Immunoglobulin kappa light chain produced by cardiomyocytes and participates in maintaining intercalated disc integrity

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Article

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Abstract

It is widely accepted that immunoglobulins (Igs) are produced only by B cells and function as antibodies. However, growing evidence has proven that almost all non-B cells also produce Igs with nonconventional roles, such as promotion of cell survival, proliferation and migration. In this study, we identified Ig light chain (κ chain) expression in mouse and human cardiomyocytes, especially on intercalated discs (ICDs). Unexpectedly, conditional knockout (cKO) of Igκ in adult cardiomyocytes in mice resulted in significant hypotension, a rapid decrease in cardiac contractility and conduction defects. Histologically, Igκ knockout in mouse cardiomyocytes led to structural disruption of intercalated discs (ICDs) and loss of localization of adhesion-related N-cadherin and CX43 on ICDs. Mechanistic investigation indicated that Igκ can bind with plectin/desmoplakin, a complex that connects desmin and desmosomes and enhances the protein stability of plectin. In conclusion, Our findings identify Igκ expressed by cardiomyocytes as a new ICD-related molecule that participates in cardiomyocyte contraction and conduction by stabilizing plectin.

Introduction

Immunoglobulins (Igs) are members of a classical immune molecule family and mainly exert antibody function. For several decades, B cells and differentiated plasma cells were thought to be the only sources of Igs. However, in recent years, many studies from our group and others have revealed that Igs are not only produced by B cells but also expressed by non-B cell lineages, including epithelial cells, neurons and germ cells of the testis1-8. In addition to serving as natural antibodies for immune defense, non-B Igs also exhibit diverse biological functions; for example, renal podocyte-derived IgG promotes cell adhesion and survival9,10. Cancer-derived IgGs are involved in the survival, adhesion, migration and metastasis of cancer cells by forming focal adhesions via binding to integrin11-13. These findings suggest that Igs are widely produced by many kinds of lineages other than B cells and that their physiological or pathological functions are diverse.

The heart is generally considered a dynamic organ and has the main function of maintaining blood circulation. So far, there is no evidence revealing that heart function is related to Igs. However, a number of clinical phenomena have shown that increased Igs are closely related to heart disease; for instance, IgE rapidly increases in the blood in acute myocardial infarction14,15, local IgM deposition occurs in injured cardiomyocytes16,17, and free Ig light chain (FLC) is frequently increased in the serum of patients with heart failure (HF) and has been suggested to be a novel marker of the prognosis of heart failure18. In particular, deposition of LC in heart tissue, which affects the mechanical and structural properties of heart tissue and exhibits direct toxicity to cardiomyocytes, is the pathogenic marker of incurable myocardial amyloidosis19-22. Although the significance of Igs in heart disease has been thoroughly studied, the origin of these Igs is currently thought to be restricted to B cells, which limits the understanding of the function of these Igs. Given that our studies and those by other groups have shown that Igs can be widely produced by non-B cells, we wondered whether cardiomyocytes can also produce Igs. In 2009, Mehta et al found Ig transcripts of both heavy chain and light chain in mouse heart tissue and HL-1 cells (mouse
myocardial cells)\textsuperscript{23}. In 2017, we reported that IgM heavy chain exists in cardiomyocytes from both WT and B cell-deficient (\(\mu\)MT) mice and that ischaemia and hypoxia promote the expression and secretion of IgM\textsuperscript{24}. These studies demonstrated that Igs can be produced by cardiomyocytes, and the physiological function of cardiomyocyte-derived Igs (CM-Igs) needs to be determined.

Classical Ig consists of two identical heavy chains and two identical light chains. There are five types of heavy chains, including IgM, IgG, IgA, IgD and IgE. There are two types of light chains, namely, k and l. Each of the light chains can form a complete Ig molecule with any of the heavy chains mentioned above, but the frequency of the two light chains in mice is different, and their ratio is 20:1. In fact, in our previous work, we found that IgM as well as IgG, IgA and IgE are expressed in cardiomyocytes. However, the main Ig light chain in cardiomyocytes is Igk. Therefore, in this study, we focused on demonstrating Igk expression by cardiomyocytes and identifying the function of Igk. We found that Igk acts as a novel cardiac ICD-related protein to maintain the structural integrity of ICDs and the cardiac functions of the adult heart.

**Experimental Procedures**

**Animal and human heart specimen**

Adult Balb/c mice and C57BL/6 mice (8 weeks) were obtained from Vital River (Charles River Laboratories) and \(\mu\)MT mice (Balb/c background) were a gift from Professor Zhihai Qin (Institute of Biophysics, Chinese Academy of Sciences). Mice were perfused with PBS for 5 minutes, and the heart was removed. Conditions of animal housing and all experimental procedures were conducted under institutional guidelines provided by the Institutional Animal Care and Use Committee of China.

A human heart autopsy specimen was a gift from Qingyuan Liao (Guilin Medical University) and human iPS-CM was a gift from Feng Lan (Anzhen hospital).

