Telocytes-derived extracellular vesicles alleviate aortic valve calcification by carrying miR-30b

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Abstract

Aims Calciﬁc aortic valve disease (CAVD) is frequent in the elderly. Telocytes (TCs) are implicated in intercellular communication by releasing extracellular vesicles (EVs). This study investigated the role of TC-EVs in aortic valve calcification.

Methods and results TCs were obtained and identiﬁed using enzymolysis method and ﬂow cytometry. EVs were isolated from TCs using differential high-speed centrifugation method and identiﬁed using transmission electron microscope, western blot, and qNano analysis. The mouse model of CAVD was established. The changes of aortic valve activity-related indicators were analysed by ultrasound, and the expressions of TC markers CD34 and vimentin in mouse valve tissues were detected using RT-qPCR and western blot. The model mice were injected with TC-derived EVs. The expressions of Runx2, osteocalcin, and caspase-3 were detected using RT-qPCR and western blot. The calcification model of valvular interstitial cells (VICs) was established. TC-EVs were co-cultured with calcified VICs, and calcium deposition was detected using alizarin red S staining. miR-30b expression in calcified valvular tissues and cells was detected after EV treatment. miR-30b expression in TCs was knocked down and then EVs were extracted and co-cultured with calcified VICs. The target of miR-30b was predicted through bioinformatics website and veriﬁed using dual-luciferase assay. The levels of Wnt/β-catenin pathway-related proteins were detected. ApoE−/− mice fed with a high-fat diet showed decreased aortic valve orifice area, increased aortic transvalvular pressure difference and velocity, reduced left ventricular ejection fraction, decreased CD34 and vimentin, and increased caspase-3, Runx2, and osteocalcin. The levels of apoptosis- and osteogenesis-related proteins were inhibited after EV treatment. TC-EVs reduced calcium deposition and osteogenic proteins in calcified VICs. EVs could be absorbed by VICs. miR-30b expression was promoted in calcified valvular tissues and cells after EV treatment. Knockdown of miR-30b weakened the inhibitory effects of TC-EVs on calcium deposition and osteogenic proteins. miR-30b targeted Runx2. EV treatment inhibited the Wnt/β-catenin pathway, and knockdown of miR-30b in TCs attenuated the inhibitory effect of TC-EVs on the Wnt/β-catenin pathway.

Conclusion TC-EVs played a protective role in aortic valve calcification via the miR-30b/Runx2/Wnt/β-catenin axis.

Keywords Aortic valve calcification; Telocytes; Extracellular vesicles; miR-30b; Runx2; Wnt/β-catenin; Apoptosis

Introduction

Calcific aortic valve disease (CAVD) is a common cardiovascular disease in the elderly population, which imposes heavy economic and social burdens.1 CAVD is primarily characterized by abnormal accumulation of calcium rich nodules on the surface of the aorta and/or in the annular region of the valve tip, resulting in sclerosis, movement restriction, and stenosis.2 There lacks of established drug therapies to prevent the progression of CAVD, and surgical valve replacement is the only available treatment option, which, however, is accompanied by insurmountable complications, economic burdens, and uncertainties in the long-term prognosis.3 The mechanism of promoting the occurrence and development of aortic valve calcification has not been fully identified, but a recent report has suggested that it is a complex process...
concerning molecular and cellular phenotypes, and bone formation. It is worth noting that the prevalence of CAVD is still rising due to the trend of aging population. Hence, a better understanding about the underlying events concerning the onset and progression of aortic valve calcification remains an urgent issue to be solved.

