

MyoR Modulates Cardiac Conduction by Repressing Gata4

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The cardiac conduction system coordinates electrical activation through a series of interconnected structures, including the atrioventricular node (AVN), the central connection point that delays impulse propagation to optimize cardiac performance. Although recent studies have uncovered important molecular details of AVN formation, relatively little is known about the transcriptional mechanisms that regulate AV delay, the primary function of the mature AVN. We identify here MyoR as a novel transcription factor expressed in Cx30.2⁺ cells of the AVN. We show that MyoR specifically inhibits a Cx30.2 enhancer required for AVN-specific gene expression. Furthermore, we demonstrate that MyoR interacts directly with Gata4 to mediate transcriptional repression. Our studies reveal that MyoR contains two nonequivalent repression domains. While the MyoR C-terminal repression domain inhibits transcription in a context-dependent manner, the N-terminal repression domain can function in a heterologous context to convert the Hand2 activator into a repressor. In addition, we show that genetic deletion of MyoR in mice increases Cx30.2 expression by 50% and prolongs AV delay by 13%. Taken together, we conclude that MyoR modulates a Gata4-dependent regulatory circuit that establishes proper AV delay, and these findings may have wider implications for the variability of cardiac rhythm observed in the general population.

The cardiac conduction system is responsible for coordinating electrical activity in the vertebrate heart (1, 2). Each heartbeat is initiated in the sinoatrial node (SAN) and propagated through the atrioventricular node (AVN), His bundle, bundle branches, and Purkinje fibers. Specifically, the AVN connects the atria with the ventricles and delays impulse propagation to optimize cardiac performance (3). Environmental or genetic factors that perturb atrioventricular (AV) delay can impact a variety of common cardiovascular diseases, such as congestive heart failure, atrial fibrillation, and AV block (4). Therefore, detailed understanding of the transcriptional mechanisms controlling formation and function of the AVN will clarify how normal rhythm is established and pathological arrhythmias arise.

Human and mouse genetic studies have demonstrated that the transcription factors Tbx5 and Nkx2.5 function during development to specify lower AV nodal cells and promote their morphogenesis (5–10). Interestingly, while Tbx5 and Nkx2.5 are expressed broadly within the developing heart to activate transcription, the inhibitory T-box transcription factors Tbx2 and Tbx3 are confined to AV node progenitor cells and repress working cardiomyocyte-specific genes (i.e., Nppa and Cx43) by antagonizing Nkx2.5 (11–15). In addition, well-characterized signaling pathways influence these transcriptional mechanisms to refine AVN morphogenesis. While Notch signaling patterns early AVN development (16), the BMP pathway regulates subsequent AVN morphogenesis through Smad-dependent activation of Tbx2 (17–19). In zebrafish, Foxn4 has also been implicated in AVC patterning via direct activation of Tbx2b (20), but it remains uncertain whether Foxn4 functions similarly in higher vertebrates. Although these studies have begun to characterize pathways that impact early AVN progenitor specification, the transcriptional mechanisms that regulate AV delay, the primary function of the mature AVN, remain to be completely elucidated.

In order to gain insight into the mechanisms that regulate AVN-specific function, we focused on transcriptional regulation of Cx30.2/Gjd3, a gap junction protein required for normal AV delay in mice (21, 22). Cx30.2 expression is controlled by a distal

upstream enhancer that directs expression to the AVN, and analysis of this element revealed that Gata4 activates Cx30.2 expression to establish normal AV delay (23). Given that Gata4 is broadly expressed within the heart and has numerous functions that are critical for normal cardiac development (24), how Gata4 regulates transcription in a tissue-specific manner remains incompletely understood. Tissue-enriched cofactors provide one potential mechanism for dictating such cell-type specificity. For example, the FOG1 and FOG2 proteins are cofactors that regulate GATA-dependent gene expression in specific tissues (25–29). Given that Gata4 plays a central and pervasive role in cardiac gene expression programs (30), we reasoned that additional Gata4-interacting cofactors must exist to regulate tissue-specific gene expression and hypothesized that an AVN-specific Gata4 cofactor could thus modulate its transcriptional activity. In the present study, we used the Cx30.2 enhancer element as a tool to identify novel transcriptional cofactors of Gata4-dependent gene expression in the AVN.

Transcription factors belonging to the beta helix-loop-helix (bHLH) family are prominent regulators of lineage commitment and tissue-specific gene expression (31). For example, the proneural bHLH transcription factors Mash1 and NeuroD turn on genes required for neuronal differentiation (32, 33). In skeletal muscle, a hierarchy of bHLH factors, including MyoD, myogenin, Myf5,

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and MRF4, orchestrate muscle differentiation (34). During heart formation, the bHLH proteins Hand1 and Hand2 are critical for morphogenesis of the left and right ventricle, respectively, while Twist1 has been implicated in outflow tract development (35, 36). Interestingly, Hand2, but not Hand1, has been shown to function as a Gata4 coactivator of heart-specific gene expression (37). Whether there exist additional cardiac tissue-enriched bHLH transcription factors that modulate Gata4 transcriptional activity, however, remains to be clarified.

MyoR (also called musculin in mice and ABF-1 in humans) is an inhibitory bHLH transcription factor that was identified based on its skeletal muscle enrichment (38–40), and combined deletion of MyoR and Tcf21/capsulin impairs formation of specific facial muscles necessary for mastication (41). Here, we show that MyoR is also expressed in the AVN of developing mice and represses Gata4-dependent activation of the Cx30.2 enhancer. We demonstrate that MyoR and Gata4 interact directly with one another, and we define their minimal interaction domains. Interestingly, MyoR binds to Gata4 via a protein domain that also binds to Tbx5, but each transcription factor utilizes a unique binding surface to interact with Gata4. We also show that MyoR contains two discrete repression domains that are required to inhibit transcriptional activity. In particular, the N-terminal MyoR repression domain can function in a heterologous context to convert Hand2 from an activator into a repressor. Consistent with our *in vitro* experiments, we find that genetic ablation of MyoR increases Cx30.2 expression and delays AV conduction. These results implicate the MyoR repressor in a Gata4-dependent transcriptional circuit that establishes normal cardiac conduction.