**HL-1 myocardial cell culture**

HL-1, an adult mouse atrial myocardial cell line, was a gift from Dr. W.C. Claycomb (Louisiana State University, New Orleans, LA)\textsuperscript{25}. HL-1 cells were cultured in Claycomb medium (Sigma-Aldrich) supplemented with 10% FBS (HyClone), 100 U/ml penicillin, 100 \(\mu\)g/ml streptomycin, 2 mM L-glutamine, and 0.1 mM norepinephrine (Sigma-Aldrich) at 37°C under 5% CO2 in 0.02% gelatin coated flasks containing 0.5% fibronectin (Sigma-Aldrich).

**Isolation and culture of neonatal mouse ventricular myocytes**

Ventricular cardiomyocytes were isolated from the left ventricles of neonatal mice using enzymatic digestions containing 0.08% trypsin (Gibco BRL) and 0.1% collagenase (Sigma-Aldrich) for 8 min at 37 °C. Isolated cardiomyocytes were re-suspended in Dulbecco’s modified Eagle’s medium (DMEM) containing 10% fetal bovine serum (FBS) and 1% penicillin–streptomycin and then plated at a density of
6 × 10^5/dish and cultured for 1.5 h. The supernatant was subsequently collected and centrifuged at 500g for 10 min and resuspended in 5 ml culture medium containing 100 mmol/L 5’BrdU (Sigma-Aldrich) and plated in 35 mm culture dishes for 48 h.

**Isolation of adult mouse ventricular myocytes and single cardiomyocyte dissociation.**

Ventricular myocytes were isolated from the Langendorff-perfused hearts of adult C57BL/6 mice. Briefly, the mice were anaesthetized with pentobarbital and the heart was removed and mounted on the Langendorff apparatus and perfused with Ca^{2+}-free Tyrode's solution to eliminate the blood. Then the heart was perfused with Tyrode's solution containing 0.67mg/ml of collagenase type II (Worthington) for 30 min for digestion. When the heart became flaccid, remove the ventricle and cut into small pieces and then pelleted by centrifugation at 500rpm for 1 min at room temperature and resuspended in Tyrode's solution. The calcium concentration was then gradually increased to 500mM over 80 min. The supernatant was discarded and the pellet was resuspended in PBS containing 1mg/ml BSA. Cardiomyocyte single cells were manually picked under the microscope by mouth pipette. To make sure that only single cells were collected, the solution was visually inspected under the microscope and was discarded if multiple cells were observed. Volume of liquid was kept below 0.5 μl. Cells were then transferred to a 0.2 ml thin-wall PCR tube containing lysate buffer.

**RNA extraction and cDNA synthesis**

Myocardial single cell RNA extraction and cDNA synthesis were carried out according to Tang’s methods as described^{26,27}. In briefly, myocardial single cells were picked by capillary pipette and reverse transcription was then performed directly on the whole cell lysate. After this, the free primers were removed by ExoSAP-IT and a poly(A) tail was added to the 3’ end of the first-strand cDNA by Terminal Deoxynucleotidyl Transferase. Then the single cell cDNAs were amplified following PCR program: 95 °C for 3 min, then 27 cycles of 95 °C for 30 sec, 67 °C for 1 min, and 72 °C for 6 min (+ 6 sec each cycle). After this step, all cDNAs have been amplified. The single cell cDNA PCR product can be saved at −80 °C for 2 months.

Total RNA was extracted from frozen heart tissue as well as HL-1 cells by using Trizol Reagent (Invitrogen). Reverse transcription (RT) was carried out with the First Stand cDNA Synthesis Kit (Thermo Scientific) according to the manufacturer’s protocol.

**PCR amplification and DNA sequencing**

After the cDNA was obtained, specific primers were used to amplify the Igk, CD20, Myh6, Myh7, cTnT, cTnI and GAPDH. The sequences of all PCR primers are listed in Supplementary Table 1. Touchdown PCR was performed for amplification. PCR products were cloned into a pGEM-T Easy Vector (Promega). These clones were analyzed by the Sanger method using an ABI 3100 Genetic Analyzer (Applied Biosystems). The sequences of VkJk were compared with those in the BLAST and Immunogenetics databases to
identify the best matches for germline gene segments and VJ junctions. All VkJk sequences were shown in supplementary table 2.

The DNA concentration of eluted PCR products were measured and 146 ng of DNA was pooled for sequencing. The following sequencing using the 2 × 250 bp Illumina MiSeq platform were performed by Novogene Corporation (Beijing, China). Single cell of WT mice (n=17) and mMT mice (n=3) was analyzed. The usage of Vk was shown in supplementary table 3.

RNA interference

The sequence for Igk siRNA was as follows: siRNA-1: GAAGATTGATGGCAGTGAA. siRNA-2: GAACGACATAACAGCTATA. These were used to transfect HL-1 cells with the Lipofectamine 3000 kit (Invitrogen) according to the manual; a mock group was transfected with control siRNA. Cells were harvested 24 h and 48 h after transfection for Igk mRNA and protein detection.

