YAP Circular RNA, circYap, Attenuates Cardiac Fibrosis via Binding with Tropomyosin-4 and Gamma-Actin Decreasing Actin Polymerization

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Cardiac fibrosis is a common pathological feature of cardiac hypertrophy. This study was designed to investigate a novel function of Yes-associated protein (YAP) circular RNA, circYap, in modulating cardiac fibrosis and the underlying mechanisms. By circular RNA sequencing, we found that three out of fifteen reported circYap isoforms were expressed in nine human heart tissues, with the isoform hsa_circ_0002320 being the highest. The levels of this isoform in the hearts of patients with cardiac hypertrophy were found to be significantly decreased. In the pressure overload mouse model, the levels of circYap were reduced in mouse hearts with transverse aortic constriction (TAC). Upon circYap plasmid injection, the cardiac fibrosis was attenuated, and the heart function was improved along with the elevation of cardiac circYap levels in TAC mice. Tropomyosin-4 (TMP4) and gamma-actin (ACTG) were identified to bind with circYap in cardiac cells and mouse heart tissues. Such bindings led to an increased TPM4 interaction with ACTG, resulting in the inhibition of actin polymerization and the following fibrosis. Collectively, our study uncovered a novel molecule that could regulate cardiac remodeling during cardiac fibrosis and implicated a new function of circular RNA. This process may be targeted for future cardio-therapy.

INTRODUCTION
Cardiac fibrosis is one of the most common contributions to cardiac dysfunction, due to pathological myocardial remodeling. This process is associated with excessive matrix (i.e., collagens) deposition and cardiac fibroblast activation, leading to a variety of cardiac remodeling and progressive cardiac dysfunction.1 Upon cardiac fibrosis, the compliance of heart tissues is decreased, and the progression of heart failure is accelerated. Cardiac fibroblasts play central roles in maintaining physiological heart functions as well as pathogenic cardiac remodeling during myocardial infarction and heart failure.2,3 Following acute cardiac injury, fibroblasts are activated. These activated fibroblasts are involved in the process of inflammation, fibrosis, and scar formation consequently. In the past prevailing theory, the activated form of fibroblasts, also known as myofibroblasts, was regarded as the key effector in the pathogenesis of cardiac fibrosis via secreting collagens and other extracellular matrix (ECM) molecules. However, recent studies revealed protective effects of myofibroblasts upon pressure overload (PO).4 There appear to be some factors regulating myofibroblast activities associated with synthesis of fibrosis markers and cardiac remodeling. Understanding the mechanism is critical for the development of safe and effective treatment targeting cardiac fibrosis.

Yes-associated protein (YAP) is an essential effector in the Hippo signaling pathway.5–8 YAP has been reported to mediate crucial pathways in heart development, heart regeneration, cardiac hypertrophy, and myocardial infarction.9–12 Activation of cardiomyocyte YAP could improve cardiac function and survival after cardiac injury.13 In addition to functioning on cardiomyocytes, the Hippo-YAP pathway has been found to affect other types of cells in the heart, including fibroblasts, vascular cells, and immune cells. The human YAP gene regulates a variety of cell activities and can generate many exons.1,14 With alternative splicing, many isoforms can be produced. Back-splicing of these exons can potentially synthesize 15 YAP circular RNA isoforms.15 It thus appears that more circular RNA isoforms than mRNA isoforms can be synthesized by the same gene. In our previous study, we found that one of the YAP circular isoforms (hsa_circ_0002320), which is generated from exons 5 and 6 of YAP pre-mRNA, played a crucial role in cell proliferation and survival.16 Interestingly, our circular RNA sequencing showed that this isoform was the most abundant YAP circular RNA (circYap) expressed in human heart tissue.
Recent studies have discovered the important roles of circular RNAs in regulating cardiovascular functions. Considering the important function of YAP, the parental gene of circYap, in cardiovascular diseases and our previous study on circYap, we were inspired to further explore the roles of circYap in regulating cardiac functions. In particular, our sequencing results indicated that investigating the potential effects of circYap hsa_circ_0002320 in cardiac remodeling and fibrosis is imperative. Our present study is designed to discover the novel role of this circular RNA in the development of cardiac fibrosis and the possible underlying mechanisms.