MATERIALS AND METHODS

Flow cytometry and microarray analysis. All Cx30.2-lacZ mice used in the present study contain a 10.8-kb fragment of genomic DNA that encompasses the endogenous Cx30.2 enhancer fused to a minimal HSP promoter and lacZ coding sequence as described previously (23). Male Cx30.2-lacZ mice were crossed to female ICR mice to generate Cx30.2-lacZ hemizygous day 12.5 embryos, and transgene positive animals were identified by PCR genotyping by using lacZ primers. Embryonic hearts were dissected from individual embryos and placed into ice-cold phosphate-buffered saline (PBS), minced extensively, and digested in 0.25% trypsin (Invitrogen) for 30 min at 37°C with gentle trituration every 10 min. Digested cells were strained and resuspended in a minimal volume of Dulbecco modified Eagle medium (DMEM; Invitrogen) and incubated with an equal volume of 2 mM fluorescein digalactoside (Invitrogen) for 1 min at 37°C. This cell suspension was diluted 10-fold with ice-cold DMEM and placed on ice for 1 h prior to fluorescence-activated cell sorting (FACS) analysis. FACS was performed at the UT Southwestern Flow Cytometry Core Facility, and Cx30.2-β-Gal⁺ and Cx30.2-β-Gal⁻ cells were sorted directly into TRIzol (Invitrogen) for RNA purification. Total RNA was submitted to the UT Southwestern Medical Center Microarray Core Facility, and microarray analysis was performed in triplicate for each sample using an Illumina Mouse-6 v1.1 BeadChip Array. Cx30.2-EGFP mice were generated in a similar fashion to the Cx30.2-lacZ transgenic line (23), except that an enhanced green fluorescent protein (EGFP) cassette replaced the lacZ reporter gene. Cx30.2-EGFP mice were processed like Cx30.2-lacZ mice, except that the AVC region was dissected prior to trypsin digestion and fluorescent labeling was not necessary. Reverse transcription-PCR (RT-PCR) analysis was performed with cDNA created by the oligo(dT) method (Invitrogen) and the indicated coding sequence primers using GoTaq polymerase (Promega, Madison, WI) in a Master-Cycler EP gradient thermal cycler (Eppendorf, Westbury, NY) for 30 to 35 cycles.

Tissue microdissection and qPCR analysis. For AVC dissection of embryonic day 10.5 (E10.5) hearts, the atria and ventricles were carefully removed. The remaining tissue, comprising the AVC, was trimmed and frozen in liquid nitrogen prior to RNA isolation and cDNA synthesis. Real-time PCR was performed as previously described (68) using gene-specific probes (ABI, Foster City, CA). PCRs were performed in an ABI Prism 7000 sequence detection system and analyzed using the $\Delta\Delta C_T$ method using GAPDH for normalization according to the manufacturer's instructions. Each quantitative PCR (qPCR) was performed in triplicate and averaged before final values were calculated.

Section immunostaining. E16.5 Cx30.2-EGFP mouse hearts were dissected in cold PBS, fixed in 4% paraformaldehyde for 30 min at 4°C, equilibrated in 30% sucrose, embedded in blocks containing tissue freezing medium (Electron Microscopy Sciences, Hatfield, PA), and snap-frozen prior to sectioning with a cryostat. Consecutive 8-μm sections were obtained, and the position of the AVN was identified based on anatomical landmarks from a survey using hematoxylin and eosin staining and Hcn4 immunostaining. Consecutive sections through the AVN were stained with GFP (Life Technologies, Grand Island, NY), Gata4 (Santa Cruz Biotechnology, Dallas, TX), MyoR (Santa Cruz Biotechnology), and Hcn4 (Alomone Labs, Jerusalem, Israel) primary antibodies, followed by detection with Alexa Fluor-labeled secondary antibodies (Life Technologies) according to the manufacturer's suggested protocol.

DNA constructs. The Cx30.2-Luc reporter construct was generated by subcloning a 4.1-kb Cx30.2 enhancer fragment (−4.1 to 0.0 relative to the translation start site) using KpnI-XhoI into pGL2 E1b LUC, which contains a minimal TATA box, followed by the luciferase coding sequence. The Gata4, MyoR, and Tbx5 GST and His fusion constructs were generated by PCR amplification of their coding regions using primers containing BamHI and EcoRI linkers. Each coding sequence amplicon was digested and cloned directly into pGEX 6P1 (GE Healthcare Life Sciences, Pittsburgh, PA) or PRSET A (Life Technologies). Gal4-MyoR fusion eukaryotic expression constructs were created by PCR amplification of the appropriate fragment using primers containing EcoRI and SalI linkers. Digested MyoR amplicons were directly cloned into the pM vector (Clontech, Mountain View, CA). For chimeric MyoR-Hand2 constructs, PCR was performed on the N terminus of MyoR using reverse primers that contained an overlapping segment of the Hand2 sequence, and this amplicon was used as a megaprimer to amplify the C-terminal portion of Hand2; the Hand2-MyoR construct was created similarly. The outside primers for each chimeric construct contained EcoRI and SalI linkers that were used to clone directly into the pM vector.

Transient transfections. Cotransfection experiments were carried out in COS7 cells with Fugene (Promega) according to the manufacturer's suggested protocol. Briefly, 100 ng of reporter plasmid was cotransfected with 30 to 100 ng of Gata4, MyoR, Hand1, LexA-VP16, or Gal4 fusion expression constructs. All experiments were carried out in triplicate at least three times. Reporter expression was normalized by cotransfection of a cytomegalovirus (CMV)-lacZ plasmid, and β-galactosidase activity was used to calculate relative luciferase units. The luciferase activity was determined at 48 to 72 h using a kit from Promega according to the manufacturer's recommendations. Expression of transfected fusion proteins was verified by Western blot analysis of cellular extracts.

Western blotting, coimmunoprecipitation, and ChIP assays. Flag- and Myc-tagged expression plasmids were transfected into COS7 cells, and whole-cell lysate was prepared 72 h posttransfection. For Western blotting, whole-cell lysates were separated by SDS-PAGE and subjected to immunoblotting with a Myc antibody. Alternatively, lysates were immunoprecipitated with Flag-agarose (Sigma, St. Louis, MO) and washed, followed by SDS-PAGE and Western blotting. Experiments were performed on immunoprecipitated lysates alongside parallel input blots with Myc and Flag antibodies. For chromatin immunoprecipitation (ChIP) experiments, COS7 cells were transfected with the indicated eukaryotic expression constructs and enhancer templates. The cells were cross-linked, harvested, and sheared according to a published protocol (42).