Immunofluorescence

Frozen sections of heart tissues, primary cultured cardiomyocytes, and HL-1 cells were fixed with acetone, and blocked with 5% FBS at room temperature for 30 minutes. The cells were then stained overnight at 4 °C with the following primary antibodies: rabbit anti-mouse Igk (1:50; Proteintech), rabbit anti-mouse Cx43 (1:100; Proteintech), rabbit anti-mouse N-cadherin (1:100; Proteintech), rabbit anti-mouse Plakoglobin (1:100, Proteintech) were added at 4°C overnight. Then, the cells were washed with PBS and incubated for 1 h with the respective secondary antibodies conjugated to 5 (1:1000; Abcam), Alexa Fluor 568 (1:1000; Abcam) or Alexa Fluor 594 (1:1000; Abcam) after which the cells were stained with Hoechst (1:10000; Sigma) for 5min at room temperature.

Western blot analysis

Total cellular protein was extracted from heart tissue and HL-1 cells using RIPA lysis buffer (Beyotime) containing a protease inhibitor cocktail. The samples were subsequently separated by 12.5% SDS-PAGE and transferred onto nitrocellulose membranes. The membranes were then incubated with the following primary antibodies: Biotin-labeled goat anti-mouse Igk (1:1000; SouthernBiotech), rabbit anti-mouse Cx43 (1:1000; Proteintech), rabbit anti-mouse N-cadherin (1:1000; Proteintech), rabbit anti Desmoplakin (1:1000, Proteintech), GAPDH (1:1000, ZSGB-Bio) and b-actin (1:1000, ZSGB-Bio). The secondary antibodies conjugated with HRP (ZSGB-Bio) were applied at a concentration of 1:10,000 and visualized using with ECL chemiluminescence (GE). Western Blots were quantified by densitometry using ImageJ software.

Generation of conditional cardiac-specific Igk-knockout mice

To generate inducible cardiac-specific Igk cKO mice, transgenic mice expressing a tamoxifen-inducible Cre recombinase protein under the control of the α-myosin heavy chain promoter, αMHC/MerCreMer mice (MCM) were intercrossed with Igk<sup>fl<sub>ox</sub>/fl<sub>ox</sub></sup> mice (obtained from Shanghai Biomodel Organism Science &
Technology Development Co., Ltd) in a C57BL/6 genetic background. To induce Cre recombination, 8- to 12-week-old male aMHC/MerCreMer; Igk<sup>flox/flox</sup> (Cre<sup>+</sup>;Igk<sup>f/f</sup>) were treated with 80mg/kg of tamoxifen (Sigma) via intraperitoneal injection once a day for 4 consecutive days. aMHC-Cre<sup>+</sup>; Igk<sup>f/f</sup> mice, aMHC-Cre<sup>+</sup> mice with tamoxifen age- and sex-matched littermates were utilized as control mice. Conditions of animal housing and all experimental procedures were conducted under institutional guidelines provided by the Institutional Animal Care and Use Committee of China.

**Construction of adenovirus**

The full-length cDNA of mouse IgkV17-121/J2/C was inserted into adeno-associated virus serotype 9 vector-cTnT (AAV9-cTnT) to construct the AAV9-cTnT-Igk virus (produced by Likeli, Beijing, China). 5x10<sup>11</sup>vg of AAV9-cTnT-Igk or AAV9-CMV-GFP were injected intravenously into 10-week-old cardiac Igk knockout mice (αMHC-Cre<sup>+</sup> Igk<sup>f/f</sup>), while the same volume of AAV9-CMV-GFP was injected intravenously into αMHC-Cre<sup>+</sup> mice. Four weeks after injection, tamoxifen was injected to induce Igk gene knock down. Western blot of myocardial tissues was performed to confirm the re-expression efficiency.

**Systolic blood pressure analysis**

Systolic blood pressure was measured in conscious mice using BP-98A Blood Pressure Analysis System (Softtron). Briefly, animals were placed in a plastic chamber maintained at 37°C, and a cuff with a pneumatic pulse sensor was attached to the tail. we measured systolic blood pressure five times per animal and recorded the mean overall blood pressure.

**Echocardiography analysis.**

To evaluate left ventricular function and dimension, transthoracic two-dimensional echocardiography was performed on mice sedated with 1.5% isoflurane using a Visual Sonics Vevo 770 ultrasound system (Visual Sonics) equipped with a 17.5-MHz linear array transducer. M-mode tracings in the parasternal short axis view were used to measure left ventricular posterior wall thicknesses (LVPW) and left ventricular inner dimension (LVID) at end-diastole, which were used to calculate left ventricular fractional shortening (FS) and the ejection fraction (EF) according to standard formulas.

**Sarcomere shortening analysis**

Isolated cardiomyocytes were placed on the experimental chamber of an inverted microscope (Olympus) with permanent perfusion of Tyrode solution containing 500mM Ca<sup>2+</sup> heated to 37°C. Myocytes were electrically stimulated (15V, 0.5hz) and sarcomere shortening was detected using a video-based detection system (IonOptix) on intact myocytes. Six twitches per myocyte were collected for each sample (n =2). Signal averaged data were analyzed to determine resting sarcomere length, peak shortening normalized for resting sarcomere length (percent peak height), time to peak shortening (TTP).