**RESULTS**

**Decreased Expression of circYap in Patients with Cardiac Diseases**

Based on the RNA sequencing results in nine human heart specimens (W.W.D. et al., unpublished data), three out of fifteen reported circYap isoforms were found to be expressed in these human hearts (Figure 1A). Among these three isoforms, hsa_circ_0002320 was the highest expressed circYap isoform in human hearts (Figure 1B). Subsequently, we measured the levels of mouse equivalent (89% homology) in different mouse organs and found that all organs analyzed expressed this circYap isoform.
with the heart being one of the organs expressing high levels (Figure 1C).

The YAP gene is known to play important role in maintaining heart functions. We examined whether hsa_circ_0002320 expressed by the YAP gene had any clinical implication. The levels of circYap were measured in the hearts of 104 patients with cardiac hypertrophy, including 25 patients with heart failure, 4 patients with aortic stenosis, 3 patients with mitral stenosis, 8 patients with hypertrophic cardiomyopathy, and 64 patients with tetralogy of Fallot. As the controls, 21 heart samples from donations were also subjected to RNA isolation for real-time PCR. We found that the levels of circYap were significantly reduced in the hearts of patients with cardiac hypertrophy compared to those in the donated normal hearts (Figure 1D).

Ectopic circYap Improved Heart Functions in the TAC Mice
To test the role of circYap in maintaining cardiac functions, we generated PO in mice by transverse aortic constriction (TAC). We found the expression of circYap was significantly decreased in the heart 8 weeks after surgery, while ectopic delivery of circYap expression plasmid increased circYap levels (Figure 2A). In the following heart functional study, we detected a significant increase in left ventricular end-systolic diameter (LVESD) and the left ventricular end-diastolic diameter (LVEDD) in the TAC mice, which were diminished by ectopic delivery of circYap expression plasmids that were generated by us previously (Figure 2B). We also detected a significant decrease in left ventricular ejection fraction (LVEF), left ventricular fractional shortening (LVFS), and contraction velocity (dp/dt) in the TAC hearts, and such reductions were prevented by the delivery of the circYap (Figure 2C). The heart weights were increased in the TAC mice but were maintained at normal levels in TAC mice with circYap plasmid delivery (Figure 2D). The M-mode pictures showed the typical increase in the left ventricular chamber of the TAC heart, and delivery of circYap reduced it to normal levels (Figure 2E).

Ectopic circYap Prevented Heart Fibrosis
Fluorescence in situ hybridization (FISH) confirmed successful delivery of circYap into mouse heart tissues (Figure 3A; Figure S1A). In addition, co-localization of circYap with cardiomyocytes and cardiac fibroblasts upon circYap plasmid delivery was observed by FISH and immunofluorescence (IF) staining (Figure S1B). H&E staining and Sirius red staining demonstrated that the hypertrophy and fibrosis in the TAC mouse hearts were prevented by the delivery of circYap plasmids (Figure 3B). Since cardiac hypertrophy and remodeling are associated with fibrosis, we stained the heart tissues with Masson trichrome and Sirius red to visualize the collagen deposition in the...
Quantitative analysis revealed that the collagen levels were significantly increased in the hearts of TAC mice, while circYap injection retained the collagen levels in TAC mice (Figures 3D and 3E). To examine the clinical relevance of these results, we measured levels of collagen-I and collagen-III in the patient heart specimens. The assays showed that the levels of collagen-I and collagen-III were significantly higher in the samples from the patients with cardiac hypertrophy compared to the hearts without cardiovascular disease (Figures 3F and 3G). In addition, the correlation of circYap and collagens in human heart tissue samples was analyzed. Moderate but significant negative correlations were observed between circYap and collagens (Figures S1D and S1E).