Sheared chromatin was immunoprecipitated with a Flag antibody (Sigma, Milwaukee, WI) that had been coupled to protein G-Dynabeads (Invitrogen). The immunoprecipitates were washed extensively, and associated genomic DNA was eluted and used as the template in qPCRs using Cx30.2 enhancer primers. For *in vivo* ChIP assays, pooled AVC and non-AVC tissue were isolated from E16.5 Cx30.2-lacZ embryos, snap-frozen, and pulverized prior to cross-linking. Preparation of sheared chromatin and subsequent immunoprecipitation were carried out essentially as described above except that a MyoR antibody (Santa Cruz Biotechnology) was used. All qPCRs were performed at least in triplicate, and the resulting average occupancy values were normalized to input and expressed as the fold enrichment relative to control. Sequential ChIP experiments were conducted essentially as previously described (43) using a Gata4 antibody (Santa Cruz Biotechnology) that was biotinylated with a commercially available kit (Thermo Scientific, Grand Island, NY) according to the manufacturer's suggested protocol. Sequentially immunoprecipitated fragments were interrogated by qPCR, and enhancer occupancy was calculated relative to mock immunoprecipitation controls and compared to the GAPDH (glyceraldehyde-3-phosphate dehydrogenase) coding sequence.

GST pulldown assays. Protein-protein interaction experiments were carried out using standard methods. Briefly, the indicated proteins were *in vitro* translated using a coupled transcription-translation kit (Promega) in a 50- μ l reaction mixture containing [³⁵S]methionine and other reagents recommended by the manufacturer. Beads containing normalized amounts of Gata4 or MyoR glutathione S-transferase (GST) fusion proteins, expressed and purified as described previously (44), were equilibrated in buffer A (50 mM NaCl, 20 mM Tris-HCl [pH 8.0], 0.05% NP-40, 0.25% bovine serum albumin [BSA], 1 mM phenylmethylsulfonyl fluoride, and 1 mM dithiothreitol) for 30 min at 4°C. Subsequently, 5 μ l of *in vitro*-translated protein was added, and the reaction mixtures were incubated at 4°C for 2 h. For competition assays, highly purified recombinant His-MyoR was added during the incubation period. The beads were then washed three times with buffer A and once with buffer A without BSA, and bound ³⁵S-labeled proteins were visualized on a 10 to 15% SDS-PAGE gel, stained and destained, submerged in fluorographic solution (Amplify; GE Healthcare Life Sciences), dried, and exposed overnight at -80°C.

Mouse breeding. All animal procedures were approved by the Institutional Animal Care and Use Committee at UT Southwestern Medical Center. Cx30.2-lacZ transgenic and MyoR-null mice have been previously described (23, 38). Cx30.2-lacZ and Cx30.2-EGFP male mice were crossed with ICR female mice, and the vaginal plug was considered to be 0.5 days postcoitus (dpc). Similarly, for experiments involving embryonic heart microdissection, wild-type C57BL/6 males and females were crossed, and the vaginal plug was considered to be 0.5 dpc. To generate MyoR^{+/+}; Cx30.2-lacZ/+ and MyoR^{-/-}; Cx30.2-lacZ/+ animals, Cx30.2-lacZ/+ mice were first crossed with MyoR^{-/-} mice to obtain Cx30.2-lacZ/+; MyoR^{+/-} mice, which were intercrossed to generate the indicated strains. MyoR^{+/+} and MyoR^{-/-} animals for qPCR and EKG analysis were generated by intercrossing MyoR^{+/-} mice.

β -Galactosidase assays. β -Galactosidase assays were performed as described previously (23). Briefly, embryos were removed at E16.5 and placed into ice-cold PBS, and AVCs were microdissected and flash-frozen in liquid nitrogen. Embryonic AVCs were subsequently lysed in 100 μ l of passive lysis buffer (Promega) and homogenized by repeated passage through a pipette tip. Protein lysates were incubated on ice for 30 min with frequent agitation and were centrifuged to remove debris. Lysate (20 μ l) was incubated with LacZ assay buffer (10 mM MgCl₂, 50 mM β -mercaptoethanol, 67 mM Na₃PO₄ and 0.88 mg/ml ONPG [*o*-nitrophenyl- β -D-galactopyranoside]) for 1 h at 37°C prior to the addition of 1 ml of stop solution (1 M Na₂CO₃) and the A₄₂₀ measurement. Reactions were performed in triplicate, and samples of the indicated genotypes were averaged together to generate the final results. A sample of each lysate (10 μ l) was used for the BCA assay (Pierce, Rockford, IL) to determine protein concentrations and normalize the β -galactosidase activity of each sample.

EKG analysis. Electrocardiograms (EKGs) were performed as previously described (23) on adult mice at 4 to 6 weeks of age anesthetized with 50 μ g/kg of pentobarbital delivered by intraperitoneal injection. Mice were laid down on a heating pad to maintain core body temperature, and limb leads were placed subcutaneously. Lead II EKGs were recorded at a sampling rate of 4,000 Hz (ADInstruments, Colorado Springs, CO) at rest. EKG intervals were measured by using the LabChart software package (ADInstruments) by an investigator who was blinded to the genotypes of the mice. Comparisons in all experiments were made between MyoR^{-/-} and wild-type age-matched littermates.

Statistics. All data are expressed as means \pm the standard errors of the mean. *P* values were determined by using the Student two-tailed *t* test, and statistical significance was considered for *P* < 0.05.

RESULTS

Identification of MyoR as an AVN-enriched transcript. We previously generated a transgenic Cx30.2-lacZ mouse line marking cells of the developing AVN (23). In order to identify genes enriched in the developing AVN, we performed microarray analysis on Cx30.2-lacZ⁺ cells at E12.5 (Fig. 1A). As expected, Cx30.2 transcripts were enriched 70-fold in isolated Cx30.2-lacZ⁺ cells, and several ion channel subunits known to be expressed in the AVN (e.g., Kcne1 and Cacna2d2) emerged at the top of our gene list (45), thereby confirming our cell isolation strategy (Fig. 1B). Moreover, Tbx2, Tbx3, and Tbx5, which are known transcriptional regulators of AVN specification and/or morphogenesis (2), were enriched in Cx30.2-lacZ⁺ cells. Unexpectedly, our microarray analysis also identified enrichment of MyoR, a transcriptional repressor known to regulate facial muscle formation (41), in Cx30.2-lacZ⁺ cells.