**Electrocardiography analysis**
Surface electrocardiogram (ECG) recordings were performed on mice anaesthetized with 1–1.5% pentobarbital sodium. Needle electrodes were used and placed in the conventional lead II position. A differential amplifier amplified the signals in the bandwidth of 0.1–1000 Hz, and signals were filtered using an adaptive 60 Hz filter. The signals were digitized at 3000 Hz, and analyzed using ECG Analysis Software (QRS phenotyping, Calgary, Canada).

**Electron microscope analysis**

The heart tissues fixed with 3% glutaraldehyde were chopped into 1 mm$^2$ x 1 mm pieces. Then, the tissues were rinsed, dehydrated, embedded, sliced, stained, and subsequently investigated under a transmission electron microscope (TEM).

**Statistical analysis**

All data are presented as the means±s.e.m. Statistical analyses were performed using Prism Software. The statistical significance of the difference between two sets of data was assessed using an unpaired t-test. Statistical significance was defined as P < 0.05. * P< 0.05, **P< 0.01, ***P< 0.001.

**Data Availability**

This study includes no data deposited in external repositories

**Results**

**Igk expression in mouse and human cardiomyocytes**

To determine whether Igk is present in cardiomyocytes in the mice, immunofluorescence and immunogold electron microscopy were performed by using anti-mouse Igk antibodies. Positive staining for Igk was clearly observed in cardiomyocytes, especially on intercalated discs (ICDs) in heart tissue (Fig 1Aa, c), and Igk was colocalized with plakoglobin, which is a specific marker of ICDs (Fig 1Ab). Given that B cells are considered the only sources of Igs, B cell-deficient mice (mMT mice) were used to exclude disturbance of B-Ig. As expected, Igk was also observed in the cardiomyocytes of mMT mice by immunofluorescence and Western blotting. As shown in Fig 1Aa and Fig 1B, Igk was found on ICDs in mMT hearts, and the signal intensity was comparable with that in WT mice. This indicates that Igk on ICDs originated from non-B cells. Subsequently, HL-1 myocardial cells and primary cultured cardiomyocytes from newborn mice were used to further confirm the existence of Igk in cardiomyocytes, as illustrated in Fig 1C and Fig 1D. Igk was localized on the cross-striations of primary cultured cardiomyocytes from neonatal mice and the cell junctions of HL-1 cells. Moreover, knockdown of Igk by siRNA resulted in reductions in Igk levels and Igk staining at cell junctions, suggesting that Igk can be produced by the cardiomyocytes of mice (Fig 1D). To demonstrate that Igk can also be expressed in human cardiomyocytes, 3 normal hearts and 3 viral myocarditis hearts obtained from autopsies were used for immunohistochemical staining, and Igk was found to be localized on ICDs of cardiomyocytes.
To further confirm that Igk is produced by human cardiomyocytes, Igk was detected in human iPS-CM (induced pluripotent stem cell-derived cardiomyocytes) by IF and RT-PCR; significant staining for Igk was observed on the cell membrane, and Igk was found to be colocalized with plakoglobin (Fig 1E).

Surprisingly, we identified free Igk (k free light chain, kFLC) as the main form of Igkin heart tissue, as revealed by nonreduced SDS-PAGE/Western blotting (Fig 1Fa). Interestingly, ischaemia promoted the secretion of k FLC in cardiomyocytes (Fig 1 Fb, c).

Igk transcripts in cardiomyocytes with restricted diversity compared with that of B cells

To further identify whether Igk was synthesized by cardiomyocytes, the rearranged transcripts of Igk were detected in cardiomyocytes by RT-PCR using primers for the variable regions and constant regions of the Igk gene. First, adult cardiomyocytes were isolated from wild-type (WT) or mMT (B cell-deficient) mice, and the rearranged Igk transcripts were successfully amplified from cDNA libraries of cardiomyocytes (Fig 2Aa, b). Sequencing analysis showed that all the cardiomyocyte-derived Igk transcripts we analysed had classical V-J rearrangements. Interestingly, unlike the B cell-derived Ig that showed high diversity, the V-J rearrangement pattern of cardiomyocytes showed very restricted diversity (Fig 2Ac, d). Similarly, the V-J rearrangement pattern in cultured cardiomyocytes isolated from neonatal mice showed a conservative sequence that was different from that in fibroblasts isolated from the same heart (supplementary Fig 1A, B). In addition, the Igk transcript was also detected in HL-1 cells, with IgkV17-121/J2 being the dominant VJ pattern (Fig 2C).