Furthermore, we analyzed collagen expression in mouse heart tissues by using real-time PCR. Collagen-I and collagen-III were significantly increased after TAC and decreased by circYap delivery (Figures 4A and 4B). The levels of both collagen-I (Figure 4A) and collagen-III (Figure 4B) were inversely correlated with circYap expression. We also measured the expression of other fibrosis markers, including transforming growth factor-β1 (TGF-β1) (Figure 4C), nerve growth factor (NGF) (Figure 4D), and connective tissue growth factor (CTGF) (Figure S1F). Expectedly, levels of all of these fibrosis markers increased significantly in the TAC mouse hearts. Ectopic delivery of circYap prevented such effects of TAC. The levels of these fibrosis markers showed inverse correlation with circYap levels. However, we did not find significant change.
of tumor necrosis factor-α (TNF-α) levels, and therefore the levels of TNF-α were not correlated with circYap expression (Figure S1G). These results demonstrated that circYap was involved in maintaining heart function via inhibiting cardiac fibrosis during PO.

To examine how circYap affected heart functions, we ectopically expressed circYap in mouse cardiac fibroblasts (MCFs) and cardiomyocytes (HL-1 and AC16). After confirming the increased expression of circYap (Figures S2A and S2B), we measured mRNA expression of collagen-III and found the collagen-III mRNA expression was significantly downregulated in the cells with circYap overexpression (Figure 4E). Since fibroblasts play the most important roles in mediating the development of cardiac fibrosis, we treated MCFs with recombinant TGF-β1 to mimic the induction of fibrosis. The results showed that TGF-β1 treatment significantly boosted the expression of collagen-I and collagen-III (Figure 4F), while overexpression of circYap could successfully block the elevation of these fibrosis markers (Figure 4G). Overexpression of circYap also inhibited the total collagen levels in the culture media and altered the secretion of some cytokines upon TGF-β1 treatment (Figures S2C and S2D). In addition, overexpression of circYap significantly increased the survival rates of cardiac cells (Figure S2E) and decreased the migratory
ability of cardiac fibroblasts (Figure S2F). We also noticed the cell morphologies were obviously changed in the cells after stable transfection with circYap under both normal (with fetal bovine serum [FBS]) and stress (without FBS) condition. Upon circYap overexpression, the cardiac fibroblasts became flat and spread out, while the cardiomyocytes like HL-1 and AC16 cell were shrunken and elongated (Figure S3).

### Interaction of circYap with TPM4 and ACTG

To uncover the molecules that mediated the function of circYap in cardiac fibrosis, we performed RNA pull-down assay by using the circYap probe in MCF and HL-1 cells. Twelve proteins were identified to be pulled down by circYap probe, among which gamma-actin (ACTG) and tropomyosin-4 (TPM4) were ranked as the top two highest counts with no change in expression levels (Figure 5A; Table S3; Figure S4A). To confirm the mass spectrometry results and test the specificity of the interaction, we generated a circYap construct without the 5' intron (no-intron), the essential component in back-splicing. Transfection with circYap increased circYap levels in both cell types significantly, but transfection with the no-intron construct had little effect (Figure 5B).

![Figure 5. The Binding of circYap with ACTG and TPM4 Proteins](image)

(A) Mass spectrometry assay showing the proteins pulled down by circYap probe. (B) Expression of circYap in MCF and HL-1 cells transfected with circYap or the plasmids lacking an intron for circularization (no-intron). n = 3. **p < 0.01. (C) The binding of circYap with ACTG and TPM4 in MCF and HL-1 cells with or without circYap and its linear precursor (no-intron) overexpression. n = 3. **p < 0.01. (D) Antibodies against ACTG and TPM4 could precipitate circYap in mouse heart tissues. n = 3. **p < 0.01. (E) The binding of circYap or YAP mRNA with TPM4 and ACTG in mouse heart tissues with or without RNase R treatment. n = 4. **p < 0.01. (F) The binding of circYap with ACTG and TPM4 in pressure-overload heart tissues with or without circYap plasmid injection. n = 4. **p < 0.01. (G) The circYap probe pulled down ACTG and TPM4 proteins in MCF cells. (H) The binding of ACTG and TPM4 proteins in MCF cells upon circYap overexpression with RNase A or RNase R treatment. RNase A and RNase R were added after immunoprecipitation. n = 3. (I) The binding of ACTG and TPM4 proteins in MCF cells upon circYap overexpression with RNase A or RNase R treatment. RNase A and RNase R were added before immunoprecipitation. n = 3.
with circYap and the no-intron constructs were subjected to immunoprecipitation assays. Antibodies against ACTG and TPM4 proteins could precipitate circYap but not the product of the no-intron construct (Figure 5C). In mouse heart tissues, we also confirmed that antibodies against ACTG and TPM4 could precipitate endogenous circYap (Figure 5D).