Although the microarray results suggested AVN enrichment of MyoR expression, we wanted to validate these results using an alternative approach. Therefore, we manually dissected the AV canal (AVC) region of E10.5 hearts to isolate tissue containing AV nodal cells. Compared to the whole heart, gene expression analysis of E10.5 AVC tissue demonstrated enrichment of AVC-specific markers (Cacna2d2, Tbx2, and Tbx3) and depletion of the chamber-specific marker Nppa (15), suggesting that microdissection was successful (Fig. 1C). Importantly, MyoR expression was ~3.5-fold enriched in E10.5 AVC, which compares favorably with the established regulator of AVN formation, Tbx3. Taken together, these results demonstrate that MyoR is expressed in the developing heart with particular enrichment in Cx30.2⁺ AV nodal cells.

To determine whether MyoR expression is developmentally regulated, we performed qPCR analysis on RNA prepared from whole hearts isolated from staged embryos and adults. Interestingly, MyoR expression increases during embryogenesis and peaks during late gestation with an ~20-fold increase at E16.5 compared to E10.5 (Fig. 1D). Although MyoR expression reaches a maximum at E16.5, it is worth noting that Gata4 and Tbx5 are ~70- and ~10-fold more abundant than MyoR, respectively, at the same developmental time point (data not shown). Given that MyoR expression peaks at E16.5, we microdissected the AVC of E16.5 Cx30.2-EGFP and isolated Cx30.2-EGFP⁺ cells by flow cytometry for gene expression analysis. Using this approach, we found that MyoR was ~14-fold enriched in Cx30.2-EGFP⁺ AV nodal cells compared to Cx30.2-EGFP⁻ AVC cells at E16.5, confirming that MyoR is enriched in Cx30.2-EGFP⁺ cells just prior to birth (Fig. 1E).

To determine whether MyoR protein has a similar expression

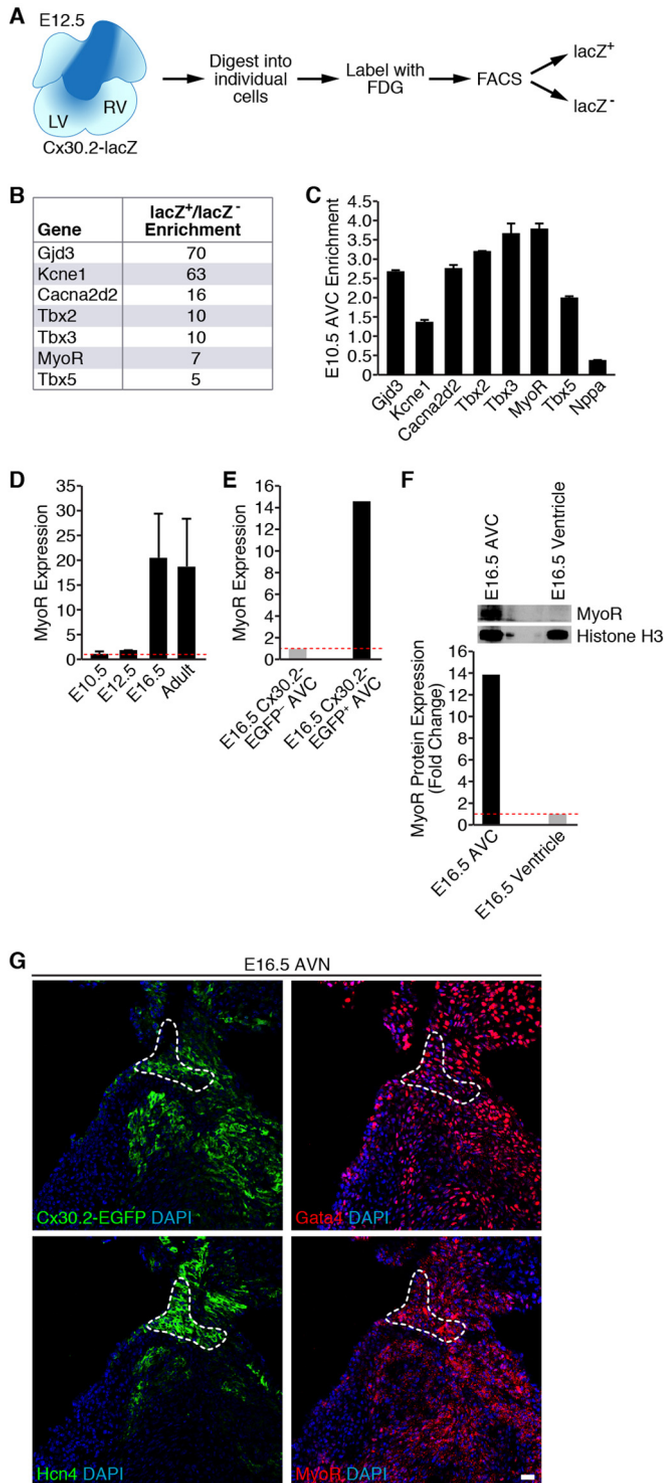


FIG 1 Transcriptional profiling of Cx30.2-lacZ⁺ cells identifies MyoR as an AVC-enriched transcript. (A) Strategy used for purifying E12.5 Cx30.2-lacZ⁺ AVC cells by flow cytometry. (B) Partial list of genes found to be enriched in microarray analysis. Fold enrichment refers to transcript levels in Cx30.2-lacZ⁺ cells relative to lacZ-null cells. Note that MyoR transcripts were 7-fold more abundant in Cx30.2-lacZ⁺ cells within the E12.5 heart. (C) qRT-PCR analysis of microdissected E10.5 AVC tissue demonstrating enrichment of MyoR. Fold enrichment refers to transcript levels in dissected AVC tissue versus whole heart. (D) Developmental analysis of MyoR expression by qRT-PCR analysis. MyoR expression in the whole heart at a particular time point

pattern, we dissected E16.5 AVC and the remaining non-AVC (atrial and ventricular) tissue for Western blot analysis. Consistent with our qRT-PCR analysis, MyoR protein was expressed in the E16.5 AVC ~14-fold more than non-AVC tissue (Fig. 1F). We also performed immunostaining of Cx30.2-EGFP E16.5 heart sections to assess the *in situ* localization of MyoR protein in relation to Cx30.2, Gata4, and the AVN marker Hcn4. This experiment revealed that Cx30.2, Gata4, and MyoR are coexpressed in the Hcn4⁺ AVN (Fig. 1G). Interestingly, we found that MyoR localizes to both the nucleus and cytoplasm, raising the intriguing possibility that nucleocytoplasmic shuttling regulates MyoR transcriptional activity. Collectively, our gene and protein expression data demonstrate that MyoR is a novel AVN-enriched transcription factor and suggest that it may function within the AVN during late gestation.

