Myocardin ablation in a cardiac-renal rat model

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Cardiorenal syndrome is defined by primary heart failure conditions influencing or leading to renal injury or dysfunction. Dilated cardiomyopathy (DCM) is a major co-existing form of heart failure (HF) with renal diseases. Myocardin (MYOCD), a cardiac-specific co-activator of serum response factor (SRF), is increased in DCM porcine and patient cardiac tissues and plays a crucial role in the pathophysiology of DCM. Inhibiting the increased MYOCD has shown to be partially rescuing the DCM phenotype in porcine model. However, expression levels of MYOCD in the cardiac tissues of the cardiorenal syndromic patients and the effect of inhibiting MYOCD in a cardiorenal syndrome model remains to be explored. Here, we analyzed the expression levels of MYOCD in the DCM patients with and without renal diseases. We also explored, whether cardiac specific silencing of MYOCD expression could ameliorate the cardiac remodeling and improve cardiac function in a renal artery ligated rat model (RAL). We observed an increase in MYOCD levels in the endomyocardial biopsies of DCM patients associated with renal failure compared to DCM alone. Silencing of MYOCD in RAL rats by a cardiac homing peptide conjugated MYOCD siRNA resulted in attenuation of cardiac hypertrophy, fibrosis and restoration of the left ventricular functions. Our data suggest hyper-activation of MYOCD in the pathogenesis of the cardiorenal failure cases. Also, MYOCD silencing showed beneficial effects by rescuing cardiac hypertrophy, fibrosis, size and function in a cardiorenal rat model.

DCM is a major cause of HF, accounting for nearly 1/3rd of total cases. Many of these patients subsequently display kidney dysfunction or injury leading to cardiorenal syndrome. More than half of the heart failure patients show renal diseases. Co-existence of cardiac and renal dysfunction in the patients increases the mortality significantly compared to cardiac or renal disease alone patients.

Various molecular pathways including Renin-angiotensin-aldosterone system (RAAS) are shown to be influencing the cardiorenal syndrome. Notably, circulating Ang II (an essential component of RAAS) affects cardiac function by, increasing systemic arteriolar vasoconstriction, vascular resistance, and cardiac afterload through AT1 receptor-mediated endothelial dysfunction.

Ang II has been shown to induce MYOCD under hypoxic condition. MYOCD is a cardiac-specific transcriptional co-activator present in cardiomyocytes and smooth muscle cells. MYOCD is involved in heart development and cardiomyocyte differentiation. Also, MYOCD is required for maintenance of structural integrity, cardiomyocyte survival, and heart function. MYOCD has been shown to promote fibroblast to myofibroblast differentiation and to inhibit cell proliferation. Forced expression of MYOCD in fibroblasts induces cardio-myogetic properties alone and/or in combination with other factors. Transforming growth factor (TGF-β) was shown to induce MYOCD expression in fibroblasts and vice-versa. TGF-β induction of MYOCD expression in the infarcted heart may have a potential function in fibroblast-to-myofibroblast transition, similar to Myocardin related transcription factor MRTF-A and MRTF-B which have been shown to be key regulator in fibroblast to

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myofibroblast differentiation induced by TGF-β1. Further, deletion of MYOCD gene in the adult murine heart resulted in dilated cardiomyopathy, and rapid death due to heart failure. Upregulation of MYOCD expression has been shown in cardiac hypertrophy and MYOCD overexpression in mouse cardiomyocytes resulted in activation of genes associated with cardiac hypertrophy. Increased cardiac MYOCD expression has been reported in various cardiac ailments including DCM patients with end-stage HF. MYOCD has been shown to be a pro-hypertrophic factor in cardiac remodeling induced in multiple models. However, there is no report so far, suggesting the role of MYOCD in cardiorenal syndrome.

In the present study, we analyzed the cardiac-specific expression of MYOCD in DCM patients with renal disease and DCM alone cases. The results showed the MYOCD is overexpressed in the DCM patients with renal disease compared to DCM alone cases. In addition, the effects of cardiac-specific silencing of MYOCD was explored in a cardiac renal syndrome rat model. The cardiac-specific silencing of MYOCD in rats decreased the expression of upregulated hypertrophic and fibrotic genes leading to restoration of left ventricular function.