Subsequently, single cardiomyocyte RNA-seq analysis was applied to further confirm Igk expression and sequence characteristics. Single cardiomyocytes from either WT or mMT mice were manually picked under microscopy by a mouth pipette. Cardiomyocytes were easily distinguished by and picked based on their unique rod-like cell shapes, and total RNA was obtained and reverse transcribed into cDNA according to Tang's masterpiece (supplementary Fig 1C). Then, CD20 transcripts were amplified to exclude single-cell cDNA contamination from B cells, and Myh6, cTnT and cTnI transcripts were amplified to identify cardiomyocyte. Additionally, Igk transcripts were sequenced by Sanger sequencing and next-generation sequencing. We isolated 56 single cardiomyocytes from both WT and mMT mice and found that 73% (42/56) of single cardiomyocytes showed typical rearranged Igk sequencing (Fig 2Da, b, supplementary Fig 1D, F). Interestingly, we found that 30% (13/42) of single cardiomyocytes expressed more than one functional VkJk pattern (Fig 2D f). We compared the frequency of Vk and Jk usage in cardiomyocytes with that of B cells and found that VJ usage was quite different between cardiomyocytes and B cells. The cardiomyocyte-derived Igk gene showed an obvious restricted pattern of VJ combinations. In detail, the dominant Vk usages in cardiomyocytes were IgkV17-121 (28.3%, 95/336), IgkV9-120 (25%, 84/336), and IgkV14-100 (15.5%, 52/336), but low frequency was used by B cells (Vk17-121 (2.1%), Vk9-120 (2.5%) and Vk14-100 (0.6%)) (data from https://www.10xgenomics.com/). However, the B cell-derived Vk pattern showed diversity, and the dominant Vk usage in B cells was Vk10-96 (6%), which was not detected in cardiomyocytes (Fig 2Dc-h). More interestingly, Vk17-121 and Vk9-120 localize next to each other on the genome, suggesting that this chromosomal region may be open and more easily
transcribed

supplementary Fig 1E). Additionally, we also analysed the sequence of the J segment usage and V-J junction between CM-Igk and B-Igk and found that the dominant usage of the J segment is Jk1 in B cells and Jk2 in cardiomyocytes (Fig 2Di). The V-J junction analysis showed that the sequences of the V-J junction in IgkV9-120/J2 and IgkV14-100/J2 were quite the same between CM-Igk and B-Igk. Interestingly, the sequences of the V-J junction in IgkV17-121/J2 were quite different between CM-Igk and B-Igk, with 63% (46/73) of the CM-IgkV17-121 segment and J2 segment showing a direct connection; however, there was a deletion of nucleotides in the V terminal and J terminal at the V-J junction of B-IgkV17-121/J2 (66/91) (supplementary Fig 1G).

Cardiomyocyte-specific Igk knockout results in decreased cardiac conduction and contractility

To evaluate Igk function in cardiomyocytes, a tamoxifen-induced Igk cKO mouse was established by intercrossing Igk<sup>f/f</sup> mice with a-MHC-Cre mice (Fig 3A). Igk cKO mice and their littermates (8- to 12-week-old), including Igk<sup>f/f</sup> and a-MHC-Cre mice, were administered tamoxifen for four consecutive days. Two days after tamoxifen treatment, we first analysed cardiac function by analysing systolic blood pressure, and unexpectedly, we found that cKO mice displayed significantly reduced blood pressure (Fig 3C), indicating that heart function was impaired. Additionally, echocardiographic analysis showed that the contractility of the cKO mice was markedly impaired compared with that of the control mice, with cKO mice showing reductions in ejection fraction (% EF; 51.87% ± 4.355 for cKO mice versus 68.26%± 2.219 for control mice) and LV fractional shortening (% FS; 24.91% ± 3.061 for cKO mice versus 36.52% ± 1.584 for control mice) (Fig 3D). Moreover, a contractile shortening assay showed that isolated single cardiomyocytes from cKO mice exhibited a reduced contractile response compared with those from control mice (0.0422± 0.006719 for cKO mice versus 0.1520 ± 0.04271 for control mice) (Fig 3E). More importantly, we found that ablation of Igk in cardiomyocytes led to the death of 40% cKO mice as early as d4 (n = 16). In contrast, tamoxifen-treated control mice (Igk<sup>f/f</sup>) appeared healthy throughout the course of tamoxifen treatment (Fig 3B). These data suggest that the expression of Igk in cardiomyocytes is critical for heart function.

Cardiac-specific Igk knockout causes structural abnormalities of intercalated discs