MyoR inhibits Gata4-dependent Cx30.2 activation. Given that MyoR is expressed in Cx30.2⁺ AV nodal cells, we hypothesized that it regulates Cx30.2 gene expression. Since Hand2, another bHLH family member, directly interacts with Gata4 to regulate cardiac gene transcription (37), we wondered whether MyoR similarly targets Gata4 to modulate Cx30.2 gene expression. In order to test this possibility, we performed transient-transfection analysis using a Cx30.2-Luc reporter construct (Fig. 2A) and various combinations of Gata4, MyoR, and Hand1 (Fig. 2B). As expected, Gata4 activated Cx30.2 reporter expression ~3.5-fold (Fig. 2B), while the addition of MyoR almost completely abolished Gata4-dependent gene activation. In contrast, Hand1, which does not interact with Gata4 (37), does not repress Gata4-dependent activation of the Cx30.2 reporter, suggesting that repression is specific to MyoR rather than a general property of bHLH factors in this system. Thus, these results demonstrate that MyoR inhibits Gata4-dependent transcription of a Cx30.2 reporter gene, which is consistent with previous experimental evidence showing that MyoR is a transcriptional repressor (38, 40).

Since MyoR was able to repress Gata4-dependent transcriptional activity, we also wondered whether MyoR and Gata4 could interact with one another. Therefore, we performed coimmunoprecipitation analysis using a Flag antibody on cells transfected with Myc-Gata4, along with empty vector or Flag-MyoR (Fig. 2C). This experiment showed that Gata4 was immunoprecipitated with a Flag antibody in a MyoR-dependent fashion, demonstrating that these two proteins do interact. Since MyoR interacts with Gata4 to repress Cx30.2 transcriptional activation, we also wanted to evaluate whether MyoR could associate with Gata4 on the Cx30.2 enhancer. Therefore, we performed ChIP assays on cells overexpressing MyoR alone or in combination with Gata4. Although MyoR weakly associated with the Cx30.2 enhancer in the

relative to E10.5 is shown. MyoR transcripts were found to peak during late gestation (E16.5). (E) qRT-PCR analysis of MyoR expression level in Cx30.2-EGFP⁺ AVC cells isolated at E16.5. MyoR expression is shown relative to ventricular levels in EGFP⁻ AVC cells. (F) Western blot analysis of AVC and ventricular nuclear extracts dissected from E16.5 embryos (top) with the indicated antibodies. Band quantitation and normalization demonstrated ~14-fold enrichment of MyoR in the E16.5 AVC relative to ventricular tissue. (G) Immunostaining of consecutive sections through an E16.5 Cx30.2-EGFP heart for GFP, Gata4, Hcn4, and MyoR, demonstrating localization of Cx30.2, Gata4, and MyoR to the Hcn4⁺ AVN (dashed outline). All sections were counterstained with DAPI to stain cell nuclei. Scale bar, 20 μ m. LV, left ventricle; RV, right ventricle; FDG, fluorescein digalactopyranoside; AVC, atrioventricular canal; AVN, atrioventricular node.