Telocytes (TCs) is a new type of interstitial cells, which are characterized by smaller cell body with long prolongations of uneven calibre. Cardiac interstitium contains typical TCs, which establish complicated spatial relationships with myocardial cells and cardiac stem cells, confirming the regulatory role of TCs in the three-dimensional organization of cardiac tissues. In pathological state, TCs can improve cardiac functions by promoting myocardial cell regeneration, enhancing angiogenesis and reducing cardiac fibrosis. A previous literature has pointed out that TCs are implicated in intercellular communication through direct gap junctions or extracellular vesicle (EV) release. Currently, TC-derived EVs have been accepted as biomarkers and transferring tools for drug, vaccine, and gene of cardiovascular diseases. New evidence has suggested that calcification-competent EVs from valvular interstitial cells (VICS) are regulators of calcification in diseased aortic valves and atherosclerotic plaques. More importantly, the intercellular communication mediated by EVs can also be achieved through the delivery of genetic information, such as microRNAs (miRs). miRs are highly conserved non protein-coding RNAs that maintain intracellular homeostasis through negative gene regulation. Deregulation of miRs is extensively reported in the progression and prognosis of CAVD. But the profiling and role of miR in TC-EVs in CAVD is less studied. Based on these findings, we investigated the role of TC-EVs in CAVD, along with its underlying miR mechanism, which shall provide impetus for the clinical management of CAVD.

**Methods**

**Ethics statement**

The study was approved by the Ethical Committee of The First Affiliated Hospital of Nanjing Medical University. All experimental procedures were performed based on the Ethical Guidelines for the study of experimental pain in conscious animals.

**Preparation of cardiac telocytes**

Female C57BL/6 mice aged 8 weeks and weighed 15–20 g [SYXX (Guangdong) 2019-0203, Ruijian Xingze Biological Medicine Co., Ltd, Dongguan, Guangdong, China] were treated with 3% pentobarbital sodium. Hearts were dissected under aseptic conditions and put into 50 mL centrifuge tubes containing cold phosphate buffer saline (PBS) added with 1% penicillin and streptomycin. Then the hearts were placed in the cell culture room immediately. Following rinsing with fresh PBS to remove the blood, the hearts were sliced into millimetre-sized pieces in a aseptic culture dish containing Dulbecco’s modified Eagle’s medium (DMEM)/F12 (12400-024, Gibco, Grand Island, NY, USA) added with 1% penicillin and streptomycin. These pieces were rinsed twice and resuspended in PBS. Next, the enzymatic digestion medium containing 1.5 mg/mL collagenase type 2 (V900892, Sigma-Aldrich, Merck KGaA, Darmstadt, Germany), DMEM/F12, and 1% penicillin and streptomycin was supplemented and the mixture was centrifuged at 180 rpm and 37°C for 40 min. The solution was filtered through a 41-μm nylon mesh (Millipore, Billerica, MA, USA), and the collected cell suspension was centrifuged at 300×g for 10 min. Thereafter, the cells were seeded in aseptic culture dishes containing 10 mL DMEM/F12, 10% EV-depleted fetal bovine serum (FBS) (System Biosciences, Mountain View, CA, USA) and 1% penicillin and streptomycin, followed by incubation in humidified air at 37°C for 1 h for fibroblast attachment. Unattached cells were transplanted into a fresh dish containing the above medium, and the medium was refreshed every 2 days.

**Immunofluorescence staining**

The phenotype of sorted cells was identified using immunofluorescence staining. In brief, the cells were seeded on cover glasses for 2 days, fixed with 4% para-formaldehyde for 10 min, and permeabilized with 0.5% Triton X-100 for 10 min. The cover glasses were cultured with the primary antibodies against α-smooth muscle actin (α-SMA), von Willebrand factor (vWF), and vimentin, and then incubated with the fluorescent-conjugated secondary antibodies. Images were captured under a fluorescence microscope (Carl Zeiss, Jena, Germany).

**Preparation of extracellular vesicles**

The supernatants of cultured TCs were collected on ice and centrifuged at 10 000×g for 30 min to remove cells and debris. Then, the supernatants were transferred to a fresh tube, filtered through a 0.22-μm membrane, and centrifuged at 120 000×g and 4°C for 2 h. The isolated EV pellet was washed once with aseptic PBS and resuspended in 200 μL PBS. In some samples, an Amicon ultra filter (Millipore) was used to reduce the volume from 50 to 1 mL to concentrate the supernatant, with a 100 000 molecular weight cutoff. Subsequently, EVs were isolated from the supernatants using the differential high-speed centrifugation method.