Since we have reported that circYap has potential binding ability with YAP mRNA, we used RNase R to treat the heart tissue lysate to exclude the possibility that circYap bound with these proteins via YAP mRNA. We found that after depletion of the potential binding of YAP mRNA with either ACTG or TPM4 by RNase R treatment, circYap could still bind to these two proteins (Figure S5E). This indicated that circYap directly bound to ACTG and TPM4 but not bound with these proteins via YAP mRNA. Moreover, we found that the binding of circYap with ACTG and TPM4 was significantly decreased in mouse heart tissues upon TAC but maintained at a higher level by restoration of circYap levels with plasmid injection (Figure 5F).

Due to the interaction of ACTG with TPM4 in a dynamic way under physiological conditions, we examined the change of interaction of ACTG and TPM4 upon circYap overexpression. In the circYap-transfected cells, the circYap probe could pull down ACTG and TPM4 (Figure 5G; Figure S4B), which indicated that circYap enhanced the interaction of ACTG and TPM4.

To determine whether the interaction of ACTG and TPM4 were bridged by circYap or a complex of circYap-ACTG-TPM4 was formed, the reaction mixture was treated with RNase A or RNase R before or after the immunoprecipitation. We found that neither RNase A nor RNase R treatment after binding reaction could reduce the binding ability of ACTG with TPM4 (Figure 5H). RNase A treatment before the binding reaction showed decreased effect of circYap on the formation of the complex (Figure 5I). These results demonstrated that circYap directly bound to both ACTG and TPM4 and facilitated the interaction of these two proteins by forming a complex.

Identifying the Binding Sites of circYap with TPM4 and ACTG

By using a computational approach, the contact sites of circYap with ACTG and TPM4 were putatively mapped (Figure S4C). The best predicted secondary structure of circYap was analyzed for its thermodynamic properties. The structure of the ACTG protein used in the docking procedure was derived from Protein Data Bank (PDB) entry PDB: 5JLH. The molecular simulation result supported that circYap could perfectly dock ACTG, while the 2- and 3-dimensional sequence of the circYap showed different structures (Figure S4D). The docking analysis predicted a minimal binding region of circYap for ACTG, as shown in Figure 6A. Due to unavailable crystallographic data on TPM4, we used a homology modeling approach for the generation of the three-dimensional target structure. The molecular simulation also supported that circYap could perfectly dock TPM4, and this predicted a minimal binding region of circYap for TPM4, as shown in Figure 6B. Graphical representation of the three-dimensional structure of circYap-ACTG (PDB: 5JLH)-TPM4 complex was shown in Figure 6C and S5A–S5C. To confirm these binding activities, we performed site-directed mutagenesis to generate mutations of the binding sites in circYap or these two proteins, respectively. First, we transfected the cells with circYap or protein binding site mutated circYap constructs (circYapmuACTG and circYapmuTPM4). We confirmed that mutation of either or both protein binding sites for circYap significantly decreased the ACTG or TPM4 antibody precipitating circYap (Figure 6D). On the other hand, circYapmuACTG and circYapmuTPM4 could not pull down ACTG or TPM4 due to the mutation of their binding sites (Figure 6E). Moreover, transfection with circYapmuACTG and circYapmuTPM4 constructs remarkably reduced ACTG antibody precipitating TPM4 and vice versa (Figure 6F).

Second, we synthesized blocking oligos, which were reverse complement to the ACTG/TPM4 protein binding sites in circYap, to block the binding of circYap with ACTG or TPM4. It showed that ACTG and TPM4 were not able to precipitate circYap in the presence of the blocking oligos (Figure 6G). In accordance, transfection with the blocking oligos inhibited the capability of circYap to pull down ACTG and TPM4 proteins (Figure 6H). In addition, the blocking oligos also decreased antibody against ACTG precipitating TPM4 and vice versa (Figure 6I).