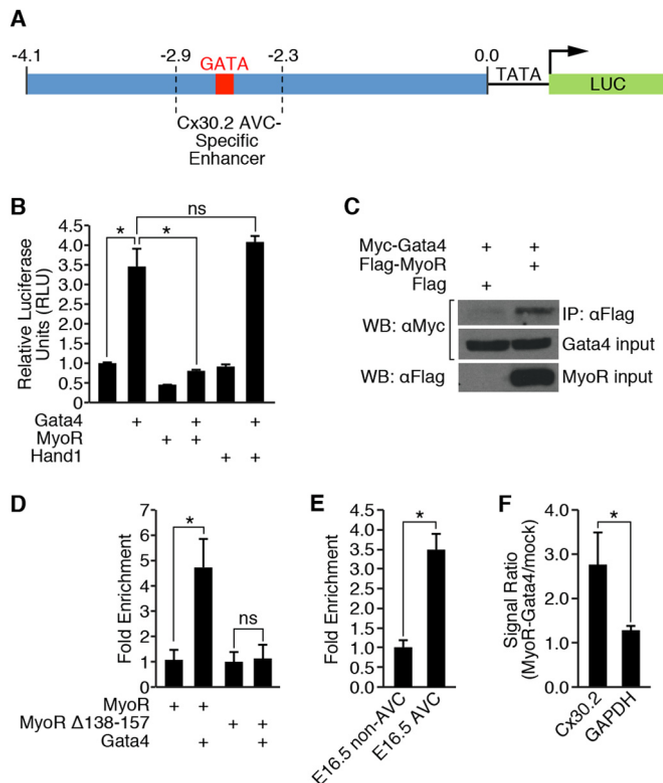


FIG 2 MyoR inhibits Gata4-dependent activation of the Cx30.2 enhancer. (A) Schematic depiction of the Cx30.2-Luc reporter construct used in panel B that contains the Cx30.2 enhancer fused to a minimal TATA box and luciferase coding sequence. The 4.1-kb enhancer fragment encompasses the $-2.9/-2.3$ region that contains the minimal AVC-specific Cx30.2 enhancer (23). (B) Transient-transfection analysis in COS cells. A total of 30 ng of Cx30.2-Luc reporter was transfected, along with 100 ng of Gata4 and/or 100 ng of bHLH transcription factor (MyoR or Hand1). A 10-ng portion of CMV-lacZ was also cotransfected as an internal control. MyoR repressed Gata4-dependent transcriptional activity driven by Cx30.2-Luc, while Hand1, another bHLH transcription factor expressed in the heart, did not. (C) Coimmunoprecipitation analysis between MyoR and Gata4. Myc-Gata4 and either Flag vector alone or Flag-MyoR were cotransfected into COS7 cells. Whole-cell lysates were precipitated with a Flag antibody and probed with either a Myc antibody (top panel). A 10% input was also run separately and probed with either a Myc antibody (middle panel) or a Flag antibody (lower panel). Myc-Gata4 was only detected after Flag immunoprecipitation in the presence of Flag-MyoR but not Flag alone. (D) A ChIP experiment was performed in COS7 cells transfected with a Cx30.2 enhancer template and the indicated proteins. MyoR associated with the Cx30.2 enhancer in a Gata4-dependent manner, but the association with the enhancer was strongly diminished in the presence of a MyoR mutant (MyoR Δ 138-157) that does not bind to Gata4. (E) MyoR ChIP from dissected E16.5 AVC and non-AVC (atrial and ventricular) tissue showed that MyoR preferentially associates with the Cx30.2 enhancer in AVC tissue compared to non-AVC tissue. (F) Sequential MyoR and Gata4 ChIP experiment from E16.5 heart tissue demonstrated that both MyoR and Gata4 simultaneously associate with the Cx30.2 enhancer *in vivo*. *, $P < 0.05$; ns, not significant.

absence of Gata4, we observed augmentation of MyoR binding (~ 4.5 -fold) in the presence of Gata4 (Fig. 2D). This result demonstrates that MyoR associates with the Cx30.2 enhancer and that this interaction is largely Gata4 dependent. Taken together, our results reveal that MyoR interacts with Gata4 on the Cx30.2 enhancer to repress Cx30.2 gene activation and suggest that MyoR mediates its transcriptional effects by interacting directly with Gata4.

While these experiments demonstrate that MyoR can associate

with the Cx30.2 enhancer, they do not address whether this interaction takes place in the developing heart. Therefore, we dissected E16.5 AVC and non-AVC tissue to perform ChIP experiments using a MyoR antibody. Consistent with our *in vitro* data, MyoR associated more strongly with the Cx30.2 enhancer in the AVC relative to the atria and ventricles (Fig. 2E). Furthermore, sequential ChIP experiments with MyoR and Gata4 antibodies demonstrated that both MyoR and Gata4 simultaneously occupy the Cx30.2 enhancer *in vivo* (Fig. 2F). Based on these results, we conclude that MyoR and Gata4 cooccupy the Cx30.2 enhancer in AVC tissue, and these findings are consistent with both MyoR enrichment in the AVN and its proposed role in AVN gene expression.

MyoR and Gata4 interact directly. Previous work has demonstrated that Hand2, but not Hand1, functions as a Gata4 coactivator due to its unique ability to interact directly with Gata4 (37). Given the tight link between Gata4 interaction and the functional impact of Hand2, we speculated that MyoR inhibits Gata4 via direct protein-protein interaction. To test this notion, we carried out GST pull-down assays using bead-bound GST-Gata4 protein and the 35 S-labeled MyoR C-terminal deletions shown in Fig. 3A. This experiment demonstrated that full-length MyoR interacts directly with Gata4 (Fig. 3B, lane 1) in the absence of additional cellular cofactors. Furthermore, while C-terminal deletion of MyoR to amino acid residue 157 (construct 1-157) did not affect its interaction with Gata4 (Fig. 3B, lane 2), deletion of an additional 19 residues (construct 1-138) abolished this interaction (Fig. 3B, lane 3), suggesting that amino acids 138 to 157 of MyoR are necessary to bind Gata4; these residues correspond to the second helix (H2) of the MyoR bHLH domain. Notably, none of the other C-terminal deletion mutant MyoR proteins bound directly to GST-Gata4 (Fig. 3B, lanes 4 to 6). To confirm these results, we created an internal deletion mutant of MyoR (1-201 Δ 138-157) that lacked the H2 segment. Although full-length MyoR was able to bind directly to Gata4, the internal deletion mutant was unable to do so (Fig. 3B, lanes 7 to 8). To further substantiate this finding, we performed ChIP assays in cells overexpressing a MyoR mutant devoid of its Gata4-interaction domain (Δ 138-157) alone or in combination with Gata4. Although full-length and mutant MyoR both bound weakly to the Cx30.2 enhancer, the interaction of MyoR Δ 138-157 with DNA was not augmented in the presence of Gata4 (Fig. 2D). Altogether, these results demonstrate that MyoR interacts directly with Gata4 via helix 2 of its bHLH domain and support the notion that this interaction is required for association of MyoR with the Cx30.2 enhancer.

In addition to identifying the Gata4-binding domain of MyoR, we wanted to map the reciprocal MyoR interaction surface on Gata4. Thus, we performed GST pull-down assays with bead-bound GST-MyoR and the 35 S-labeled Gata4 deletion constructs depicted in Fig. 4A. Again, we confirmed that GST-MyoR interacts directly with full-length Gata4 (Fig. 4B, lane 1). Deletion of the C-terminal 68 residues of Gata4 (1-332) was dispensable for binding to MyoR (Fig. 4B, lane 2), but removal of the next 38 residues or more (constructs 1-294, 1-260, and 1-177) severely diminished this interaction (Fig. 4B, lanes 3 to 5). Additional Gata4 deletion proteins that retained residues 294 to 332 (177-332, 177-332 Δ 265-294, 1-332 Δ 265-294, and 1-332 Δ 178-261) were still able to bind MyoR (Fig. 4B, lanes 6 to 9), suggesting that the Gata4 protein domain just C-terminal to the second zinc finger is necessary for this interaction. To further confirm these find-