Intercalated discs are made up of short processes extending from two adjacent branches of myocardial fibers, which can spread excitation from one cell to another and facilitate synchronous contraction of myocardial fibers. Subsequently, we evaluated the electrophysiological activities of cardiomyocytes by electrocardiogram (ECG) and found that cKO mice exhibited a significant reduction in resting heart rate and QRS interval prolongation (Fig 3F). To further examine the significance of the reduced Igk levels and cardiac dysfunction triggered by cardiac stress, cKO and control mice were subcutaneously injected with isoproterenol (ISO). As expected, ISO injection induced more frequent atrioventricular (AV) block in cKO mice (6/6) than in control mice (1/6). Furthermore, unlike in control mice, ISO did not significantly increase the heart rate of cKO mice, which indicated that Igk deficiency decreases the response of cardiomyocytes to ISO.
Given that Igk is localized on ICDs and Igk deficiency induced severe conduction dysfunction, we next used transmission electron microscopy (TEM) to analyze the ultrastructural of ICDs in the hearts of cKO and control mice. We found that the structures of the ICDs of cKO mice were highly convoluted and disorganized with widened gaps from day 2 after tamoxifen administration (Fig 4A). However, normal ICD structures with clear adherens junctions and desmosomes were visible in the hearts of control mice. The results indicated that Igk plays a very important role in maintaining ICD structure. The components of ICDs, namely, plakoglobin (a cytoplasmic desmosomal protein), N-cadherin (an adherens junction protein) and connexin 43 (a gap junction protein), were examined by immunofluorescence staining and stimulated emission depletion microscopy (STED). Our results showed that depletion of Igk in ICDs resulted in a significant decrease in N-cadherin and Cx43 staining in ICDs, while plakoglobin staining was increased in the ICDs of cKO mice compared to control mice (Fig 4B, C, D, E, supplementary Fig 2). Interestingly, an obvious decrease in N-cadherin occurred on the third day after tamoxifen administration, while Cx43 was decreased on the ICD on the fourth day. Moreover, Western blot analysis further confirmed a reduction in Cx43 and N-cadherin in the hearts of cKO mice compared with control mice (Fig 4F). These results indicated that Igk is required to maintain the localization of N-cadherin and Cx43 on ICDs.

To further confirm Igk function in ICDs, we overexpressed Igk 4 weeks before tamoxifen administration by generating AAV9-cTnT-Igk in Igk{f/f};a-MHC-Cre mice. Igk was knocked out in the cardiomyocytes of Igk{f/f};a-MHC-Cre mice with tamoxifen. Rescue of Igk by AAV9-cTnT-Igk significantly improved cardiac function by increasing the % EF (% EF; 44.04 ± 9.591 for cKO mice versus 70.57 ± 4.090 for cKO mice injected with AAV9-cTnT-Igk and 73.34 ± 4.473 for control mice) and % FS (% FS; 21.89 ± 5.217 for cKO mice versus 40.01 ± 3.379 for cKO mice injected with AAV9-cTnT-Igk and 42.81 ± 3.645 for control mice) (Fig 5A) and reversed the destruction of ICD structure by upregulating the protein levels of Igk, DSP, N-cadherin and Cx43 (Fig 5B, C, supplementary Fig 3), confirming that dysfunction of cardiomyocytes and abnormal ICD structure of was caused by Igk deficiency.

Cardiac-specific Igk knockout induces degradation of plectin, which causes disconnection of desmin and desmosomes

To understand the mechanism of how Igk affects the structure and function of ICDs, we wanted to identify the proteins that interact with Igk in cardiomyocytes. Total heart tissue lysates were extracted, GST-Igk or GST (as a control) was used to identify the proteins that interact with Igk by GST pull down, and protein mass spectrometry analysis was used to identify the potential interacting proteins. Unlike in the control group, desmoplakin and its binding protein plakoglobin bound with GST-Igk in the ICDs of Igk cKO mice (Fig 6A), suggesting that Igk interacts with desmoplakin. Desmoplakin associates with plakoglobin, plakophilin, b-catenin and a-catenin to form a complex that connects desmosomes and adherens junction (AJs) within the ‘area composite’ and work together to connect ICDs to the cytoskeleton, especially actin and intermediate filaments. Disconnection of actin or desmin with ICDs leads to a loss of desmosomes and AJ components 28,29. Therefore, we also determined the localization of desmin and actin in cardiomyocytes, and the immunofluorescence results showed that staining for
desmin and actin on ICDs was diminished in heart tissues from Igk cKO mice compared to control mice (Fig 6Ba, c). Similarly, the filamentous structures of desmin and F-actin in HL-1 cells was abolished when Igk was knocked down by specific siRNA (Fig 6Bb, d), suggesting that a lack of Igk led to the lost connection between two cytoskeleton components actin and desmin and ICDs. In our previous finding, we found that cancer cell-derived Igk interacts with the cytolinker protein plectin and that a lack of Igk leads to the disappearance of plectin and disassembly of actin as well as intermediate filaments. Plectin is a 500-kDa cytoskeleton-associated protein, and the C-terminus of Plectin can link the intermediate filaments to the transmembrane proteins of hemidesmosomes and maintain cell-to-cell connectivity. Moreover, the N-terminus of plectin can link actin to promote its assembly, and a lack of plectin can lead to disassembly of intermediate filaments and actin\textsuperscript{30,31}. Moreover, it has been reported that plectin can connect desmin with desmoplakin on ICDs\textsuperscript{32,33}. Thus, we analysed whether there is an interaction between Igk and plectin on ICDs. The immunofluorescence results revealed that Igk and plectin were colocalized on the ICDs of cardiomyocytes, as well as on the cell junctions of HL-1 cells (Fig 6Da). Knockdown of Igk led to decreased plectin and loss of localization on ICDs (Fig 6Db-d). Next, we established an expression vector containing the C-terminus of plectin, and IP assays showed that Igk interacts with both desmoplakin and plectin to form a complex (Fig 6C).The results of our protein stability experiments using cyclohexane (CHX) to inhibit protein synthesis revealed that Igk can maintain the stability of plectin, which can explain how Igk knockdown results in the lost connection between cytoskeleton and ICDs (Fig 6E).