Last but not least, to further confirm the binding sites, we designed His-tagged ACTG or TPM4 with circYap binding sites mutation (ACTGmut and TPMmut) (Figure S5D). By using His-tag, the potential influence of site mutation on antibody or primer recognition of ACTG and TPM4 was avoided. We found that circYap could not pull down ACTG or TMP4 with the mutation of circYap binding sites (Figure 6). In addition, ACTG or TPM4 with or without mutation were precipitated via His-tag by using Ni-NTA agarose beads. The mutated ACTG or TPM4 precipitated significantly less circYap compared to those without mutation (Figure S5E). These results confirmed that the binding of circYap and ACTG or TPM4 was abrogated by ACTGmut or TPMmut. In mouse heart tissues, the binding of ACTG and TPM4 were significantly suppressed in TAC mice that expressed decreased levels of circYap, while the interaction of ACTG and TPM4 in TAC mice with the ectopic delivery of circYap was maintained at similar levels to the sham mice (Figure 6K). These results indicated these protein binding sites were crucial for the binding of circYap with ACTG and TPM4, and circYap was involved in the interaction between ACTG and TPM4.

circYap Decreased Actin Polymerization

Actin polymerization plays important roles in regulating cell activities. Tropomyosin is an essential regulator of actin polymerization. Under physiological conditions, TPM proteins bind to actin to prevent the assembly of new actin filaments and actin polymerization. Our results showed that overexpression of circYap significantly inhibited the rate of actin polymerization (Figure 7A). In addition, we found that inhibition of ACTG by small interfering RNA (siRNA) could significantly reduce the migration rate of cardiac fibroblasts,
while silencing TPM4 by siRNA could remarkably accelerate the migration (Figure 7B; Figure S6A). This result confirmed the inhibitory effects of TPM4 on ACTG-mediated cell activities by polymerization. We further validated that interrupting the binding of circYap to ACTG and TPM4 by mutation on the binding sites significantly increased the motility of cardiac fibroblasts (Figure 7C; Figure S6B) and elevated the collagen-I and collagen-III levels (Figures 7D and 7E). In addition, blocking the binding of circYap to ACTG and TPM4 by either mutation, the binding sites or transfection with the blocking oligos decreased viability of MCF or AC16 cells (Figures S6C and S6D).

Finally, we tested whether circYap-mediated actin polymerization could occur in vivo. Heart lysates were prepared from sham and
TAC mice that showed decreased expression of circYap. An increase of actin polymerization was found in TAC mouse heart with the reduction of circYap expression. Ectopic delivery of circYap could inhibit the actin polymerization (Figure 7F). Therefore, we concluded that circYap facilitated the binding of TPM4 with ACTG, leading to the increased inhibitory effect of TPM4 on actin polymerization and the following anti-fibrotic effects (Figure 7G; Figure S6E).

**DISCUSSION**

Cardiac fibrosis is a worldwide health problem associated with nearly all etiologies of heart diseases. However, the molecular mechanisms underlying cardiac fibrosis remain unclear. Our study indicates that circYap is a critical regulator in binding to tropomyosin and actin, decreasing actin polymerization and cardiac fibrosis. Our study also suggests that we could target the process of actin polymerization to decrease cardiac fibrosis and improve heart functions by ectopic delivery of nanoparticle-conjugated circYap expression plasmids.

