolic process; the top three significant MFs related to those shared DEGs were interleukin-1 binding, ubiquitin ligase-substrate adaptor activity, and growth factor binding; the top three significant C Cs in which these shared DEGs participated were proteosome complex, endopeptidase complex, and peptidase complex (Figure 4A). KEGG pathway analysis showed those DE-mRNAs to be significantly enriched in proteasome, spinocerebellar ataxia, and Epstein–Barr virus infection pathway (Figure 4B).

**PPI Network Analysis and CytoHubba Gene Identification**

The interaction network between proteins coded by shared DEGs, which was composed of 417 nodes and 922 edges, was constructed and visualized by STRING (Figure 5A). Next, we used the cytoHubba plugin to identify hub genes. The top 10 hub
genes (Figure 5B) were identified by intersecting the results from the five algorithms of cytoHubba including Degree, MCC, MNC, DMNC, and Clustering Coefficient. Finally, we got the top 10 hub genes: PSMC4, PSMD7, PSMA5, PSMA7, PSMB1, PSMD4, PSMD11, PSMB4, PSMD9, and NFKBIA. Details of these hub genes are shown in Table 2. These genes are the most important in the PPI network and may play an important role in the pathogenesis of SCM.

Identification of Immune-Related Differentially Expressed Genes in SCM

We intersected 453 shared DEGs with 782 IRGs and obtained 10 immune genes present in SCMs: STX16, CEP68, SMAD1, ZBTB16, PSMD4, HSD11B1, PPARG, SLC7A8, THBS1, and ASPSCR1 (Figure 6A). Details of the 10 IRGs are proven in Table 3. This suggests that these IRGs play crucial roles in the immune response of SCM. Because SCM is an acute aggravating ailment induced via circulatory dysfunction and immune gadget imbalance brought on by way of infection, the steady-state rules and protection feature of the immune machine significantly have an effect on the prognosis of SCM. To locate the key targets of SCM, we intersected 10 immune genes and 10 hub genes of cytoHubba. Finally, we obtained an IRHG-PSMD4, which was upregulated in SCM (FC = 1.43, \( P = 0.00089 \) in GSE179554; FC = 2.61, \( P = 0.00565 \) in GSE40180) (Figures 6B, C). To additionally verify whether or not PSMD4 is a normal gene in SCM ailment states, we validated PSMD4 with the GSE171546 and GSE12624 datasets.
The GSE171546 dataset has a total of 20 samples, and we used five normal myocardial tissue data and five CLP-induced sepsis mouse model myocardial tissue data to verify the expression of PSMD4. GSE12624 collected 70 human samples, including 36 control samples and 34 SCM samples. The results showed that PSMD4 had a trend consistent with previous results, being upregulated and statistically significant in both the mouse dataset GSE171546 (FC = 1.98, \( P = 0.00011 \)) (Figure 6D) and the human dataset GSE12624 (FC = 1.65, \( P = 0.00964 \)) (Figure 6E).

Validation of Immune-Related Hub Genes

We assessed the expression of the IRHG-PSMD4. qRT-PCR was used to detect the gene expression of PBMCs in three patients with SCM and three normal people. The results of qRT-PCR had a consistent trend with the analysis results: as shown in Figure 6F, the expression level of PSMD4 in PBMCs of patients with SCM was significantly higher than that of normal controls (\( P < 0.05 \)).

Prediction of Target miRNAs and Construction of the Co-Expressed Network

We used three online miRNA databases: starBase, miRWalk, and DIANA-TarBase, to predict the target miRNAs of PSMD4. The miRNA prediction results of PSMD4 are shown in Supplementary Table S1. The miRNA prediction results showed that miR-324-5p coexisted in the three databases (Figure 7A), suggesting that miR-324-5p is an effective miRNA of PSMD4. Thus, the co-expressed network of PSMD4 and target miRNAs was constructed: miR-324-5p/PSMD4.
Construction of Immune-Related IncRNA–miRNA–IRHG Regulatory Network
miRNAs are well known to induce gene silencing and downregulate gene expression by binding mRNAs. However, its upstream molecules, IncRNAs, can affect the function of miRNA by combining MREs, thus upregulating gene expression. This interaction between RNAs is called a ceRNA network (Salmena et al., 2011). Next, we used the online database RNA22 to predict IncRNAs that interact with miR-324-5p. The prediction results of IncRNAs are shown in Supplementary Table S2. Through database prediction, we obtained 2,955 IncRNAs with potential binding sites for miR-324-5p. We then validated the IncRNAs using dataset GSE159309, which includes five lipopolysaccharide (LPS)–induced sepsis rat myocardial tissue data and five normal myocardial tissue data. Finally, after validation on the GSE159309 dataset, we obtained two IncRNAs for miR-324-5p from 2,955 predicted IncRNAs. One IncRNA (Kantr, FC = 0.19, P = 0.0015) (Figure 7B) was downregulated in SCM and one IncRNA (Thumpd3-as1, FC = 5.87, P = 0.0253) (Figure 7C) was upregulated in SCM. On the basis of the ceRNA hypothesis, we selected IncRNAs upregulated in SCM from these validated IncRNAs for ceRNA network construction. The binding sites of miR-324-5p and PSMD4 are shown in Figure 7D. Finally, an immune-related Thumpd3-as1/miR-324-5p/PSMD4 ceRNA network involved in the regulation of SCM mechanisms was constructed (Figure 7E), and the workflow of our study is summarized in Figure 8.