**Discussion**

We reported for the first time that the Igk light chain, which is currently believed to be a classical immune molecule, can be widely expressed in both mouse and human cardiomyocytes. Functionally, cardiomyocyte-derived k-type light chain, as a novel structural protein of cardiomyocytes, mainly plays an important role in maintaining the normal structure of cardiomyocyte intercalated discs and normal myocardial contraction and conduction.

In fact, there are many clues that cardiomyocytes can produce Igs, but due to the limitations of classical immunological theory, researchers still believe that Igs are produced by B cells. However, there are many contradictory phenomena that cannot be well explained by classical immunological theory. For example, in acute myocardial infarction, increased free Ig light chain or IgE appears rapidly in the blood, which is not consistent with the law of antibody production of B cells; some myocardial injury diseases are associated with high levels of "autoantibodies against cardiac structural proteins", but to date, scientists have not found a B cell clone that can produce these autoantibodies. Our findings suggest that the "autoantibodies against cardiac structural proteins" in myocardial injury diseases may be produced by injured cardiomyocytes and may be involved in unidentified activities rather than acting as autoantibodies. Our previous work revealed that cardiomyocytes can widely express different classes of Ig, including IgM\textsuperscript{24}, IgG and IgE, as well as the Igk light chain, but their physiological function remains unclear. Therefore, in this study, we focused on Igk to reveal its physiological function in cardiomyocytes.
In this study, we first showed evidence of Igk expression in cardiomyocytes at the protein and transcription levels. Our results revealed that Igk was widely expressed in mouse and human cardiomyocytes; moreover, Igk was mainly located on the intercalated disc. Subsequently, we analysed the characteristics of variable regions of cardiomyocyte-derived Igk. It is well known that the Ig variable region produced by B cells is infinitely diverse because its main function is to bind to a variety of antigen molecules, such as antibodies or BCR. Therefore, we first analysed whether the characteristics of the cardiomyocyte-derived Igk variable region are different from those of Igk in B cells. Our single cell sequencing results showed that, unlike B cell-derived Igk, which shows diversity, cardiomyocyte-derived Igk exhibited a very conservative VkJk usage and pattern and tended to express IgkV17-121 (28.3%, 95/336), IgkV9-120 (25%, 84/336), and IgkV14-100 (15.5%, 52/336) in cardiomyocytes in adult mice. It is worth noting that the conservative characteristics of cardiomyocyte-derived Igk were similar to those of Igk in hepatocytes and other non-B cells, especially IgkV9-120, which has been found to be highly frequently expressed in other non-B cell lineages. However, the expression frequency of the above VkJk usages in B cells was less than 3%. This limited diversity suggests that cardiomyocyte-derived k light chain may not participate as an antibody in humoural immunity but may play another important physiological role.

Next, we constructed conditional mice with conditional Igk knockout in cardiomyocytes (Igk^{f/f}; aMHC-Cre^+ mice) to determine the physiological function of Igk. To prevent the occurrence of light chain redundancy in the early development of mouse cardiomyocytes, the conditional Igk knockout was inducible, allowing the knockout of the Igk gene from cardiomyocytes at any stage of mouse development by tamoxifen treatment. In this study, we selected 8-week-old mice for conditional knockout. We first found that compared to control (Igk^{f/f} or aMHC-Cre^+) mice, Igk knockout mice exhibited significant decreases in blood pressure, cardiac ejection ability, and heart rate from day 2 after the administration of tamoxifen. Isoproterenol (ISO) was used to increase cardiac oxygen consumption and heart rate and led to atrioventricular block. Importantly, Igk knockout resulted in the death of over 40% of Igk cKO mice from day 4 after the administration of tamoxifen, suggesting that Igk is involved in maintaining normal cardiac contraction and conduction function.

To evaluate the mechanism, we observed changes in myocardial structure in Igk cKO mice. Our results showed that Igk knockout led to disordered intercalated disc structure and loss of localization of N-cadherin, an adhesion junction effector, and Cx43, a gap junction effector, at intercalated discs, suggesting that the abnormal myocardial contraction and conduction caused by Igk knockout may have been caused by structural disorder of the intercalated discs. To further confirm this result, we used AAV9-cTnT-Igk to overexpress Igk. The results showed that when cardiomyocytes overexpressed Igk, knockout induced by tamoxifen did not cause disorder of cardiomyocyte structure or function, suggesting that the above phenomenon was indeed caused by Igk knockout in cardiomyocytes.