The promising potential of RNA-targeting therapeutics for human diseases has started to be unveiled in recent years. Circular RNAs have been reported to affect progression of heart disease and cancer. Circular RNAs are potential targets in RNA therapeutics. The circYap is one of the circRNAs that we found to have protective effects in both cancer and cardiovascular disease. Potentially, the human YAP gene can produce 15 isoforms of circular RNAs. However, the expression of circYap could be tissue specific, since our circular RNA sequencing result showed that only three isoforms were detected in human heart tissues. Therefore, we selected the one with the highest level in our study. We found that the level of this circYap isoform was significantly decreased in the hearts of patients with cardiac hypertrophy. In addition, our results revealed that the expression of circYap was negatively correlated with fibrosis factors including collagen-I and -III, TGF-β, CTGF, and NGF in the heart of TAC-induced PO mouse model. In the present study, we successfully restored the circYap levels in the hearts of mice with TAC by delivery of circYap plasmids, which...
were validated by real-time PCR and FISH. Notably, the impaired heart function and the cardiac fibrosis that resulted from PO were significantly improved after ectopic circYap plasmid delivery. These results provided a clue for the potential clinical application of circYap in RNA therapeutics. Regarding the cellular target of circYap in the heart, our present study showed that circYap affected the survival and cell morphology of both fibroblasts and cardiomyocytes in in vitro cell models. The fibroblasts are acknowledged to be the major effectors of cardiac fibrosis. It has been reported that fibroblasts can modulate cardiomyocyte survival via transduction of hypertrophic signal or secretion of pro-apoptotic, pro-inflammatory mediators and exosomes.34 Our results from FISH in mouse heart tissue indicated circYap is expressed in both cardiomyocytes and cardiac fibroblasts. However, the function of circYap on myocytes and fibroblasts may be slightly different due to the different roles of these two types of cells in cardiac remodeling. The roles of circYap on specific cell types warranted further investigation in future studies.

In our previous study, we discovered that circYap inhibited YAP protein expression in breast cancer.17 YAP protein expression was increased in tumor tissues/cells compared to the benign tissue/normal cells. Elevation of circYap could effectively reduce YAP protein expression to the level of control tissues or cells. In the current study, however, YAP mRNA and protein were found to be decreased in parallel to circYap in TAC mouse heart; therefore, circYap is unlikely to antagonize YAP expression itself. The mechanisms of circYap on the YAP signaling pathway in cancer cannot explain the beneficial effects of circYap on heart function. Therefore, other potential mechanisms that mediated the function of circYap in the PO model were investigated in the present study. Here, we demonstrated that circYap inhibited cardiac fibrosis via controlling the interaction of ACTG and TPM4 and the following actin polymerization. TPM plays a critical role in actin polymerization by stabilizing actin filaments. It has been reported that mutation of TPM could result in hypertrophic cardiomyopathy, dilated cardiomyopathy, or left ventricular noncompaction.35 TPM also inhibited new actin filament assembly via restricting the ability of the Arp2/3 complex to nucleate actin polymerization.20 In our study, we found that circYap could bind with both TPM4 and ACTG to form a complex and significantly enhanced the inhibitory effect of TPM4 on actin polymerization. By binding with circYap, the interaction of TPM and ACTG was significantly enhanced. Therefore, TPM played its critical role in actin filament assembly and thus suppressed actin polymerization. Upon TAC-induced PO, actin polymerization in the mouse heart tissue was accelerated, leading to the development of collagen deposition and fibrosis. Ectopic delivery of circYap in our study significantly decreased this process. It has been reported that enhancement of actin polymerization could activate the process of lung fibrosis.27 Here, our study confirmed the link between actin polymerization and cardiac fibrosis and identified circYap as the potential inhibitor to prevent the development of cardiac fibrosis via deactivating actin polymerization.

Although our results showed that circYap could bind to ACTG and TPM4, we did not examine the sizes of the binding complexes. Since actin can form filaments and tropomyosin can bind to actin, there is a possibility that the complexes of circYap–ACTG–TPM4 have different sizes, which depends on the interaction of actin–actin and actin–tropomyosin. The sizes of the complexes may contribute to actin polymerization and cardiac fibrosis. Additional technologies with innovation are needed to uncover the details of the interaction in the future. Due to the potential formation of large complexes of these three molecules, it is also possible that some circYap-binding proteins detected by mass spectrophotometry may be under-representative. Whether these proteins are involved in circYap-regulating cardiac fibrosis warrants further investigation. Moreover, we demonstrated that ectopic circYap elevated the circYap levels in both cardiomyocytes and cardiac fibroblasts in vivo and altered the morphology of cardiomyocytes and cardiac fibroblasts in vitro. Since cardiac fibroblasts are acknowledged to play the major role in cardiac fibrosis,38 we consider the protective effects of circYap on heart function and cardiac fibrosis here is mainly mediated by the inhibition of actin polymerization in cardiac fibroblasts. Our present study may offer a novel and promising approach to impede cardiac fibrosis and improve the heart function by targeting circRNA.