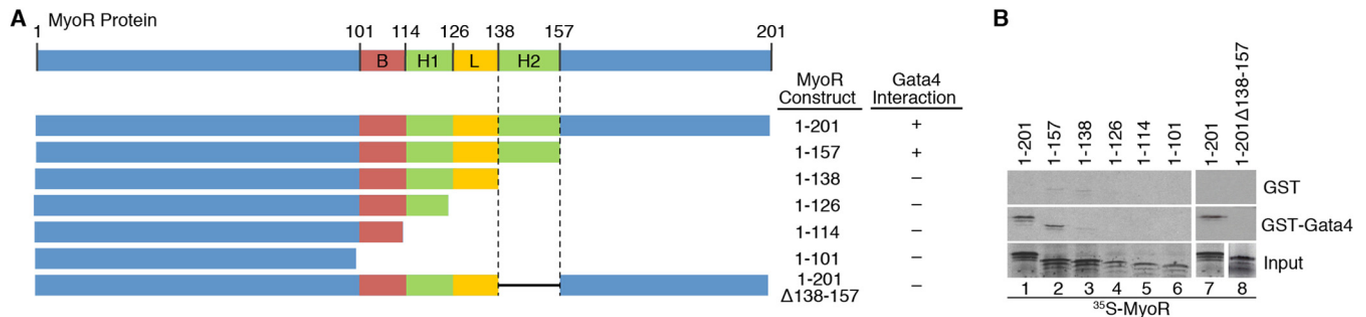


FIG 3 Gata4 interacts directly with the H2 domain of MyoR. (A) Schematic diagram of the MyoR deletion proteins used in GST pull-down experiments with summarized Gata4 interaction data. (B) *In vitro*-translated ³⁵S-labeled MyoR proteins were incubated with either GST alone or GST-Gata4 to test for direct interaction. Input proteins are shown in the bottom panel. Full-length MyoR and a C-terminal deletion to residue 157 interact directly with Gata4 (lanes 1 and 2). MyoR C-terminal deletion beyond amino acid 157 or internal deletion of residues 138 to 157 abrogates interaction with Gata4 (lanes 3 to 8), demonstrating that helix 2 (H2) of the bHLH domain mediates direct contact with Gata4. B, basic domain; H1, helix 1 domain; L, loop domain; H2, helix 2 domain.

ings, we generated a Gata4 internal deletion mutant protein lacking the presumptive MyoR interaction domain (1-440Δ294-332) and performed a GST pull-down assay with full-length MyoR. As expected, this experiment showed that a mutant Gata4 lacking residues 294 to 332 failed to interact with MyoR (Fig. 4B, lanes 11 and 12). Taken together, these studies demonstrate that MyoR interacts directly with Gata4 via amino acid residues 294 to 332, which comprise a domain with unknown structure lying adjacent to its C-terminal zinc finger.

MyoR and Tbx5 bind to unique sites on Gata4. Although the MyoR interaction site on Gata4 (positions 294 to 332) does not belong to any known protein domain family, this region has been shown previously to interact with Tbx5 (46). A specific GATA4 point mutation (G296S) segregates with autosomal dominant atrial septal defects (ASDs) and disrupts its interaction with Tbx5 (46). This observation provided strong genetic and biochemical evidence for transcriptional synergy between Gata4 and Tbx5 during cardiac morphogenesis. Interestingly, we have demonstrated that Gata4 and Tbx5 coordinately activate the Cx30.2 enhancer (23), suggesting that MyoR may inhibit Gata4-dependent transcriptional activation by perturbing the interaction between Gata4 and Tbx5. Given that Tbx5 interacts with Gata4 via a critical protein-protein interaction domain centered on the orthologous mouse amino acid residue 295, we wanted to test whether MyoR inhibits Cx30.2 transcription by competing with Tbx5 for interaction with Gata4. Since the G295S mutation abrogates the ability of Gata4 to interact with Tbx5 (46), we hypothesized that this mutation alters a common Gata4 binding surface utilized by Tbx5 and MyoR. Surprisingly, the Gata4 G295S mutant protein retained the ability to bind MyoR (Fig. 4B, lane 10), suggesting that Tbx5 and MyoR interact with distinct binding surfaces on Gata4. To confirm this result, we took the complementary approach of performing a GST pull-down competition assay. We first added ³⁵S-labeled full-length Gata4 to bead-bound GST, GST-Tbx5, or GST-MyoR, confirming that Gata4 interacts directly with Tbx5 and MyoR (Fig. 4C, lanes 1, 2, and 4). To test whether MyoR and Tbx5 compete for a common Gata4 binding site, we added unlabeled recombinant MyoR to each binding reaction. Although the addition of unlabeled MyoR displaced ³⁵S-labeled Gata4 (³⁵S-Gata4) from bead-bound GST-MyoR (Fig. 4C, lanes 4 and 5), it was unable to displace ³⁵S-Gata4 from GST-Tbx5 (Fig. 4C, lanes 2 and 3), supporting the notion that Tbx5

and MyoR bind to distinct surfaces of Gata4. Taken together, our results strongly suggest that direct competition with Tbx5 is not a major mechanism by which MyoR represses Gata4-dependent transcriptional activity.

MyoR contains two distinct repression domains. Although previous studies have established that MyoR is a transcriptional repressor (38, 40), the mechanism by which MyoR inhibits transcription remains unknown. Given that MyoR does not appear to inhibit Cx30.2 expression by competing with Tbx5 for binding to Gata4, we sought to test whether MyoR contains a discrete repression domain. To address this question, we used a *trans*-repression assay (Fig. 5A) using a synthetic promoter bearing adjacent LexA and GAL4 binding sites driving luciferase (L8G5-Luc) expression. As expected, transfection of LexA-VP16 resulted in robust activation of the L8G5-Luc reporter (Fig. 5B). When Gal4-MyoR was cotransfected with LexA-VP16, we observed potent transcriptional repression, suggesting that MyoR possesses a *trans*-acting repression domain. In order to map the putative MyoR repression domain, we generated several Gal4 MyoR fusion proteins (Fig. 5B). Deletion of the N-terminal 75 residues (Gal4-MyoR 75-201) had a negligible effect on the ability of Gal4-MyoR to inhibit transcriptional activation by LexA-VP16. Similarly, deletion of the C-terminal 44 residues (Gal4-MyoR 1-157) did not significantly impair the ability of Gal4-MyoR to repress LexA-VP16, suggesting that residues 75 to 157 contain the MyoR repression domain(s). Consistent with these findings, Gal4-MyoR 75-157 retained the ability to repress LexA-VP16-dependent activation. Further deletion of residues 138 to 157 (Gal4-MyoR 75-138) or 75 to 100 (Gal4-MyoR 101-157) decreased, but did not completely abolish, the ability of Gal4-MyoR to repress transcriptional activation by LexA-VP16. These results indicate that MyoR contains two separate *trans*-repression domains. To confirm and extend these findings, we generated a Gal4-MyoR construct lacking both putative repression domains (75 to 100 and 138 to 157) and assessed its ability to repress LexA-VP16-dependent transcriptional activation; this MyoR mutant protein was unable to repress activation by LexA-VP16. Taken together, these results demonstrate that MyoR contains two discrete repression domains comprised of residues 75 to 100 and 138 to 157 that are both required for transcriptional inhibition.

