Myocardial fibrosis disrupts normal myocardial architecture, interferes with normal electrophysiological activity, and can also interfere with the rhythmic activity of the myocardium at the corresponding level; this aggravates the pathologic state of heart diseases and causes new pathological changes [1]. Myocardial fibrosis has been shown to significantly increase the mortality of fatal diseases, including atrial fibrillation, sudden death, and heart failure [2]. Cardiac fibroblasts are important factors in the progression of myocardial fibrosis [3].

Pyroptosis is primarily dependent on caspase-1 activation by inflammasomes and specific pattern recognition receptors on the surface or inside of cells that bind to specific ligands, and successively bind to other proteins to form multimeric protein complexes inflammasomes (pyroptosomes), including NOD-like receptor family protein 3 (NLRP3) inflammatory vesicles, and so on [4]. Evidence shows that NLRP3-mediated pyroptosis contributes to myocardial fibrosis [5]. However, the role of pyroptosis signalling in cardiac fibrosis regulation remains unknown, and more research is required to shed light on this.

LncRNAs are long noncoding RNAs that are longer than 200nt (nucleotides). LncRNAs exert critical roles in chromatin remodelling, transcriptional activation, transcriptional interference, and intranuclear trafficking [6]. LncRNAs also contribute to the initiation and progression of pyroptosis [7]. The antisense non-coding RNA (ANRIL) in the INK4 gene is a lncRNA with 19 exons [8]. ANRIL spans the genome at 126,300 base pairs and is transcribed as a 3834-nt lncRNA [9). ANRIL is found at 9p21 in the antisense direction within the p15-p16-p14 gene cluster [10]. Experiments demonstrated that LncRNAs abnormalities can cause various diseases, particularly fibrotic diseases [11]. The present experiment revealed that LncRNA-ANRIL was down-regulated in myocardial fibrotic tissue and pyroptotic cardiac fibroblasts. However, the molecular mechanisms underlying the effect of LncRNA-ANRIL on myocardial fibrosis remain largely unknown.

1. Introduction

Myocardial fibrosis disrupts normal myocardial architecture, interferes with normal electrophysiological activity, and can also interfere with the rhythmic activity of the myocardium at the corresponding level; this aggravates the pathologic state of heart diseases and causes new pathological changes [1]. Myocardial fibrosis has been shown to significantly increase the mortality of fatal diseases, including atrial fibrillation, sudden death, and heart failure [2]. Cardiac fibroblasts are important factors in the progression of myocardial fibrosis [3].

Pyroptosis is primarily dependent on caspase-1 activation by inflammasomes and specific pattern recognition receptors on the surface or inside of cells that bind to specific ligands, and successively bind to other proteins to form multimeric protein complexes inflammasomes (pyroptosomes), including NOD-like receptor family protein 3 (NLRP3) inflammatory vesicles, and so on [4]. Evidence shows that NLRP3-mediated pyroptosis contributes to myocardial fibrosis [5]. However, the present investigation aims to reveal how DNMT1/lncRNA-ANRIL/NLRP3 influences fibrosis and cardiac fibroblast pyroptosis. Here, we used ISO to induce myocardial fibrosis in mice, and LPS and ATP to induce myocardial fibroblast pyroptosis. The results showed that DNMT1, Caspase-1, and NLRP3 expression were significantly increased in fibrotic murine myocardium and pyroptotic cardiac fibroblasts, whereas LncRNA-ANRIL expression was decreased. DNMT1 overexpression decreased the level of LncRNA-ANRIL while increasing the levels of NLRP3 and Caspase-1. Contrarily, silencing DNMT1 increased the LncRNA-ANRIL and decreased the levels of NLRP3 and Caspase-1. Silencing LncRNA-ANRIL increased the levels of NLRP3 and Caspase-1. The present findings suggest that DNMT1 can methylate LncRNA-ANRIL during the development of myocardial fibrosis and CFs cell scorching, resulting in low LncRNA-ANRIL expression, thereby influencing myocardial fibrosis and cardiac fibroblast pyroptosis.
Expression of LncRNAs is regulated by DNA methylation. The methylation of LncRNA-ANRIL promoter is a biomarker for predicting cardiovascular disease risk [12]. DNA methyltransferases catalyze and maintain CpG island hypermethylation in the gene promoter region. DNA methyltransferases 1 (DNMT1) is a DNA methylation transferase that acts on a double DNA strand with one methylated strand to completely methylate it and can participate in the methylation of new synthetic strands in the double strand of DNA replication [13]. Mounting evidence shows that LncRNA binding to DNMT1 potentially influences the methylation of genes involved in myocardial fibrosis [14, 15]. However, whether LncRNA-ANRIL can mediate DNMT1 and affect myocardial fibrosis and the associated mechanisms are elusive.