Material and Methods
Study Population. Thirty consecutive biopsies were taken from left ventricle region from idiopathic DCM (IDCM) patients, attending Cardiology Clinic at the Department of Cardiology, Postgraduate Institute of Medical Education and Research, Chandigarh, India between Jan 2011–2014. Inclusion criteria for recruitment of DCM patients, diagnosed after echocardiography, defined by left ventricular ejection fraction (LVEF) ≤40% and chronic mild to severe HF (NYHA functional class II to IV). All patients underwent left cardiac catheterization and coronary angiography before their inclusion in the study. Exclusion criteria were: the presence of significant coronary artery disease defined as lumen stenosis in >50% of any coronary artery, severe primary valve disease, uncontrolled systemic, hypertension, hypertrophic or restrictive cardiomyopathy, chronic systemic disease like myocarditis, thyrotoxicosis, HIV disease and drug abuse. All recruited IDCM subjects were on optimal medication, angiotensin-converting enzyme inhibitors, and beta-blockers but had persistently low LVEF despite drug regime at the time of biopsy. Endomyocardial biopsy from left ventricle region (n = 15) taken from subjects undergoing surgery for ventricular septal defect (VSD), served as controls. The VSD patients recruited in the study have normal LVEF with no right or left ventricular dysfunction. The study was approved by the Institutional Ethics Committee (8443-PG-1TRg-10/4497), Postgraduate Institute of Medical Education and Research, Chandigarh and written informed consent was taken from all patients for participation in the study. We confirm that all experiments were performed in accordance with relevant guidelines and regulations of Institutional Ethics Committee, Postgraduate Institute of Medical Education and Research, Chandigarh. Inclusion and Exclusion criteria is given in Supplementary Methods. Baseline data of patients and control subjects is available in our earlier publication.

Renal artery ligation model of cardiac renal syndrome (RAL). RAL was developed in adult wistar rats as described earlier. Briefly, right renal artery of 20-week-old male rats (250–300 g; n = 8) was ligated under anaesthesia, rats that underwent similar procedure without ligation served as sham-operated control group (n = 8). Animals were sacrificed 15 days after surgery and hearts were removed and either fixed in Karnovsky’s fixative for histological study or immediately stored in liquid nitrogen for RNA and protein isolation. Cardiac hypertrophy was measured at the gross by heart weight (HW in mg) to body weight (BW in g) ratio. Expression of cardiac fibrosis (Col-1a, Col-4a CTGF, and FGF) and hypertrophy (ANP and β-MHC) genes at mRNA and protein levels was studied by qRT-PCR and western blotting respectively. Histopathological analysis (H&E staining and MT staining) was done in formalin fixed cardiac tissues to assess cardiac remodeling (hypertrophy and fibrosis). This study was approved by Institutional Animal Ethics Committee (IAEC), Postgraduate Institute of Medical Education and Research, Chandigarh and all experiments were done as per the guidelines and regulations of the committee.

Modulation of Myocardin Gene. Myocardin (MYOC) was transiently over-expressed in H9c2 cells or CFs by transfection with 2 µg of pcDNA3-MYOC-HA_v1, using lipofectamine 2000 transfection reagent (Invitrogen) according to the manufacturer’s instructions. For MYOC cellular knockdown, H9c2 cells or CFs were transfected with 1 µg/ml of MYOC siRNA (Table S1 in Supplementary Data) which targets both the cardiac isoforms (Silencer® Select pre-designed MYOC siRNA; ID s140959; Ambion, USA). Cells transfected using scrambled sequence anti-miR (Ambion, USA) served as a negative control. Transfection efficiency was measured using Cy3 labelled negative siRNA by FACS Canto II Cell Analyser and by Olympus IX 73 Fluorescent microscopy.