The quality of EVs was identified using qNano analysis (Izon Science, Burnside, New Zealand). The protein of EVs containing cold phosphate buffer saline (PBS) added with 1% penicillin and streptomycin. The supernatants of cultured TCs were collected on ice and centrifuged at 10 000×g for 30 min to remove cells and debris. Then, the supernatants were transferred to a fresh tube, filtered through a 0.22-μm membrane, and centrifuged at 120 000×g and 4°C for 2 h. The isolated EV pellet was washed once with aseptic PBS and resuspended in 200 μL PBS. In some samples, an Amicon ultra filter (Millipore) was used to reduce the volume from 50 to 1 mL to concentrate the supernatant, with a 100 000 molecular weight cutoff. Subsequently, EVs were isolated from the supernatants using the differential high-speed centrifugation method.

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was quantified using the bicinchoninic acid (BCA) assay kit (Thermo Fisher Scientific, Rockford, IL, USA). EVs were observed using transmission electron microscopy (TEM) (CM120, Philips, Andover, MA, USA). The expression levels of CD63, CD81, and calnexin were detected using western blot. The supernatant supplemented with GW4869 acted as the negative control (NC) of EVs.

Extracellular vesicles used in this study were assigned into four groups: NC group (the supernatant of TCs supplemented with GW4869), TC-EVs group, in-miR-30b/TC-EVs group (TCs were transfected with miR-30b inhibitor and then EVs were extracted), and in-NC/TC-EVs group (TCs were transfected with inhibitor NC and then EVs were extracted). For the cell transfection, TCs in the logarithmic growth phase were seeded into 6-well plates (1.0 × 10^5 cells per well), and then transfected with miR-30b inhibitor or inhibitor NC using Lipofectamine 2000 (Invitrogen Inc., Carlsbad, CA, USA) upon 60% confluence. The cells were harvested for subsequent experiments 48 h later. miR-30b inhibitor and inhibitor NC were synthesized and purified by GenePharma (Shanghai, China).

**Reverse transcription quantitative polymerase chain reaction**

The total RNA were extracted from tissues and cells using the Trizol reagent (Invitrogen) and reverse transcribed into cDNA using the Toyoba reverse transcription kit (Fermentas Inc., Hanover, MD, USA). RT-qPCR was performed in line with the instructions of the ABI PRISM 7900 system (Applied Biosystems, Carlsbad, CA, USA). Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) and U6 acted as the internal reference for genes and miR-30b, respectively. PCR mixture included the SYBR Green/Fluorescein QPCR Master Mix (2X) (Fermentas), cDNA, and the age-miR-30b LNA™ PCR primer set (UniRT) for miR-30b. The primer sequences are illustrated in Table 1. The relative gene expression was calculated based on the 2^-ΔΔCt method. The experiments were conducted three times independently.

**Western blot analysis**

The total protein was extracted from tissues and cells using radio-immunoprecipitation assay buffer (Shanghai Beyotime Biotechnology Co. Ltd., Shanghai, China) containing protease inhibitors (Roche, Basel, Switzerland). The concentration of protein was evaluated using the BCA assay kit (Beyotime). Then, the protein was separated on sodium dodecyl sulfate polyacrylamide gel electrophoresis and transferred onto polyvinylidene fluoride membranes. The membranes were blocked for 1 h and cultured with primary antibodies CD14 (1/10 000, ab221678, Abcam), vimentin (1/10 000, ab92547, Abcam), caspase-3 (0.7 μg/mL, ab184787, Abcam), Runx2 (1 μg/mL, ab23981, Abcam), osteocalcin (1/1000, ab184787, Abcam), Wnt3 (1/10 000, ab172612, Abcam), and β-catenin (0.5 μg/mL, ab16051, Abcam) at 4°C overnight. Afterwards, the membranes were cultured with the secondary antibody immunoglobulin G (IgG) (1/2000, ab205718, Abcam). The membranes were developed and analysed using Image Lab™ software (BioRad Laboratories, Hercules, CA, USA).