Therefore, the present study intends to validate whether DNMT1 can modify LncRNA-ANRIL via methylation during the development of myocardial fibrosis and CFs cell scorching, resulting in LncRNA-ANRIL low expression and successively influencing myocardial fibrosis and cardiac fibroblasts pyroptosis.

2. Materials and Methods

2.1. Animal model

Male C57BL/6 SPF grade mice weighing 20±2 g were obtained from Shanghai Slack Laboratory Animals Co., Ltd. (experimental animal production license number: SCXK Hu 2015-0002) and housed at the Experimental Animal Center of the Jiangsu Provincial Academy of Chinese Medicine. The 40 mice were divided randomly into two groups of 20 each, one for control and one for experimentation [16]. The model group (N=20) received daily subcutaneous injections of isoproterenol (ISO; National Medicine Standard: H50020020, purchased from Nanjing, Jiangsu) (5 mg/kg body weight), whereas the control group (N=20) received an equal volume of saline solution. Mice were sacrificed after 14 days, and their hearts were removed. Every heart was divided into two parts, one of which was stored in liquid nitrogen, and the other was fixed in 10% formalin.

2.2. HE staining

A sufficient amount of the fixed sample was placed in a 4% paraformaldehyde solution in the embedding box and then placed in an automatic dehydrator for dehydration. Following complete dehydration, paraffin embedding was performed, and the paraffinized tissue sections were obtained using a manual rotary pathological microtome. The paraffinized tissue slices were placed on an automatic slicer for baking and, 1 hour later, the slices were considered ready. Dewaxed sections were soaked in citrate buffer for 10 minutes before being heated to 100 °C for 10 minutes and blocked in 5% Bovine albumin (Bovine serum albumin solution (BSA) for 1 hour, followed by overnight incubation with primary antibodies at 4°C. Then, the incubated primary antibody was recovered, and the slides were washed with phosphate-buffered saline (PBS) (5 min/cycle, for 3 cycles). After that, the slides were then incubated for 1 hour with the secondary antibody at room temperature. Then, the incubated secondary antibody was recovered, and the slides were washed with PBS (5 min/cycle, for 3 cycles), and successively stained with DAB for 1 min, and re-dyed with hematoxylin. Finally, the sections were dehydrated in ethanol solutions of different concentrations and sealed with neutral resin. Sections of stained specimens were examined at high magnification using a microscope. Five fields were assigned randomly to each segment. ImagePro Plus 6.0 software (USA) was used to quantify the percentage of positive area. All antibody brands and dilution concentrations were as follows: Caspase-1 (83383, CST, 1: 500), DNMT1 (ab188453, Abcam, 1: 500), NLRP3 (15101, CST, 1: 300), Type I collagen (ab260043, Abcam, 1: 500) and Type III collagen (22734-1-AP, Proteintech, 1: 200).

2.3. Masson staining

A sufficient amount of the fixed sample was placed in a 4% paraformaldehyde solution in the embedding box and then placed in an automatic dehydrator for dehydration. Following complete dehydration, paraffin embedding was performed, and the paraffinized tissue sections were obtained using a manual rotary pathological microtome. The paraffinized tissue slices were placed on an automatic slicer for baking and, 1 hour later, the slices were considered ready; slices were stained using the Masson staining kit (BP-DL021, SenBeiJia Biological Technology Co., Ltd., Nanjing, China). Finally, the slices were automatically machine-mounted, placed in a fume hood to air dry, taken out the next day, placed in a slicing box, stored at room temperature, and observed and photographed under an optical microscope. We checked at least five random areas of each section and used an automated photo and image analysis system (Image-Pro Plus 6.0, USA) for semi-quantitative evaluation.