Cardiac specific silencing of MYOCD Gene. Cardiac MYOCD gene expression was inhibited in RAL rats using MYOC specific siRNA, conjugated with a cardiac specific 20mer homing peptide sequence WLSFAPVVTYRALRGTGSW encapsulated in stearic acid modified carboxy methyl chitosan as described earlier, as per protocol shown in Supplementary Fig. 1. Animals were sacrificed using sodium pantothenate as anaesthesia and performing cervical dislocation. Hearts were removed, examined grossly, and histopathological analysis (H&E staining and MT staining) was performed. RNA and protein was isolated from left ventricular tissue. Expression of hypertrophic (ANP and β-MHC) and pre-fibrotic genes (Col-1a, Col-3a, Col-4a CTGF, TGF β and FGF) was measured at mRNA and protein levels.

Histological analysis. Tissues were fixed in 4% paraformaldehyde, embedded in paraffin, and sectioned at 4-µm thickness. Hematoxylin/eosin for cell size and morphology and Masson’s trichrome for collagen staining were performed using standard procedures. Nuclei were visualized using haematoxylin stain. Images were obtained by Olympus IX 73 microscope.
MYOCD antibody specificity experiment. MYOCD antibody specificity was confirmed by immunostaining in cell lines which have high expression and sparse or null expression of MYOCD to ascertain that it is not binding to other family members of Myocardin like proteins and specific for MYOCD (Supplementary Fig. 2). Imaging was done on Olympus FV1000 confocal microscopy.

Angiotensin II (Ang II) induced cardiomyocyte hypertrophy. H9c2 rat cardiomyocytes were cultured in Dulbecco's Modified Eagle's Medium (DMEM), supplemented with 10% foetal bovine serum under 5% CO2 at 37 °C. After 12 hrs of serum starvation, H9c2 cardiomyocytes were stimulated with 1 µM angiotensin II (Ang-II) (Sigma Aldrich) or 7.5 ng/ml transforming growth factor-β (TGF-β) (Peprotech) for 24 hrs, 48 hrs and 72 hrs. Expression of hypertrophic genes atrial natriuretic protein (ANP) was measured at mRNA and protein levels to confirm the development of hypertrophy.

Ang II induced fibrosis in cardiac fibroblasts. Primary adult rat cardiac fibroblasts (CFs) were isolated from ventricular tissues of 6 to 8 week old male Wistar rats, (150–200 gm) as previously described. After 12 hrs of serum starvation, CFs were stimulated with 1 µM Ang II for 24 hrs (these were optimal conditions for time and dose of Ang II as standardized in our laboratory for induction of fibrosis). Expression of various fibrotic genes (Col-1a, Col-3a, Col-4a, CTGF, and FGF-b) was measured at mRNA levels. Institutional Animal Ethics Committee, Postgraduate Institute of Medical Education and Research, Chandigarh approved all animal experiments were performed in accordance with regulations specified by CPCSEA.

Western Blotting. Cells and tissue protein lysate was prepared using RIPA lysis solution and 1X Protease inhibitor (Cat no.87785, Thermo Scientific), followed by sonication at 30% amplitude repeated for 3–4 times, 20 secs each with a 10 secs interval. Protein concentration was estimated by quick start Bradford dye reagent, 1x (Cat no. 500–0205, Bio-Rad). Laemmli Buffer was used for preparation of proteins. Protein lysate (50 µg/well) was then separated on the basis of their molecular weight on SDS-PAGE Bio-Rad mini SDS-PAGE gel instrument. 50 µg of protein sample along with 5 µl of Page Ruler pre-stained protein ladder (Cat no. 26617, Thermo Scientific), was loaded and gel was run initially at 50 volts until the tracking dye crosses the stacking gel, then at 100 volt till the dye come at the bottom of the gel. After SDS-PAGE, PVDF membrane was activated in 100% methanol and equilibrated in 1X transfer buffer along with SDS-PAGE gel and blotting sheets, at 4 °C for 15 mins. Protein wet transfer was carried out at 30 mA for 10 hr in Mini-Protean Tetra System (Bio-rad). Ponceau staining was performed in order to check the protein transfer on PVDF membrane. PVDF membrane was then washed with 1X TBST buffer and kept in blocking solution (5% BSA in TBST buffer) for 2 hrs on shaker at RT. After blocking, PVDF membrane was washed 3 times in TBST buffer (1x) for 10 mins each and then incubated in primary antibody (List of the antibodies used in the study with the working dilution, source and catalogue number are provided as Table S2 in Supplementary Data) for 4°C overnight. PVDF membrane was then washed 3 times with 1X TBST buffer for 10 mins each and then incubated with HRP labelled secondary antibody for 40 mins at RT. PVDF membrane was then washed 3 times with 1X TBST buffer for 10 mins each and target protein reactive with primary antibodies was detected. After incubation with the appropriate secondary antibodies, blots were visualized using the EZ-ECL detection system (Cat no. 20–500–120A, Biological Industries) and the FluorChem M luminescent image analyzer (Protein Simple, Biotene brand, USA). Densitometry analysis of band intensity was performed using Image J software (NIH, USA).