**Establishment of a mouse model of calcific aortic valve disease**

Apoe^-/- mouse model of atherosclerosis induced by a high-fat diet was used as the reference to establish the CAVD model. Briefly, Apoe^-/- mice aged 7–8 weeks were fed with a high-fat diet containing 1.25% cholesterol and 7.5% saturated fat (CAVD group). Wild type C57BL/6 mice fed with a normal diet and a high-fat diet were set as the control (WT group). After 10 months of feeding, the mice were subjected to echocardiography examination. Additionally, after the mice were fed with 10 months of high-fat diet for calcification induction, they were injected with 50 μg TC-EVs via tail vein (CAVD + TC-EVs group), and the mice injected with equal amounts of cell supernatants containing GW4869 were set as the controls (CAVD + NC group). After 7 days of normal feeding, the mice were euthanized by intraperitoneal injection of pentobarbital sodium ≥100 mg/kg. In this study, six C57BL/6 mice and 24 Apoe^-/- mice were used for the establishment of CAVD model, with six mice in each group. After euthanasia, the valve tissues were collected for RT-qPCR and western blot analysis.

**Table 1 Primer sequence for RT-qPCR**

| Gene    | Primer sequence |
|---------|-----------------|
| miR-29a | F: 5'-ACCAAGTTTCACTTGGTAAAC-3' |
|         | R: 5'-TCCAAGTCGCTGAAATGAAC-3' |
| CD34    | F: 5'-TCTGGTATGCCCCATACCTCC-3' |
|         | R: 5'-TATGGGACGTAGGGAGCCACT-3' |
| Vimentin| F: 5'-CAGATGCGTGAGATGGAAGA-3' |
|         | R: 5'-TCCAGCGGTCTCTAGTGA-3' |
| Caspase-3| F: 5'-AGAGCTGGGGGCTCAAGTGA-3' |
|         | R: 5'-AGAACCACTGGCACTATGGA-3' |
| Runx2   | F: 5'-ATGAGACGCCCACACTCCTCG-3' |
|         | R: 5'-CACCTCTAGACGGGGGCGTAAAG-3' |
| Osteocalcin| F: 5'-GGGACCCAAAGGGGCTCAT-3' |
|         | R: 5'-GAGTCCTCCACGACATACAA-3' |
| GAPDH   | F: 5'-GGCTCGGCGACGACATATAC-3' |
|         | R: 5'-CAATGGGAACGCTCGACAGA-3' |

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Echocardiography

Echocardiography was performed using the cardiac ultrasound instrument (Vivid E95, General Electric Company, Schenectady, NY, USA), with the transducer frequency of 2.0–5.0 MHz. The mice were anaesthetised with ether. The chest and upper abdomen of mice were depilated, coated with conductive adhesive, and fixed on the test table. The left, right, and apex of mouse sternum were scanned. The valve orifice area, aortic transvalvular pressure difference and velocity, and left ventricular ejection fraction (LVEF) were recorded. Vivid paediatric heart ultrasound probe M5Sc-D was used. Echocardiography parameters were evaluated by radiologist independently in a blind manner.

Calcification induction of valvular interstitial cells

The VICs at passage 3–7 (Beijing Qbioscience Biotechnology Co., Ltd, Beijing, China) were detached with trypsin to prepare cell suspension. Cells were seeded in 6-well plates at 1 x 10^5/mL. The interstitial cells in the blank group were cultured in the standard medium (DMEM + 10% FBS), and interstitial cells in the CAVD group were incubated in the calcification medium (0.1% FBS, 50 ng/mL BMP-2, 100 nM dexamethasone, 50 μg/mL ascorbic acid and 5 mM β-glycerophosphate). The medium was refreshed every 2 days for totally 7 days of culture. Additionally, the CAVD + NC group and CAVD + TC-EVs group were set, and NC was the cell supernatant supplemented with GW4869.