DISCUSSION
In the 1980s, Parker et al. evaluated cardiac characteristics in patients with septic shock and found that patients with septic cardiomyopathy had reduced LVEF and organic left ventricular enlargement (Parker et al., 1984). The high mortality rate of septic cardiomyopathy is due to the fact that it is a multi-organ–related infectious disease that is harmful to the whole body, resulting in dysfunction of the immune system and multiple organs (Hollenberg and Singer, 2021). At the same time, the activation of the body’s inflammatory response further impairs cardiomyocyte function (Brown and Jones, 2004). Therefore, the search for specific and highly sensitive diagnostic biomarkers is crucial for the early intervention of SCM. In recent years, target gene therapy has achieved remarkable results in the early diagnosis, treatment, and prognosis prediction of the disease (Dmitrieva et al., 2020; Sharma et al., 2020), enlightening that we can explore the early diagnosis and treatment biomarkers of SCM at the gene level.

Immune response is an important pathological process of SCM and participates in the regulation of SCM process (Zhang et al., 2020). However, the roles of IRGs in the development of SCM are currently unclear. Therefore, we combined shared DEG profiles in septic cardiomyopathy with IRGs to obtain IRHGs and immune-related ceRNA regulatory networks in SCM to deepen our understanding of the molecular mechanisms of septic cardiomyopathy.
After the construction of the PPI network, we identified 10 differentially expressed hub mRNA (PSMC4, PSMD7, PSMA5, PSMA7, PSMB1, PSMD4, PSMD11, PSMB4, PSMD9, and NFKBIA) using cytoHubba. These hub genes may be potential targets for the regulation in SCM. After verification of IRGs, PSMD4 was finally identified as IRHGs. PSMD4 (also known as Rpn10, official full name: proteasome 26S subunit ubiquitin receptor, non-ATPase 4) is a subunit of regulatory granules, is the main ubiquitin (Ub) receptor for the 26S proteasome. Our results showed that PSMD4 is not only a central gene but also an IRG in SCM. At present, there are several studies showed that PSMD4 could affect the development of most tumor disease. For example, the study of Jiang et al. showed that PSMD4 can regulate the level of hypoxia-inducible factor 1α through the PTEN/Akt signaling pathway, thereby promoting the occurrence of hepatocellular carcinoma (Jiang et al., 2019); PSMD4 can inhibit endoplasmic reticulum (ER) stress-induced cells apoptosis, advertising the development of esophageal cancer (Ma et al., 2019); in addition, Xu et al. (2021) showed that PSMD4 is a potential target in mouse colorectal cancer, mediating the enhancement of immunoproteasome activity by atractylenolide I (ATT-I), making the body more responsive to immune checkpoint blocking therapy. However, studies on PSMD4 in septic cardiomyopathy are still insufficient. Therefore, we believe that PSMD4 holds promise as a novel immune-related biomarker and therapeutic target in SCM.

In addition, although immune responses are known to be an important pathological process in SCM, it is uncommon to construct immune-related ceRNA networks in SCM. Here, we constructed a ceRNA network targeting PSMD4 to explore the potential regulation of PSMD4 in SCM immunity. In this study, Thumpd3-as1/miR-324-5p/PSMD4 signaling pathway was identified by bioinformatic analysis.

**Thumpd3-as1/miR-324-5p/PSMD4 Axis**

After confirming that PSMD4 is the hub gene of SCM, we estimated its target miRNA and finally got miR-324-5p. Because the main way that miRNA regulates target genes is to inhibit the expression of mRNA or promote the degradation of mRNA (Correia de Sousa et al., 2019), that is, miRNA is negatively correlated with mRNA expression. Our results showed that the IRG-PSMD4 was upregulated in SCM. Therefore, PSMD4 upstream miRNAs should be at low levels. Wang et al. found that the level of miR-324-5p in hypoxic/reoxygenated cardiomyocytes decreased with prolonged hypoxia, and miR-324-5p altered mitochondrial morphology and cardiomyocyte death by acting on Mtf1 (Wang et al., 2015);
the results in the work of Chen et al. confirmed that the expression level of miR-324-5p in peripheral blood endothelial progenitor cells of patients with ST-segment elevation myocardial infarction (STEMI) was significantly lower than that of healthy volunteers (Chen et al., 2019). In addition, Helwak et al. confirmed that miR-324-5p and PSMD4 have direct binding sites by CLASH technique (Helwak et al., 2013). These studies all demonstrate that it is realistic to construct a miR-324-5p/PSMD4 regulatory pathway. However, the specific mechanism of miR-324-5p/PSMD4 in SCM has not been reported yet, and further studies are needed.