MATERIALS AND METHODS

Human Heart Specimens
Heart samples were obtained from 104 patients with cardiac hypertrophy resulting from heart failure, aortic stenosis, mitral stenosis, hypertrophic cardiomyopathy, or tetralogy of Fallot, and 21 organ donations without heart disease records (trauma or non-cardiac causes). The clinical characteristics and etiology of these patients are described in Table S1. The donation hearts from individuals who died from non-cardiac causes were collected 0.5–6 h after patient’s death. At the time of surgery, the heart tissue was removed from the left ventricular free wall or right ventricular outflow tract and snap-frozen in liquid nitrogen for RNA or protein isolation. The study was carried out in accordance with the Ethics Code of the World Medical Association (Declaration of Helsinki). All patients selected in this study gave formal informed consent prior to enrollment.

Animal Models
The PO model was induced by modified TAC in mice as previous described.39 A successful PO model was confirmed with both visual confirmation of differential carotid pulpalibility and measuring the carotid artery flow velocities by Doppler. Those mice with a right carotid (RC)/left carotid (LC) flow ratio >5 were included for further experiments. The sham mice underwent surgery at the same time points with the similar process of anesthesia and other operation except aortic banding. All animal experiments were performed in accordance with relevant guidelines and regulations approved by the Animal Care Committee of Sunnybrook Research Institute.

For circYap administration, 8-week-old C57BL6 mice were processed to TAC and the plasmids of control vector or circYap (50 μg each) were injected intraperitoneally twice per week for 8 weeks. The plasmids of circYap and vector control were generated as described in the following section of construct generation. The plasmids were
conjugated with polyethylene glycol (PEG) and AU nanoparticles before injection as previously described. Groups of sham and TAC mice without injection served as controls. After 8 weeks of injection, mice were sacrificed following cardiac function assessment. Hearts and other organs were harvested. Part of the heart tissue was fixed with 10% buffered formalin and embedded in paraffin and then sectioned to 5 μm slides. H&E staining, Masson’s trichrome staining, and Sirius red staining were performed as described.

Mice were anesthetized with 2% isoflurane inhalation to undergo transthoracic echocardiography and invasive hemodynamic assessment. Transthoracic echocardiography was performed and analyzed in a blinded manner, using a Vevo 2100 high-resolution imaging system equipped with a 40-MHz transducer to measure LVEDD, LVEDP, LVFS, and dp/dt. A 1.4-Fr high-fidelity pressure catheter (SPR-671, Millar Instruments, Houston, TX, USA) was inserted into the LV via the right carotid artery to measure left ventricular pressure (LVSP) and dp/dt using PowerLab system (AD Instruments).

For Sirius red staining, mouse heart sections were stained with Weigert’s hematoxylin for 8 min after being dewaxed and hydrated. Then, the slides were stained in 0.1% picrosirius red for another 1 h and washed in 0.1% acetic acid. Masson’s trichrome staining was performed with Masson’s trichrome staining kit from American Master Tech.

**FISH**

In FISH, Alexa 488 labeled DNA oligo probes against circYap were generated by fluorescence PCR labeling kit from Biolyx. The labeled probes were heated at 95°C for 5 min and chilled on ice immediately to prevent reannealing. A scramble sequence was used as a negative control. After dehydration and air drying, the fixed samples were pretreated with hybridization solution in 55°C for 0.5 h. The pre-hybridized slides were incubated with 50 nM fluorescence-labeled DNA oligo probes in hybridization buffer at 55°C for 1 h followed by serial washes with saline-sodium citrate (SSC) buffers. DAPI staining was done for another 20 min after washing with PBS before mounting.