Valvular interstitial cells used in this study were assigned into six groups: blank group, CAVD group, CAVD + TC-EVs group (CAVD VICs were treated with 50 μg TC-Evs), CAVD + NC group (CAVD VICs were treated with equal amounts of NC), CAVD + in-miR-30b/TC-EVs group (CAVD VICs were treated with equal amounts of in-miR-30b/TC-EVs), and CAVD + in-NC/TC-EVs group (CAVD VICs were treated with equal amounts of in-NC/TC-EVs). After 24-h incubation, the cells were harvested for subsequent experiments.

Von Kossa staining

The valve tissue sections were immersed in 2% silver nitrate for 20 min under good sunlight or ultraviolet lamp, washed with distilled water for 5 min, treated with 5% sodium thiosulfate solution for 2 min, and counterstained with 0.1% nuclear solid red solution for 1 min. Following washing, the calcified tissues were black under the light microscope.

Alizarin red S staining

After staining with alizarin red S solution (40 mM; Sigma-Aldrich), the calcified VICs were placed for 30 min. The cells were washed and then observed under the light microscope.

Dual-luciferase reporter gene assay

The 3’UTR sequences of Runx2 containing miR-30b binding site were synthesized. Wild-type (WT) plasmids (Runx2-WT) and mutant-type plasmid (Runx2-MUT) were constructed. Then the constructed vectors were mixed with mimic-NC or miR-30b mimic and transfected into 293T cells (American Type Culture Collection, Manassas, Virginia, USA). After 48 h of transfection, luciferase activities were detected using the dual-luciferase reporter Gene assay kit (BioVision, SanFrancisco, CA, USA) on the Glomax20/20 luminometer (Promega, WI, USA).

Statistical analysis

SPSS 21.0 software (IBM Corp., Armonk, NY, USA) was utilized for data analysis. Data were expressed as mean ± standard deviation. Comparisons of data between two groups were analysed using the t-test. One-way analysis of variance (ANOVA) was employed for comparisons among multi-groups, followed by Tukey’s multiple comparison test. The P value was obtained from a two-tailed test, and a value of P < 0.05 meant a statistical difference.

Results

Culture of telocytes and extraction of extracellular vesicles

Single cell suspension was obtained using enzymatic hydrolysis, and the cells were seeded in the culture flask after adherence and gradient centrifugations. After 96 h of culture, a large amount of cell debris could be observed under the light microscope. After 7 days, the cell monolayer showed clonal phenomenon with the size of 10–30 μM. After 14 days, the cells reached 80% confluence and cell populations arranged in the form of paving stones; podoms and podomers appeared alternately with uneven length (Figure 1A). The cardiac cells expressing CD34 and CD117 were sorted using flow cytometry (Figure 1B). The level of vimentin was detected using immunofluorescence staining. Telopodes and podomers also showed vimentin-positive, and the nucleus was blue after DAPI staining (Figure 1C).

Then, EVs were isolated from TCs using high-speed centrifugation and identified using TEM, western blot analysis, and qNano analysis. The isolated EVs were cup-shaped or round vesicles (Figure 1D). EV markers CD63 and CD117 were sorted using flow cytometry (Figure 1B). The level of vimentin was detected using immunofluorescence staining. Telopodes and podomers also showed vimentin-positive, and the nucleus was blue after DAPI staining (Figure 1C).

The diameter of most EVs ranged from 50 to 150 nm (Figure 1F).
Telocyte-extracellular vesicles prevented aortic valve calcification and apoptosis of valvular interstitial cells

ApoE−/− mice were fed with a high-fat diet for the establishment of CAVD model. After 10 months of high-fat diet feeding for calcification induction, echocardiographic analysis was performed, and the results showed that the aortic valve orifice area was decreased, the aortic transvalvular pressure difference and velocity were increased, and LEVF was reduced significantly (Figure 2A), indicating that the mouse model of CAVD was successfully established. Moreover, the expressions of TC markers CD34 and vimentin in mouse valve tissues were detected using RT-qPCR and western blot, and the results showed after 10 months of calcification induction in ApoE−/− mice. The expressions of CD34 and vimentin in mouse valve tissues were significantly decreased (Figure 2B,C), indicating that TCs played a positive role in myocardial injury.