ICDs are composed of three major complexes: desmosomes, adherens junctions (AJs) and gap junctions (GJs). Desmosomes and AJs provide mechanical attachment between two cardiomyocytes by anchoring the actin cytoskeleton and intermediate filaments (IFs), enhancing the stability of the ICD. GJs connect
the cytoplasm of adjacent cardiomyocytes metabolically and electronically to enable the propagation of electrical stimuli throughout heart muscle cells. These three complexes work together to incorporate the mechanical and electrical functions of the ICD. Notably, components of desmosomal and adherens junctions are intimately associated within the area composita and work together to facilitate mechanical coupling. In the area composita, desmosomes are considered to be associated with the intermediate filament desmin through binding desmoplakin, and AJs are considered to be associated with actin through binding a-catenin, b-catenin and vinculin. However, how does Igk knockout affect the localization and expression of ICD-related proteins? Our study found that Igk can interact with desmoplakin. Igk knockout first led to a reduction in N-cadherin and then in Cx43 and a loss of localization on ICDs, suggesting that Igk knockout first affects desmosome and AJ function and then secondarily affects GJs. To address why Igk knockout results in instability of ICDs, we further determined the intracellular localization of both F-actin and desmin and found that Igk knockout also resulted in disassembly of both F-actin and desmin. The results suggest that disassembly of both F-actin and desmin may be the cause of instability of ICDs resulting from Igk knockout. To date, no evidence has indicated that desmoplakin function is related to actin, which suggests that there may be other proteins related to Igk; this study suggested that desmoplakin interacts with Igk. Plectin is a large cytolinker protein, and its main function is to interact and promote the assembly of actin and intermediate filaments. It has been reported that plectin is localized on ICDs and connects intermediate filaments/desmin to desmosomes by interacting with desmoplakin. Moreover, our previous work demonstrated that cancer cell-expressed Igk can interact with plectin (unpublished). Based on the above evidence, we hypothesize that cardiomyocyte-expressed Igk can interact with plectin, which may be involved in the stability of ICDs. Our results showed that Igk can interact with plectin and that Igk gene knockout results in a significant loss of plectin and disassembly of desmin and F-actin. Moreover, a loss of plectin leads to disruption of AJs and GJs, which leads to abnormal electrical coupling of cardiomyocytes and reduced conduction velocity. Our mechanistic study revealed that Igk can promote the stability of plectin on ICDs. However, the mechanism by which Igk regulates plectin and its role in ICD assembly still require further investigation.

Our study has only broken the ice, and it is believed that there are many physiological and pathological functions of cardiomyocyte-expressed Igs that have yet to be revealed. For example, myocardial amyloidosis is a common heart disease that is caused by pathogenic FLCs. At present, it is generally accepted that these pathogenic Ig light chains are secreted and deposited into myocardial tissue by multiple myeloma cells; however, in most cases, no malignant proliferation of plasma cells is found in myocardial amyloidosis. In contrast, we found that cardiomyocytes can secrete FLC under conditions of cell stress, such as hypoxia. Our study provides novel insight into the source of abnormally elevated FLC in myocardial amyloidosis and myocardial injury, such as myocardial infarction and viral myocarditis. However, the exact pathological mechanism of cardiomyocyte-derived Igs is still unclear and requires further investigation.

Declarations
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Author contributions

ZZ and QXY conceived the study and designed the workplan. Experimental work: ZZ, SZ, ZWJ. Development and contribution of reagents: LF, LQY. Manuscript writing: ZZ and QXY.

Conflict of interest

None

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**Supplementary Materials**

**Supplementary Fig 1. Igk transcript was found in cardiomyocyte**

(A) a. Isolated cardiomyocyte of neonatal mice. Myosin was used to identify cardiomyocyte. Bar=20mm.
   b. Transcript of Igk variable region and constant region in cardiomyocyte and fibroblast by RT-PCR. Bar=20 mm.

(B) Igk knockdown on mRNA levels in HL-1 cells by two independent siRNA and Igk transcript was detected by RT-PCR.

(C) a. Schematic overview of the single-cell sequencing workflow in cardiomyocyte. b. single cardiomyocyte was picked by a mouth pipette under microscopy.

(D) a. Transcript of Igk was detected in single cardiomyocyte. Spleen of WT mice was used as positive control. b. Transcript of CD20 was detected in single cardiomyocyte. Spleen of WT mice was used as positive control. c. Transcript of cTnT was detected in single cardiomyocyte. Heart tissue of WT mice was used as positive control.

(E) Igk V segment distribution on the chromosome.

(F) Sequence of Igk J region to C region which was amplified in sFig 1C.

(G) Sequence of Igk CDR3 region of CM-Igk and B-Igk.

**Supplementary Fig 2. Colocalization of N-cadherin/Cx43 and Plakoglobin on the ICD of Igk cKO and control mice.**

(A) a. Colocalization of N-cadherin and plakoglobin on the ICD of cKO and control mice for consecutive four days after tamoxifen injection by confocal. B. Colocalization of Cx43 and plakoglobin on the ICD of cKO and control mice for consecutive four days after tamoxifen injection by confocal. Bar=25 mm.

**Supplementary Fig 3. Colocalization of N-cadherin/Cx43 and Plakoglobin on the ICD of cKO mice, control mice and cKO mice injected with AAV9-cTnT-Ig Igk vector.**

(A) a. Colocalization of N-cadherin and plakoglobin on the ICD of cKO mice, control mice and cKO mice injected with AAV9-cTnT-Ig Igk. b. Colocalization of N-cadherin and plakoglobin on the ICD of cKO mice, control mice and cKO mice injected with AAV9-cTnT-Ig Igk.