**RT-PCR and RNase Treatment**

Total RNA was extracted from cells or tissues using a kit from Geneaid TrRNA isolation kit or TRIzol. 1 μg RNA was subjected to reverse transcription and quantitative PCR (qPCR) using iScript RT kits and SYBR green master mix (Bio-Rad). U6 or GAPDH were used as internal controls. The sequences of primers are listed in Table S2. RNase treatment was conducted as previously described. In brief, 1 μg RNase R (Epicenter) or RNase A (QIAGEN) was added in the mixture either before or after immunoprecipitation and incubated at 37°C for 15 min.

**RNA and Protein Interaction**

For immunoprecipitation, 100 μL magnetic beads were washed in PBS-T (PBS + 0.1% Tween-20) and incubated with 1 μg primary antibody at room temperature for 10 min. The total protein lysates were collected, and the protein concentrations of different samples were equalized. Then, the protein lysate was incubated with antibody-containing beads for 1 h. The magnetic beads were washed 3 times with PBS-T and were resuspended in either TRizol (for RNA extraction) or 2 × Laemmli buffer (0.125 M Tris-HCl, 4% SDS, 20% glycerol, 10% 2-mercaptoethanol, 0.004% bromphenol blue [pH 6.8] for protein isolation). The isolated RNAs were subjected to real-time PCR, while the isolated proteins were subjected to western blotting.

**Bioinformatics Prediction**

To determine the possible interaction of circYap with ACTG1/TPM4, 20,000 models were generated using NPDock server, a protein-RNA docking analysis tool. NPDock server combines GRAMM for global macromolecular docking, scoring with a statistical potential, and clustering, followed by refinement of best-scored docked complexes from the three biggest clusters. Distance-based and residue-level resolution contact maps of circYap-ACTG1 docked complex and circYap-TPM4 docked complex were determined using RNAmap2D and COCOMAPS tools. Contact distances were computed between Cα atoms of protein residues and O5′ atoms of RNA strands. Two residues are in contact when their O5′-Cα distance is less than 10 Å. A distance-based approach was used to identify the binding site residues/nucleotides for the protein-RNA complexes using a specific cutoff value. Two atoms (one in RNA and another in protein) are considered to be interacting with each other if the distance between them is <3.5 Å.

**Actin Polymerization Assay**

Actin polymerization was conducted with the Actin Polymerization Biochem kit (Cytoskeleton). Briefly, the pyrene actin was dissolved in G-buffer containing ATP at 0.4 mg/mL and incubated on ice for 1 h to depolymerize actin oligomers, followed by centrifuge at 14,000 rpm at 4°C for 30 min. The supernatant was transferred to a 96-well plate (200 μL per well). Meanwhile, the cells were lysed in actin-compatible buffer containing 20 mM HEPES, 20 mM NaCl, and protein inhibitors and subjected to centrifugation at 150,000 × g at 4°C for 1 h. 20 μL of the supernatant was added in each well containing G-actin stock. Then, actin polymerization buffer was added in the wells to start the reaction. Kinetic for over 120 cycles, 60 s interval time was set on Multiscan Spectrum (BioTek Synergy I) with the ex.350 nm and em.410 nm.
Statistical Analysis
All experiments were performed in triplicate, and numerical data were subject to independent sample t test for two groups and one-way ANOVA for three or more groups. The levels of significance were set at \( p < 0.05 \) and \( **p < 0.01 \).

Declaration
Our study complies with the Declaration of Helsinki; the Animal Care Committee of Sunnybrook Research Institute has approved the research protocol and informed consent has been obtained from the subjects. Analysis of human heart tissues was approved by the Ethics Committee of Guangdong General Hospital.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.jmther.2020.12.004.

ACKNOWLEDGMENTS
This work was supported by grants from Canadian Institutes of Health Research (PJT-153105 and PJT-155962) to B.B.Y.

AUTHOR CONTRIBUTIONS
B.B.Y. supervised the project. N.W. and B.B.Y. designed the experiments. S.W. and J.X. collected patient samples. N.W., W.W.D., X.L., F.L., S.M., E.E., J.L., L.Z., K.Z., and A.A. performed the experiments. F.M.A. performed computational analysis of the interaction between circYap and the proteins. N.W. and B.B.Y. wrote the paper.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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