Figure 1. Myocardin (MYOCD) mRNA and protein expression in myocardial tissues of DCM patients with and without renal failure and controls. (a) Quantitative mRNA expression of MYOCD in DCM cardiac tissues (n = 30) with and without renal failure and control group (n = 15) as determined by qRT-PCR using Taqman probe chemistry. (b) Representative western blot for MYOCD and GAPDH proteins in control (C1–C4) and DCM patients (P1–P5) in LV myocardium samples. Semi-Quantitative protein expression of MYOCD in DCM patients and controls as estimated by software ImageJ. Normalization of total RNA and protein was done using GAPDH as an internal control. Data given is Mean ± SEM; *p < 0.05 Control vs DCM; ***p < 0.001 Control vs DCM or DCM with cardio-renal syndrome.
Statistical Analysis. The statistical analysis was carried out using Statistical Package for Social Sciences (SPSS Inc., Chicago, IL, and version 21.0 for Windows)/Graph Pad Prism 5.0. All quantitative variables were estimated using mean and measures of dispersion (standard deviation and standard error). Normality of data was checked by measures of skewness and Kolmogorov Smirnov tests of normality. Expression of genes were compared using Mann Whitney or Student's t-test depending on the normality of data. One way/Two way ANOVA with appropriate post-hoc tests were performed when multiple comparisons were done within various experimental groups. All statistical tests were two-sided and performed at a significance level of $p \leq 0.05$, $p \leq 0.01$, and $p \leq 0.001$.

Results
Cardiac MYOCD expression level is significantly increased in cardiorenal syndrome patients. We analyzed the mRNA and protein expression levels of MYOCD in endomyocardial biopsies of DCM patients with and without renal failure. The mRNA and protein expression levels of MYOCD was increased in endomyocardial biopsies obtained from DCM patients compared to healthy controls. However, the increase is more pronounced in DCM patients with renal failure compared to DCM alone patients (Fig. 1a,b).

MYOCD activation induces hypertrophic and fibroblast associated genes. To know the functional relevance of increase of MYOCD in cardiorenal patients, we treated cardiomyocytes cells (H9c2) and fibroblasts with Ang II (a well-known modulator of cardiorenal disease). The treatment showed increased mRNA and protein expression of MYOCD in both cell types (Supplementary Fig. 3a–c), along with hypertrophy markers including Atrial Natriuretic factor (ANP), Beta-Myosin heavy chain (β-MHC) and fibrotic genes at mRNA levels such as Collagen (Col) 1a, Col 3a, Col 4a, CTGF, TGF-β and FGF-β respectively (Supplementary Fig. 4a–c). Pre-treatment with...
Losartan (an Ang II type I receptor antagonist) reversed these effects by increasing the levels of MYOCD, ANP and β-MHC mRNA levels in cardiomyocytes, whereas, treatment with known Ang II analogue had no significant effect on MYOCD, ANP and β-MHC expression in these cells (Supplementary Fig. 4d).