Then, we further explored the lncRNAs that bind to miR-324-5p. We obtained two consensus lncRNAs (Kantr, FC = 0.19, \( P = 0.0015 \); and Thumpd3-as1, FC = 5.87, \( P = 0.0253 \)) by RNA22 database prediction and GSE159309 dataset validation. Finally, Thumpd3-as1, a lncRNA with high expression in SCM, can bind to miR-324-5p according to the ceRNA network hypothesis (Smillie et al., 2018).

Previous research on Thumpd3-as1 (THUMPD3 antisense RNA 1) is mainly in oncology and osteoarthritis. Hu et al. confirmed that Thumpd3-as1 can regulate the self-renewal of non–small cell lung cancer cells (Hu et al., 2019); the findings of Wang et al. confirmed that Thumpd3-as1 overexpression enhanced the inflammatory response of chondrocytes in osteoarthritis (Wang et al., 2021); the study of Xia et al. confirmed that Thumpd3-as1 is a lncRNA related to the immune microenvironment of colon adenocarcinoma (Xia et al., 2021). All of these suggest that Thumpd3-as1 has strong feasibility in cellular characterization and immune regulation. However, the specific effects of Thumpd3-as1 and Thumpd3-as1/miR-324-5p/PSMD4 ceRNA network in the mechanism of SCM still need to be further explored.

As we pointed out in Table 4, two previous papers reported that identification of hub genes in human septic cardiomyopathy using bioinformatic analysis (Chen et al., 2020; Kang et al., 2020). Here, we presented that several findings in our study are significantly novel in comparing to what these two published papers. First of all, we used two datasets for DEGs acquisition and dataset validation for targets, which was solid. Furthermore, we identified IRHGs rather than common hub gene in SCM; in addition, a novel immune-related ceRNA network, Thumpd3-as1/miR-324-5p/PSMD4, was constructed and validated, which was different from the previous papers.

Above all, this study explored biomarker genes in SCM using multiple GEO datasets of septic cardiomyopathy mouse heart tissue. In addition, we innovatively combined the hub gene of septic cardiomyopathy with the immune gene, obtained the important immune hub gene PSMD4 in SCM, and constructed the immune-related Thumpd3-as1/miR-324-5p/PSMD4 ceRNA network. It is suggested that PSMD4 and Thumpd3-as1/miR-324-5p/PSMD4 ceRNA network may be important biomarkers and potential therapeutic targets in SCM.

However, there are still some limitations of this study that need to be addressed. First, in our study, datasets of human and animal SCM model samples were used for analysis, but the sample size of peripheral blood from patients with SCM used for evaluation and validation in this study was small. Further increasing the sample size and conducting a potential cohort study to demonstrate our findings is what we need to do in the future. Second, although human samples were used for
FIGURE 8 | The technical workflow of this article and we proposed the novel ceRNA network. The red word means upregulation, and the green word means downregulation. IRHG3s, immune-related hub genes.
validation in this study, we used only peripheral blood single nucleated cells from patients with SCM for analysis due to the difficulty in obtaining live myocardial tissue. Obtaining cardiac tissues from patients with SCM and performing gene expression analysis under possible future conditions would improve the credibility and authenticity of the results of this study. Last, the association of immune genes with disease and disease severity requires further estimation due to lack of data including disease phenotype and disease activity status.

### DATA AVAILABILITY STATEMENT

Publicly available datasets were analyzed in this study. This data can be found at https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE179554, GSE40180, GSE171546, SE159309 and GSE12624.

### AUTHOR CONTRIBUTIONS

All authors have made significant contributions to the design and conception of the study, as well as data acquisition, analysis, and interpretation, and participated in the drafting or revision of the manuscript. For the final version to be published, all authors have made the final approval, and have reached an agreement with the journal to which the article is submitted, and are willing to be responsible for all aspects of the work to ensure the accuracy or completeness of any part of the article. All the authors are willing to cooperate in the investigation and resolution of related issues in the follow-up of the article. All authors contributed to the article and approved the submitted version.

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### SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcimb.2022.912492/full#supplementary-material

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### TABLE 4 | Several findings in this study are significantly novel in comparing to what is published.

| Items | Identification of Hub Genes in SCM Based on Bioinfomatics Analysis | Cell Paper (PMID:31424269) | Cell Paper (PMID:31794266) |
|-------|---------------------------------------------------------------|-----------------------------|-----------------------------|
| Years | 2022                                                          | 2019                        | 2020                        |
| Test set | GSE401807 and GSE179554                                      | GSE79962                    | GSE79962                    |
| Species/tissue | Mice/Heart                                      | Human/Heart                | Human/Heart                |
| Hub genes | IRHG: PSMD4                           | MYC, SERPINE1, CCL2                 | NDUF5B, TIMMDC1, VDAC3             |
| Validation set | GSE171546 and GSE159309                      | –                          | –                          |
| Verification | Human PBMC: qRT-PCR                              | –                          | –                          |
| Mechanism | IncRNA–miRNA–IRHG ceRNA network:                | TF-miRNA-mRNA network      | –                          |
| WB, Western blot. |                                                                                      |                            |                            |