2.4. Sirius red staining

A sufficient amount of the fixed sample was placed in a 4% paraformaldehyde solution in the embedding box and then placed in an automatic dehydrator for dehydration. Following complete dehydration, paraffin embedding was performed, and the paraffinized tissue sections were obtained using a manual rotary pathological microtome. The paraffinized tissue slices were placed on an automatic slicer for baking and, 1 hour later, the slices were considered ready; slices were stained using the Sirius red staining kit (BP-DL030, SenBeiJia Biological Technology Co., Ltd., Nanjing, China). Finally, the slices were automatically machine-mounted, placed in a fume hood to air dry, taken out the next day, placed in a slicing box, stored at room temperature, and observed and photographed under an optical microscope. We checked at least five random areas of each section and used an automated photo and image analysis system (Image-Pro Plus 6.0, USA) for semi-quantitative evaluation.

2.5. Immunohistochemistry (IHC)

The paraffinized tissue slices were placed on an automatic slicer for baking and, 1 hour later, the slices were considered ready. Dewaxed sections were soaked in citrate buffer for 10 minutes before being heated to 100 °C for 10 minutes and blocked in 5% Bovine albumin (Bovine serum albumin solution (BSA) for 1 hour, followed by overnight incubation with primary antibodies at 4°C. Then, the incubated primary antibody was recovered, and the slides were washed with phosphate-buffered saline (PBS) (5 min/cycle, for 3 cycles). After that, the slides were then incubated for 1 hour with the secondary antibody at room temperature. Then, the incubated secondary antibody was recovered, and the slides were washed with PBS (5 min/cycle, for 3 cycles), and successively stained with DAB for 1 min, and re-dyed with hematoxylin. Finally, the sections were dehydrated in ethanol solutions of different concentrations and sealed with neutral resin. Sections of stained specimens were examined at high magnification using a microscope. Five fields were assigned randomly to each segment. ImagePro Plus 6.0 software (USA) was used to quantify the percentage of positive area. All antibody brands and dilution concentrations were as follows: Caspase-1 (83383, CST, 1: 500), DNMT1 (ab188453, Abcam, 1: 500), NLRP3 (15101, CST, 1: 300), Type I collagen (ab260043, Abcam, 1: 500) and Type III collagen (22734-1-AP, Proteintech, 1: 200).

2.6. Cell culture and treatment

One-three day old neonatal C57BL/6J mice were killed and disinfected with 75% alcohol for 10 minutes. The heart was retrieved by opening the chest and cleaned thrice with PBS. Extraneous tissue and vasculature were removed. Mouse heart tissue was minced with scissors. Next, we added 0.25
percent trypsin and 0.1 percent type II collagenase, and the mixture was shaken for 30 minutes in a 37°C thermostat. The suspension was filtered through a 200-mesh filter after digestion. Next, the filtrate was collected and centrifuged. The cells were centrifuged for 5 min at 1000rpm and supernatant was discarded. The cells were grown in a DMEM medium containing 10% foetal bovine serum at 37°C and 5% CO₂. After two hours of culture, the cells were removed from the medium and replaced with new ones. Cells were divided into control and model groups. After 5.5 hours, 1 μg/ml LPS was added to the model group cells, followed by 5 ng/ml ATP. Cells in the model group were incubated in the same amount of PBS for the same amount of time.

2.7. Plasmid transfection and siRNA transfection

siRNA targeting ANRIL and DNMT1 mRNA, as well as plasmids overexpressing ANRIL and DNMT1, were designed and constructed by Shanghai Jikai Gene Technology Co., Ltd. The cells were digested before they were transferred to six-well culture dishes for incubation. Cells were transfected with RNAiMax Lipofectamine (Invitrogen, Carlsbad, CA, USA) at various dosages of siRNA or promiscuous ribonucleic acids as directed by the manufacturer.