Figure 3. Effect of MYOCD siRNA treatment on expression of hypertrophic genes (ANP and β-MHC) and on cardiomyocyte size in RAL rats (n = 8). (a) mRNA expression of MYOCD, ANP and β-MHC in RAL and MYOCD siRNA treated RAL rats. (b) Representative western blot for MYOCD, ANP and GAPDH proteins in RAL and MYOCD siRNA treated RAL rats (before and after ligation). (c) Haematoxylin and Eosin (H & E) stained photomicrographs (10X) of rat hearts: (I) Control group, (II) RAL group (III) Scrambled treated RAL group IV) MYOCD siRNA treated (before ligation) RAL Group V) MYOCD siRNA treated (after ligation) RAL group. Bar diagram shows mean of cardiomyocyte area (n = 50) as measured by Image J analyser. Normalization of total RNA and protein was done using GAPDH as an internal control. Data given is Mean ± SEM; **p < 0.01; ***p < 0.001 for Control vs RAL or scrambled; ##p < 0.01 for RAL or Scrambled vs RAL treated with MYOCD siRNA before ligation (Pre-ligation); $p < 0.05, $$$p < 0.01 $$$p < 0.001 for RAL or Scrambled vs RAL treated with MYOCD siRNA after ligation (Post-Ligation).
Figure 4. Effect of MYOCD siRNA treatment on expression of fibrotic genes (Col 1α, Col 3α, Col 4α, TGF β, CTGF and FGF β) in RAL rats (n = 8). (a) mRNA expression of Col 1α, Col 3α, Col 4α, TGF β, CTGF and FGF β in RAL rats. (b) Representative western blot for CTGF, FGF and GAPDH proteins in RAL rats. (c) mRNA expression of Col 1α, Col 3α, Col 4α, TGF β, CTGF and FGF β in MYOCD siRNA treated RAL rats (before and after ligation). (d) Representative western blot for CTGF, FGF and GAPDH proteins in MYOCD siRNA treated RAL rats (before and after ligation). (e) Photomicrographs (10X) of rat heart tissues with Masson's Trichrome staining which is collagen specific and evaluates fibrosis: (I) control, (II) RAL rats with extensive collagen deposition (III) Scrambled treated RAL group III) MYOCD siRNA (before ligation) treated RAL rats show decreased collagen deposition (IV) MYOCD siRNA (after ligation) treated RAL rats show decreased collagen deposition. Normalization of RNA and protein was done using GAPDH as an internal control. Data given is Mean ± SEM; *p < 0.05; **p < 0.01 for Control vs RAL or scrambled; #p < 0.05; ##p < 0.01 for RAL or scrambled vs RAL treated with MYOCD siRNA before ligation; $p < 0.05; $$p < 0.01 for RAL or scrambled vs RAL treated with MYOCD siRNA after ligation.
In vitro silencing of MYOCD inhibited Ang II induced cardiomyocyte hypertrophy and fibrosis. To examine the effect of MYOCD silencing on cardiomyocyte hypertrophy, Ang II treated H9c2 cells were transfected with MYOCD siRNA. siRNA transfection significantly reduced the MYOCD levels (Supplementary Fig. 5a,b), leading to the reduction of ANP and β-MHC levels (Fig. 2a,b).

Also, treatment of MYOCD siRNA significantly decreased MYOCD mRNA expression levels in the cardiac fibroblasts (Fig. 2c), whereas scrambled siRNA also showed some effect on MYOCD expression in these cells but the changes found in MYOCD siRNA treated cells was much significant. MYOCD siRNA treatment resulted in a significant decrease in mRNA expression of Col 1a, Col 3a, Col 4a, TGF-β, FGF-β and CTGF genes in Ang II treated cardiac fibroblasts (Fig. 2d).

Cardiac-specific inhibition of MYOCD gene in a cardiorenal rat model improves left ventricular functions. To know whether MYOCD inhibition in a cardiorenal rat model can rescue the above observed in vitro phenotype, we performed renal artery ligation (RAL) in rats. This model has been shown to an excellent model for cardiac-renal heart failure20. RAL in rats resulted in cardiac hypertrophy and systolic dysfunction (Fig. 3a,b; and Table 1).