Telocyte-extracellular vesicles can reduce myocardial fibrosis, improve cardiac function, and increase angiogenesis. Hence, we hypothesized that TC-EVs had an effect on aortic valve calcification and apoptosis of valvular interstitial cells.
valve calcification. To confirm this hypothesis, model mice were treated with TC-EVs, and then the expression levels of Runx2, osteocalcin, and caspase-3 were detected using RT-qPCR and western blot. The results revealed that the expressions of Runx2, osteocalcin, and caspase-3 in calcified valves of model mice were notably higher than those in EV-treated mice (Figure 2D,E), indicating that TC-EVs treatment reduced the calcification of aortic valve and alleviated the apoptosis of VICs.

Telocyte-extracellular vesicles prevented valves from transforming into contractile and osteogenic phenotypes

To further determine the effect of TC-EVs on valve calcification, we conducted in vitro experiments. A calcification model of VICs was established in vitro, and the appearance and phenotype changes of calcified VICs were observed. VICs were cultured and identified. After 12 h of culture,
the primary VICs began to adhere to the wall and formed a continuous monolayer in about 1 week (Figure 3A). VICs were positive for α-SMA and vimentin and negative for vWF, as immunofluorescence staining indicated (Figure 3B). The aforementioned markers were quantified using flow cytometry. Vimentin was highly expressed in VICs (99%), and α-SMA-positive cells accounted for about 50% of all cells (Figure 3C) (both $P < 0.01$). Subsequently, calcification induction was performed. The growth morphology of the cells induced by calcification was similar to that of control cells, with spindle-shape, clear boundary, and radial or swirling growth arrangement (Figure 3D). TC-EVs were cultured with calcified valve cells, and then the calcium deposition was detected. The results showed that EV treatment reduced the formation of calcium deposition significantly in calcified valve cells (Figure 3E). The levels of Runx2 and osteocalcin were increased in calcified valve cells, and EV treatment notably decreased the levels of osteogenic-related proteins in calcified valve cells (Figure 3F,G) (all $P < 0.01$).

Figure 3 TC-EVs prevented aortic valves from transforming into contractile and osteogenic phenotype. (A) Appearance of calcified valve interstitial cells was observed under the light microscope; (B) levels of α-SMA, vimentin, and vWF in interstitial cells were observed; (C) quantitative analysis of vimentin and α-SMA-positive cells in valve interstitial cells was conducted using flow cytometry; (D) cell growth morphology before and after calcification induction was observed; (E) formation of calcium deposition was detected using Alizarin red S staining; (F, G) levels of Runx2 and osteocalcin were detected using RT-qPCR and western blot analysis. The cell experiment was repeated three times independently. Data are presented as mean ± standard deviation. Data in panel E/F/G were analysed using one-way ANOVA, followed by Tukey’s multiple comparison test. **$P < 0.01$. 

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Telocyte-extracellular vesicles were the communication media of miR-30b during aortic valve calcification

It has been reported that miR-30b has a certain effect on aortic valve calcification. We speculated that TC-EVs protected aortic valve calcification by releasing miR-30b. PKH26-labelled EVs were cultured with VICs for 48 h, and the cytoplasm of cells incubated with PKH26-labelled EVs showed red fluorescence (Figure 4A), suggesting that TC-EVs could be absorbed by VICs. Then, miR-30b expression in the supernatants treated with TC-EVs or GW4869 was detected using RT-qPCR, along with miR-30b expression in calcified valvular tissues and VICs after EV treatment. The results showed that miR-30b mainly existed in EVs, and miR-30b expression in calcified valvular tissues and VICs was notably increased after EV treatment (Figure 4B) (all \( P < 0.01 \)). It was indicated that TC-EVs were the communication media of miR-30b during aortic valve calcification.