2.8. qRT-PCR analysis

RNAiso Plus (9190, TAKARA, Shiga, Japan) was used to lyse the ground mouse heart tissue, which was mixed with chloroform and centrifuged at 12000 rpm for 15 min. After extraction, the supernatant was mixed with isopropanol and centrifuged at 12,000 rpm for 10 min. The supernatant was discarded, and 75% alcohol was used for washing the precipitate. After washing, the supernatant was discarded again and the extracted RNA was dissolved in diethyl pyrocarbonate water. PrimeScript™ RT reagent Kit with gDNA Eraser (RR047A, TAKARA, Shiga, Japan) was used to reverse-transcribe mRNA into cDNA. Finally, the TB Green Premix Ex TaqII kit (RR420A, TAKARA, Shiga, Japan) was used to analyze the gray value of the protein band, and GAPDH was used to measure the gray value of the protein. The sequencing of primers was as follows: β-actin (Forward, 5'-CTACCTCATGAAAGATCCTGACC-3'; Reverse, 5'-CACACTCTTCTCTTGTATGTCAC-3'), ANRIL (Forward, 5'-TAGCCTCAGAGATTATCCTGTT-3'; Reverse, 5'-GCAGAAGTCACCAAGCCTACC-3'), COL1A1 (Forward, 5'-TGAACGTTGAGCAGGAC-3'; Reverse, 5'-CCATCCTTTACCAAGGAGAC-3'), COL3A1 (Forward, 5'-GAAAGATGGGAGACTGGAC-3'; Reverse, 5'-CAGGTGTTGACCTGGTCG-3'), NLRP3 (Forward, 5'-GAGACGAAAAACGACACGTAAA-3'; Reverse, 5'-TACCAGGTATGCCTTGTAATCC-3'), Type I collagen (1: 1000), Type II collagen (1: 1000), and Type III collagen (1: 1000).

2.10. Hoechst staining

Apoptotic cells were stained with the Hoechst 33258 staining kit (Beyotime). Cells were plated in six-well cell culture plates. The cells were stained using the detection kit according to the manufacturer's instructions. Stained cells were examined using a fluorescence microscope. Five fields from each section were selected randomly. Image-Pro Plus 6.0 software (USA) was employed to quantify densitometric.

2.11. Immunofluorescence (IF)

Cells were cultivated to create cell climbing pieces. After 4% paraformaldehyde treatment, the cells were permeabilized with 0.5% Trition X-100 for 15 min, blocked for 1 hour with 5% BSA, and then incubated with primary antibodies overnight at 4°C. Then, the incubated primary antibody was recovered, and the slides were washed with PBS (5 min/cycle, for 3 cycles). After washing the membrane, the corresponding secondary antibody was added to the box and incubated at room temperature for 2 h. After the incubation was completed, TBST was used to clean the PVDF membrane again (5 min/cycle, for 3 cycles). Finally, ECL luminescent solution was used to detect the expression of the corresponding protein bands in the bioimage. ImageJ software was used to analyze the gray value of the protein band, and GAPDH was used as an internal reference to calculate the value. All antibody bodies and dilution concentrations were as follows: Caspase-1 (1: 1000), DNMT1 (1: 1000), NLRP3 (1: 1000), Type I collagen (1: 1000) and Type III collagen (1: 1000).

3. Results

3.1. Pathological and fibrotic changes in myocardial tissue of mice in an ISO-induced myocardial fibrosis model

We constructed a mouse myocardial fibrosis model by in-
DNMT1 methylation of ANRIL in myocardial fibrosis.

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jecting ISO subcutaneously. After two weeks, the HE staining (Fig. 1A), Masson staining (Fig. 1B), and Sirius red staining (Fig. 1C) revealed that the fibrous connective tissue in the ISO-induced myocardial fibrosis proliferated significantly, myocardial cells were significantly deformed, and collagen deposition increased when compared to the control group. Myocardial fibrosis-associated proteins COL1A1 and COL3A1 were detected by qRT-PCR, WB, and IHC. The results revealed that the mRNA expression of COL1A1 and COL3A1 in the myocardial tissue was significantly increased in the mice in the model group two weeks after modeling (Fig. 1E). Moreover, the protein expression of COL1A1 and COL3A1 in the myocardial tissue was significantly increased in mice from the model group (Fig. 1F-G). These findings support the ability of ISO to induce myocardial fibrosis.

3.2. Expression of DNMT1, LncRNA-ANRIL, Caspase-1, NLRP3 and COL1A1, COL3A1 regulation in ISO-induced myocardial fibrosis model in mice

Following ISO induction, qRT-PCR was used to verify LncRNA-ANRIL expression in cardiac tissues. Moreover, the expression of DNMT1, myocardial pyroptosis-related Caspase-1, and NLRP3 was verified by qRT-PCR, western blotting, and immunohistochemistry. The results of qRT-PCR revealed that LncRNA-ANRIL expression in the model group was significantly lower than that in the control group. However, mRNA expression of DNMT1, Caspase-1 and NLRP3 was significantly increased in the cardiac tissue of mice in the model group (Fig. 2A). WB and IHC analysis demonstrated that the model group significantly increased the protein expression of DNMT1, Caspase-1, and NLRP3 (Fig. 2B-C). These findings demonstrate that DNMT1 in myocardial fibrotic tissue can reduce LncRNA-ANRIL expression via methylation, thereby promoting myocardial tissue pyroptosis.