Next, we injected the MYOCD siRNA conjugated with homing peptide through tail veins of the RAL rats as described earlier to be cardiac-specific and having no bystander effects18. To study the effect of heart-specific MYOCD silencing on cardiac function, echocardiography and HW/BW ratio were measured in MYOCD siRNA treated RAL rats 15 days post ligation. HW/BW ratio was significantly reduced in MYOCD siRNA treated RAL rats (Table 1). A significant decrease in LVEDD and increase in LVEF was also seen in these rats (Table 1) indicating improvement in left ventricular function by MYOCD inhibition.

MYOCD inhibition in the RAL model attenuated cardiac hypertrophy and fibrosis. Ablation of MYOCD gene with MYOCD siRNA conjugated with homing peptide in RAL rats resulted in significant reduction in cardiac MYOCD expression (Fig. 3a,b), whereas scrambled siRNA did not affect the MYOCD expression. Also, a profound reduction in the expression of hypertrophy markers, ANP and β-MHC (Fig. 3a,b) and a decrease in cardiac myocyte size in MYOCD siRNA treated RAL rats compared to untreated group were observed (Fig. 3c).

There was no evidence of inflammation or coronary occlusion in these rats, suggesting absence of ischemic lesions (Fig. 3c).

This model showed cardiac fibrosis along with hypertrophy (Fig. 4a,b). Furthermore, MYOCD siRNA treatment, resulted in a significant reduction in cardiac fibrotic gene expressions including Col 1a, Col 3a, Col 4a, TGF-β, CTGF and FGF-β compared to untreated RAL rats (Fig. 4c,d). Histopathological examination showed decreased collagen deposition in the siRNA treated RAL cardiac tissues (Fig. 4e). Collectively, these data suggested that the MYOCD inhibition can reverse the heart failure in RAL model.

Discussion
MYOCD role have been reported in various animal models of HF and in end-stage heart failure patients14,15. However, its role in cardiorenal patients remains unexplored. We observed a significant upregulation of cardiac MYOCD in endomyocardial biopsies from cardiorenal patients. This is the first observation on the MYOCD expression levels in the cardiorenal failure patients.

Cardiorenal syndrome patients display a significant amount of circulating renin with Ang II production, leading to arteriolar constriction and high venous pressure. Ang II along with aldosterone release result in systemic pressure through increased sodium re-absorption in the distal nephrons. These ultimately worsen the patient cardiac functions25. Thus, consistent activation of RAAS contributes to poor prognosis of the cardiorenal patients. RAAS activation is shown to be associated with upregulation of MYOCD related transcription factors leading to increased transforming growth factor-β (TGF-β) and fibronectin14,22. In the present study, direct inhibition of MYOCD in Ang II treated cardiomyocytes and fibroblasts resulted in decreased expression of hypertrophic and pro-fibrotic markers including TGF-β. These data suggest RAAS activation can be mitigated by MYOCD inhibition.

Torrado et al. showed that moderate inhibition of MYOCD using siRNA in doxorubicin-induced HF (DHF) model could partly restore diastolic dysfunction and decreased the DHF animal mortality23. However, such an effect remains to be explored in a cardiorenal model. We explored whether heart-specific MYOCD targeting could ameliorate cardiac remodeling and improve cardiac function in a rat renal artery ligation (RAL) model. RAL is a well-established model for studying the cardiac remodeling in renal dysfunction associated heart failure20.