Inhibition of miR-30b attenuated the protective effect of telocyte-extracellular vesicles on aortic valve calcification

We knocked down the miR-30b expression in TCs and then extracted the EVs to confirm the protective effects of EVs on aortic valve calcification by releasing miR-30b. miR-30b expression in EVs extracted from TCs with knockdown of miR-30b was significantly reduced (Figure 5A). EVs with low-expressing miR-30b were cultured with VICs, and the results showed that inhibition of miR-30b could weaken the inhibitory effects of TC-EVs on the formation of calcium deposition, and the levels of Runx2 and osteocalcin (Figure 5B–D).

Telocyte-extracellular vesicles transferred miR-30b to inhibit Runx2 and inactivate the Wnt/β-catenin pathway

From the aforementioned results, we knew that TC-EVs played a protective role in valve calcification by transferring miR-30b, but the downstream mechanism of miR-30b remained unclear. The bioinformatics website (http://starbase.sysu.edu.cn/agoClipRNA.php?source) showed that there were several target genes of miR-30b, including Runx2. We found in the aforementioned results that miR-30b negatively regulated Runx2, so we hypothesized that there was a target relationship between miR-30b and Runx2. The target relationship between miR-30b and Runx2 was verified using dual-luciferase reporter gene assay (Figure 6A). Inactivation of the Wnt/β-Catenin pathway can inhibit valve calcification, and Runx-2 acts coordinately with components of the Wnt/β-catenin signalling pathway to enhance gene expression of both osteogenic bone markers and mineralization. Hence, the levels of Wnt/β-catenin pathway-related proteins in tissues and cells were detected. The Wnt/β-catenin pathway was activated in CAVD group compared with the blank group, and EV treatment inhibited the Wnt/β-catenin pathway; knockdown of miR-30b attenuated the inhibitory effect of TC-EVs on the Wnt/β-catenin pathway (Figure 6B).

Discussion

Telocyte-extracellular vesicles participate in the progression of myocardial infarction, systemic sclerosis, and heart failure by delivering miRs, which can become a breakthrough in the management of CAVD. Intriguingly, miRs are critical factors in regulating gene expression in various diseases. In this study, we found that TC-EVs were the communication media of miR-30b and could inhibit Runx2 and inactivate the Wnt/β-catenin pathway, which might be a potential therapeutic target for aortic valve calcification.
in the initiation and prognosis of CAVD. Our results elucidated that TC-EVs ameliorated aortic valve calcification via the miR-30b/Runx2/Wnt/β-catenin axis.

In the human heart, TCs widely exist in epicardium, myocardial interstitium, and aortic valves, where they contribute to the modulation of cardiac homeostasis and regeneration. Through the release of EVs or atypical junctions, TCs form a cardiac network and manipulate the long-distance heterologous communication in normal or diseased hearts. A previous study has reported that TC-EVs transfer macromolecular signals to adjacent cells to stimulate neovascularization in infarcted myocardium. In the present study, we extracted and identified TC-EVs, and then established the mouse model of CAVD. Our results
found that 10 months of calcification induction reduced the expressions of TC markers CD34 and vimentin in mouse valve tissues, indicating that TCs played a protective role against myocardial injury. The decrease of TCs in adult hearts may be one of the reasons for the limited cardiac regeneration ability in the elderly. TC-shuttled exosomes attenuate cardiac fibrosis and reduce collagen deposition. Accordingly, we hypothesized that TC-derived EVs also exerted some
effects on aortic valve calcification. Hence, model mice were treated with TC-EVs, and the results found that TC-EVs decreased the expressions of osteoblast specific transcription factor Runx2, the most abundant non-collagen in bone matrix osteocalcin, and the key gene of apoptosis in calcified valves of mice caspase-3. Briefly, the results demonstrated that TC-EVs could reduce the degree of aortic valve calcification and the apoptosis of VICs.