Primary cardiac fibroblasts were extracted, and LPS and ATP were used to construct a cellular pyroptosis model. The expression of COL1A1 and COL3A1 was validated using the model. qRT-PCR results demonstrated significantly lower mRNA levels of COL1A1 and COL3A1 in pyroptotic cells (Fig. 2D). WB and IF assays indicated significantly more protein levels of COL1A1 and COL3A1 in pyroptotic cells (Fig. 2E-F). These findings demonstrate that LPS and ATP cause CF cellular scorching and promote myocardial fibrosis.

3.3. Expression of DNMT1, LncRNA-ANRIL, and pyroptosis-related proteins in the LPS and ATP-induced pyroptosis model in cardiac fibroblasts

Hoechst 33258 was used to analyze apoptotic cells in mouse cardiac fibroblasts after they were treated with LPS, that the analysis revealed that compared to the control group, cardiac fibroblasts induced with LPS and ATP resulted in significantly more apoptotic cells (Fig. 3A). Subsequently, LncRNA-ANRIL expression in cardiac fibroblasts was verified by qRT-PCR after LPS and ATP induction. The expression of DNMT1, Caspase-1 and NLRP3 was verified using qRT-PCR, western blotting, and immunohistochemistry. qRT-PCR analysis revealed significantly lower expression of LncRNA-ANRIL in the model group than in the control group. However, pyroptotic cells significantly increased the mRNA expression of DNMT1, Caspase-1, and NLRP3 in cardiac fibroblasts (Fig. 3B). WB and IF detection results demonstrated that DNMT1, Caspase-1, and NLRP3 protein expression levels were significantly higher in pyroptotic cells (Fig. 3C-D). These findings demonstrate the methylation-induced role of DNMT1 in regulating LncRNA-ANRIL expression during cardiomyocyte pyroptosis.

3.4. Changes in DNMT1, LncRNA-ANRIL, expression, and pyroptosis-related proteins in cardiac fibroblasts after overexpression and silencing of DNMT1

DNMT1 was overexpressed in cardiac fibroblasts using
pEGFP-C1-DNMT1 and then treated the cells with an empty vector plasmid. qRT-PCR analysis revealed significantly reduced expression of LncRNA-ANRIL in the pEGFP-C1-DNMT1 group following DNMT1 overexpression, whereas the mRNA expression of DNMT1, Caspase-1, and NLRP3 was significantly increased in cardiac fibroblasts (Fig. 5A). WB and IF analysis revealed that the protein expression of Caspase-1 and NLRP3 was dramatically lower in the pEGFP-C1-ANRIL group compared to pyroptotic cells (Fig. 5B-C). Next, siANRIL was used to silence LncRNA-ANRIL in cardiac fibroblasts. qRT-PCR analysis revealed significantly reduced expression of LncRNA-ANRIL in the siANRIL group after silencing DNMT1, whereas the mRNA expression of Caspase-1 and NLRP3 was significantly increased in cardiac fibroblasts (Fig. 5D). WB and IF analysis demonstrated that the protein expression of Caspase-1 and NLRP3 was significantly reduced in the siANRIL group (Fig. 5E-F). These findings suggest that LncRNA-ANRIL can suppress pyroptosis in cardiac fibroblasts.

3.5. Changes in LncRNA-ANRIL expression and pyroptosis-related proteins in cardiac fibroblasts after overexpression and silencing of LncRNA-ANRIL.

We used pcDNA3.1-ANRIL to overexpress LncRNA-ANRIL in pyroptotic cells. qRT-PCR analysis revealed that LncRNA-ANRIL expression was dramatically increased, whereas Caspase-1 and NLRP3 mRNA expression was significantly decreased in cardiac fibroblasts (Fig. 5A). WB and IF analysis showed that the protein expression of Caspase-1 and NLRP3 was dramatically lower in the pcDNA3.1-ANRIL group compared to pyroptotic cells (Fig. 5B-C). Next, siANRIL was used to silence LncRNA-ANRIL in cardiac fibroblasts. qRT-PCR analysis revealed significantly reduced expression of LncRNA-ANRIL in the siANRIL group after silencing DNMT1, whereas the mRNA expression of Caspase-1 and NLRP3 was significantly increased in cardiac fibroblasts (Fig. 5D). WB and IF analysis demonstrated that the protein expression of Caspase-1 and NLRP3 was significantly reduced in the siANRIL group (Fig. 5E-F). These findings show that increased DNMT1 expression plays an important role in decreasing methylation-regulated LncRNA-ANRIL expression and can cause cardiac fibroblast pyroptosis.