| Treatment Groups | LVDD (mm) | FS (%) | LVEF (%) | HW/BW ratio |
|------------------|-----------|-------|---------|-------------|
| Group I (Sham Controls) | 4.34 ± 0.20 | 62.08 ± 1.18 | 78 ± 8.8 | 3.2 ± 0.5 |
| Group II (Ligated) | 5.83 ± 0.38* | 34.6 ± 3.8* | 55 ± 6.7* | 4.6 ± 0.7* |
| Group III (Ligated + post-mycd siRNA) | 4.95 ± 0.67* | 56.8 ± 1.97* | 72 ± 5.9* | 3.95 ± 0.3* |
| Group IV (Ligated + pre-mycd siRNA) | 5.05 ± 0.45* | 50.6 ± 2.85* | 61 ± 7.2* | 3.7 ± 0.4* |
| Group V (Ligated + scrambled siRNA) | 5.6 ± 0.4 | 38 ± 2.4 | 52 ± 4.4 | 4.7 ± 0.5 |

Table 1. Echocardiographic parameters and heart weight (HW) to body weight (BW) ratio in RAL model treated with myocd siRNA. LVDD: Left Ventricular Diastolic Diameter; FS: Fraction Shortening; LVEF: Left Ventricular ejection fraction. *p < 0.05 Group I vs Group II or V; †p < 0.05 Group II or V vs Group III; ‡p < 0.05 Group II or V vs Group IV.
Notably, we explored if MYOCD silencing pre- or post-ligation by siRNA could have effective in a RAL rat model showing increased expression of cardiac MYOCD with extensive cardiac hypertrophy and fibrosis. A cardiac specific homing peptide conjugated MYOCD siRNA to specifically inhibit cardiac MYOCD without any specific bystander effects was employed. We observed that this cardiac specific MYOCD silencing resulted in attenuation of cardiomyocyte hypertrophy and cardiac fibrosis when given at either both at pre-ligation or post-ligation stages.

Further, we compared whether siRNA inhibition was more effective at pre- or post-ligation to ascertain the efficacy of MYOCD inhibition on cardiac hypertrophy and fibrosis under these two conditions. In pre-ligation treatment, MYOCD siRNA was introduced before the initiation of cardiac remodeling process, and in post-ligation treatment, MYOCD siRNA was given after the initiation of cardiac remodeling. Both the treatment groups showed significant reduction in the cardiac remodeling and improvement of cardiac function, but post-ligation treatment showed a greater reduction. These results suggest that MYOCD inhibition is more effective at post ligation stage, probably as MYOCD upregulation is maximal at this stage.

Our data suggest cardiac specific MYOCD silencing in cardiorenal rats model (RAL) resulted in the significant reduction in expression of hypertension markers, ANP and β-MHC and decrease in cardiomyocyte size. Further, the histopathological analysis also revealed reduction in cardiomyocyte size will be beneficial in ameliorating the process of cardiac hypertrophy. A significant decrease was observed in the cardiac expression of Col 1α, Col 3α, Col 4α, TGF-β, CTGF and FGF-β genes in MYOCD siRNA treated RAL rats. Histopathological examination corroborated the beneficial effect of MYOCD siRNA, as myocardial collagen deposition decreased in siRNA treated rats, suggesting the potential in controlling the cardiac fibrosis. MYOCD silencing was done in a controlled manner so there are less chances of bystander effects on the normal physiology of the myocytes and fibroblasts. Cardiomyocytes were found to have normal morphology and functional capacity after the MYOCD siRNA treatment.

Our data document for the first time, the upregulation of cardiac MYOCD in cardio-re nal patients and cardiac-specific MYOCD silencing in a cardiorenal model has an integrated beneficiary effect on cardiac hypertrophy, fibrosis and function. Further studies along with newly reported MYOCD antibody will strengthen the therapeutic role of MYOCD in the therapeutic role of MYOCD in reversing cardiac remodeling in the cardiorenal syndrome.