Valvular interstitial cells are the most common cells in aortic valves, which regulate normal valve functions and repair diseased valves. In addition, the in vitro experiments were conducted to further determine the effect of TC-EVs on valve calcification. TC-EVs were co-cultured with calcified VICs, and then the phenotypic change of VICs was observed. Valve calcification generally has a pathophysiological process involving the osteogenic differentiation of VICs, lipoprotein deposition, and inflammation. The osteogenic differentiation of VICs is ultimately accounted for calcium deposition and osteogenic nodule formation. Runx2, a transcription factor, is indispensable for the osteoblast differentiation and chondrocyte maturation. Osteocalcin is a type of osteoblast specific secreted protein, which serves as a hormone by triggering insulin generation and promoting energy consumption in target organs. The levels of osteogenic proteins, such as Runx2 and osteocalcin, are highly expressed in calcific aortic valves of human or mice. EV treatment notably decreased the levels of Runx2 and osteocalcin in calcified valve cells. Consistently, Liu et al. have demonstrated that multiple myeloma-derived EVs can lead to a reduction of Runx2 and osteocalcin in bone marrow stromal cells.

Extracellular vesicles derived from TCs can transfer macromolecular signals such as miRs to neighbouring cells and change their transcriptional activity. It has been reported that miRs have great potentials as a biomarker for the diagnosis and treatment of CAVD. miR-30b acts as an osteogenic inhibitor and exerts effects on aortic valvular calcification and apoptosis in CAVD by modulating Runx2 and caspase-3. Vishal Nigam et al. have indicated that overexpression of miR-30b can reduce the levels of calcification-related genes in aortic VICs. However, whether TC-EVs play a protective role in CAVD via transferring miR-30b remains unclear. The results of our experiments indicated that TC-EVs could be absorbed by VICs, and miR-30b expression in calcified valvular tissues was notably increased after EV treatment. Furthermore, inhibition of miR-30b could attenuate the protective effect of TC-EVs on aortic valve calcification, indicating that TC-EVs were the communication media of miR-30b in aortic valve calcification. Subsequently, we focused on the downstream mechanism of TC-EVs in CAVD via transferring miR-30b. Bioinformatics website predicted that there were multiple target genes of miR-30b, including Runx2. It has been reported that simvastatin can reduce the calcium deposition in aortic VICs in vitro by decreasing LPS-induced Runx2 expression, suggesting the involvement of Runx2 in aortic valve calcification. The target relationship between miR-30b and Runx2 was verified using dual-luciferase reporter gene assay in this study. Consistently, a previous literature has also exhibited that miR-30 family members can suppress osteoblast differentiation via inhibiting Runx2 in MC3T3-E1 cells. Wnt/β-catenin pathway is implicated in the differentiation of VICs into osteoblast cells, eventually contributing to aortic valve calcification. Cai et al. have demonstrated that Wnt/β-catenin pathway can enhance the osteogenic differentiation and calcification of vascular smooth muscle cells by regulating Runx2. Our results showed that EV treatment suppressed the Wnt/β-catenin pathway, and EVs extracted form miR-30b-silencing TCs attenuated the inhibiting effects on the Wnt/β-catenin pathway. Overall, TC-EVs could transfer miR-30b to inhibit Runx2 level and the Wnt/β-catenin pathway in calcified valve tissues.

To conclude, we are the first to find that TC-EVs alleviate valve calcification via the delivery of miR-30b and the Runx2/Wnt/β-catenin axis. This fundamental information may suggest potential therapy for CAVD in the clinical trial. In the future, we shall carry out more prospective trials on the feasibility and safety of TC-EVs in the treatment of CAVD, so as to refine our clinical guidance.

Conflict of interest

The authors declared that they have no competing interests.

Author contributions

R.Y. is the guarantor of integrity of the entire study; R.Y. contributed to the study concepts, study design, definition of intellectual content, literature research, manuscript preparation, and manuscript editing and review; Y.H.T. contributed to the clinical studies; R.Y. and X.W.C. contributed to the experimental studies and data acquisition; Y.Y. contributed to the data analysis and statistical analysis. All authors read and approved the final manuscript.

Data availability statement

All the data generated or analysed during this study are included in this published article.
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