4. Discussion

LncRNAs are involved in the development of various disorders and have been linked to cardiovascular disease [17-19]. In epigenetics, DNA methylation affects not only LncRNA expression but also myocardial fibrosis disease, which involves LncRNA [20].

There is no known link between LncRNA and pyroptosis in cardiac fibroblasts during myocardial fibrosis. The goal of this study was to figure out how DNMT1 methylation influences LncRNA-ANRIL regulation, impacting pyroptosis in cardiac fibroblasts. The analysis demonstrated that DNMT1 could inhibit the expression of LncRNA-ANRIL via methylation while also promoting the expression of NLRP3, a pyroptosis-associated inflammasome that contributes to cardiac fibroblast pyroptosis.

Fig. 3. Expression of DNMT1, LncRNA-ANRIL, Caspase-1, and NLRP3 in the LPS and ATP-induced pyroptosis model in cardiac fibroblasts. (A) Analysis of apoptosis by Hoechst staining (objective: 100×). (B) were qRT-PCR detection of LncRNA-ANRIL level, and mRNA levels of DNMT1, Caspase-1, and NLRP3 in cardiac fibroblasts. (C) WB detection of protein levels of DNMT1, Caspase-1, and NLRP3 in cardiac fibroblasts. (D) IF detection of protein levels of DNMT1, Caspase-1, and NLRP3 in cardiac fibroblasts (objective: 100×). Data were shown as mean ± SD (N = 3, per group). *p<0.05, **p<0.01 vs Control.

Fig. 4. Changes in DNMT1, LncRNA-ANRIL, Caspase-1, and NLRP3 expression in cardiac fibroblasts after overexpression and silencing of DNMT1. (A) qRT-PCR detection of LncRNA-ANRIL level, and mRNA levels of DNMT1, Caspase-1, and NLRP3 in cardiac fibroblasts. (B) WB detection of the protein levels of DNMT1, Caspase-1, and NLRP3 in cardiac fibroblasts. (C) IF detection of the protein levels of DNMT1, Caspase-1, and NLRP3 in cardiac fibroblasts (objective: 100×). Data were shown as mean ± SD (N = 3, per group). *p<0.05, **p<0.01 vs Control.
confirmed that DNMT1 methylation promotes NLRP3-mediated cardiac fibroblast pyroptosis inhibiting LcnRNA-ANRIL. These findings contribute to the rapidly expanding field of myocardial fibrosis and lay the groundwork for a new theoretical foundation and research path for myocardial fibrosis diagnosis and treatment.

5. Conclusion
The present findings suggest a role for DNMT1 in LcnRNA-ANRIL methylation and increasing the production of NLRP3, a pyroptosis-associated inflammasome, thereby promoting cardiac fibroblast pyroptosis and the progression of myocardial fibrosis.

Informed Consent
The authors report no conflict of interest.

Availability of data and material
We declared that we embedded all data in the manuscript.

Authors’ contributions
LZ and GL conducted the experiments and wrote the paper; KC and CX analyzed and organized the data; SK and WW conceived, designed the study and revised the manuscript.

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References

Fig. 5. Changes in LcnRNA-ANRIL, Caspase-1, and NLRP3 expression in cardiac fibroblasts after overexpression and silencing of LcnRNA-ANRIL. (A) qRT-PCR detection of LcnRNA-ANRIL level, and mRNA levels of Caspase-1 and NLRP3 in cardiac fibroblasts. (B) WB detection of protein levels of Caspase-1 and NLRP3 in cardiac fibroblasts. (C) IF analysis of the protein levels of Caspase-1 and NLRP3 in cardiac fibroblasts (objective: 100×). (D) qRT-PCR detection of the level of LcnRNA-ANRIL, Caspase-1, and NLRP3 mRNA in cardiac fibroblasts. (E) Western blot analysis of the Caspase-1 and NLRP3 protein levels in cardiac fibroblasts. (F) IF analysis of the protein levels of Caspase-1 and NLRP3 in cardiac fibroblasts (objective: 100×). Data were shown as mean ± SD (N = 3, per group). *p<0.05, **p<0.01 vs Control.