References
1. Hershberger, R. E., Hedges, D. J. & Morales, A. Dilated cardiomyopathy: the complexity of a diverse genetic architecture. *Nat. Rev. Cardiol.* 10, 531–547 (2013).
2. Crowley, S. D. Angiotensin II causes hypertension and cardiac hypertrophy through its receptors in the kidney. *Proc. Natl. Acad. Sci.* 103, 17985–17990 (2006).
3. Chiu, C.-Z., Wang, B.-W., Chung, T.-H. & Shyu, K.-G. Angiotensin II and the ERK pathway mediate the induction of myocardin by hypoxia in cultured rat neonatal cardiomyocytes. *Clin. Sci. Lond. Engl.* 1979(119), 273–282 (2010).
4. Wang, D. et al. Activation of cardiac gene expression by myocardin, a transcriptional cofactor for serum response factor. *Cell* 105, 851–862 (2001).
5. Huang, J. et al. Myocardin is required for cardiomyocyte survival and maintenance of heart function. *Proc. Natl. Acad. Sci.* 106, 18734–18739 (2009).
6. Zhang, X., Zhang, Y., & Wei, J. Y. Identification of a novel serum response factor cofactor in cardiac gene regulation. *Cell* 37, 255–264 (1989).
7. Wang, D. et al. Myocardin induces cardiomyocyte hypertrophy. *Circ. Res.* 98, 1089–1097 (2006).
8. Wang, K., Long, B., Zhou, J. & Li, P.-F. miR-9 and NFATc3 regulate myocardin in cardiac hypertrophy. *Circ. Res.* 103, 2184–2190 (2017).
9. Morita, T. et al. Myocardin silencing reduces cardiomyocyte hypertrophy. *J. Biol. Chem.* 282, 21846–21855 (2007).
10. Wang, J. et al. Myocardin silencing reduces cardiomyocyte hypertrophy. *J. Biol. Chem.* 282, 21846–21855 (2007).
11. Mittal, A., Sharma, R., Prasad, R., Bahl, A. & Khullar, M. Role of cardiac TBX20 in dilated cardiomyopathy. *J. Mol. Med. Berl. Ger.* 81, 566–577 (2003).
12. Kontaraki, J. E., Parthenakis, F. I., Nyktari, E. G., Patrianakos, A. P. & Vardas, P. E. Myocardial gene expression alterations in dilated cardiomyopathy. *J. Mol. Cell. Biol. Tiss. Res.* 5, 411–420 (2003).
13. Official website, https://www.nature.com/articles/cr20095
14. Grossman, R. C. Experimental models of renal disease and the cardiovascular system. *Open Cardiovasc. Med. J.* 4, 257–264 (2010).
15. Clemmer, J. S., Pruett, W. A., Coleman, T. G., Hall, J. E. & Hexter, R. L. Mechanisms of blood pressure salt sensitivity: new insights from mathematical modeling. *Am. J. Physiol.-Regul. Integr. Comp. Physiol.* 312, R451–R466 (2017).
16. TGF-β-induced epithelial to mesenchymal transition | Cell Research. Available at, https://www.nature.com/articles/cr20095 (Accessed: 18th April 2018).
17. Rana, S. et al. A spatio-temporal cardiomyocyte targeted vector system for efficient delivery of therapeutic payloads to regress cardiac hypertrophy abating bystander effect. *J. Control. Release Off. J. Control. Release Soc.* 200, 167–178 (2015).
18. Shivakumar, K., Nair, R. R. & Valiathan, M. S. Paradoxical effect of cerium on collagen synthesis in cardiac fibroblasts. *Arch. Biochem. Biophys.* 351, 281–286 (1998).
19. Lu, Q. et al. CRISPR-Cas9-Mediated Epitope Tagging Provides Accurate and Versatile Assessment of Myocardin-Brief Report. *Arterioscler. Thromb. Vasc. Biol.* 38, 2184–2190 (2018).
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Author Contributions
Anupam Mittal and Perundurai Dhandapany conceived the idea for the study. Anupam Mittal, Madhu Khullar and Perundurai Dhandapany was involved in analysis, interpretation of data and drafting of the manuscript. Anupam Mittal performed majority of the experiments. Santanu Rana helped with RAL model development. Satish K Raut helped in providing the animal data. Akhilesh Kumar, Rajni Sharma, Ajay Bahl, Uma Nahar Saikia, Rishikesh Prasad and Sagartirtha Sarkar provided reagents.

Additional Information
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