The N terminus of MyoR converts Hand2 into a repressor. The bHLH family comprises a diverse set of transcription factors

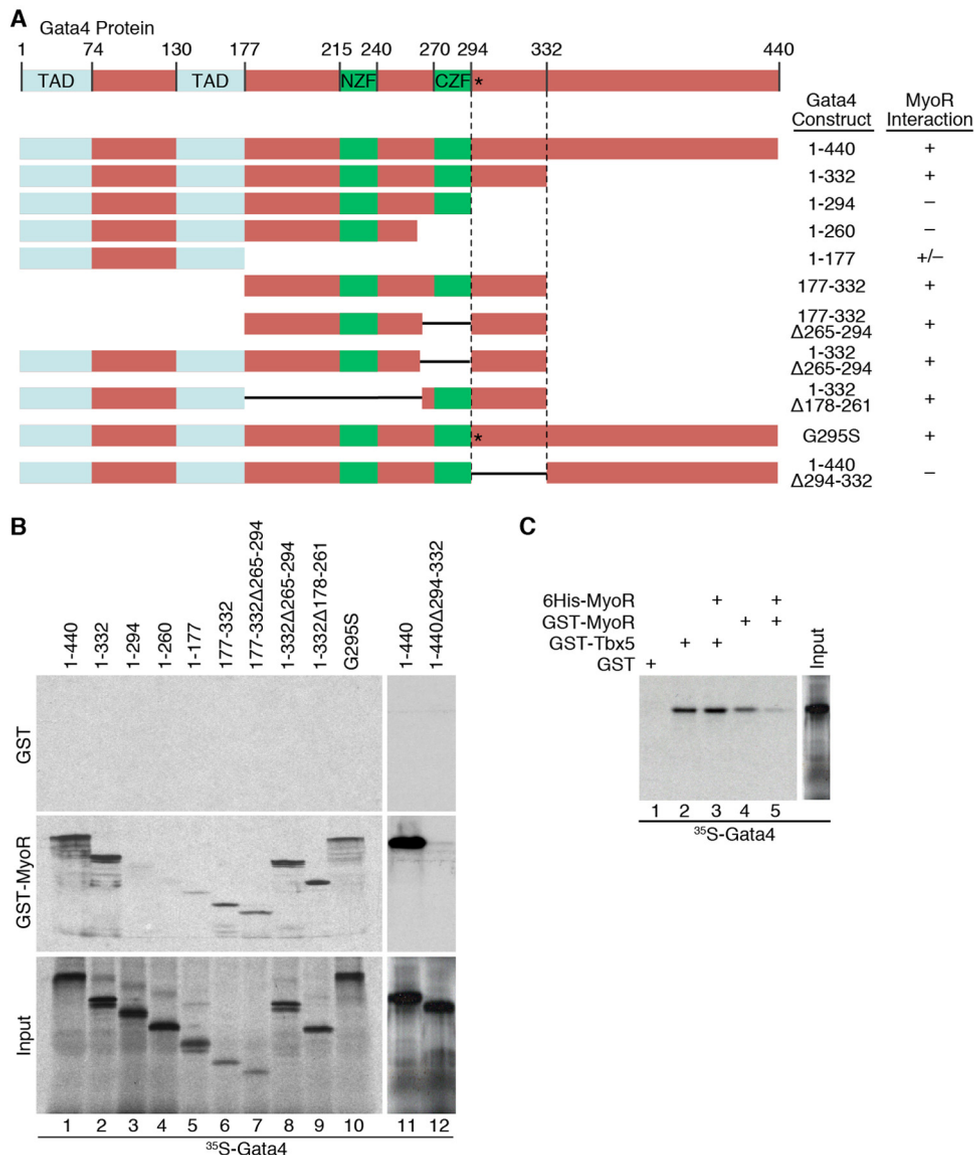


FIG 4 MyoR interacts directly with Gata4 adjacent to the CZF and separate from the Tbx5 binding surface. (A) Schematic diagram of the Gata4 deletion proteins used in GST pull-down experiments with summarized MyoR interaction data. (B) *In vitro*-translated ^{35}S -labeled Gata4 proteins were incubated with either GST alone (upper panel) or GST-MyoR (middle panel) to test for direct interaction. Input proteins are shown in the lower panel. Full-length Gata4 and deletions that preserve residues 294 to 332 interact directly with MyoR (lanes 1 and 2 and lanes 6 to 9). C-terminal deletion of Gata4 beyond amino acid 332 or internal deletion of residues 294 to 332 eliminates interaction with MyoR (lanes 3 to 5 and 12), showing that the region adjacent to the C-terminal zinc finger (CZF) is responsible for interacting with MyoR. In addition, the Gata4 G295S mutant retains the ability to interact with MyoR (lane 10), suggesting that MyoR and Tbx5 contact Gata4 via distinct binding surfaces. (C) GST pull-down competition assay with ^{35}S -labeled Gata4 interacting with GST, GST-Tbx5, or GST-MyoR in the presence or absence of unlabeled His-tagged MyoR recombinant protein (lanes 1 to 5). Although unlabeled MyoR competes efficiently with GST-MyoR for Gata4 interaction (lanes 4 and 5), it is unable to prevent binding between GST-Tbx5 and Gata4 (lanes 2 and 3), providing additional evidence that the binding of Gata4 to MyoR or Tbx5 is not mutually exclusive. TAD, transactivation domain; NZF, N-terminal zinc finger; CZF, C-terminal zinc finger.

that can activate and/or repress transcription (31). Hand1 and Hand2 are expressed in the developing heart and play crucial roles during ventricular morphogenesis (35). Although Hand2 binds Gata4 to cooperatively activate transcription on several cardiac promoters (37), Hand1 does not interact with Gata4 and has no effect on Gata4-dependent transcription (Fig. 2B). Since MyoR and Hand2 are bHLH family members with opposing effects on Gata4-dependent transcription, we could directly test additional characteristics of the MyoR repression domains. Thus, we generated chimeric MyoR-Hand2 proteins to assess (i) the ability of

both MyoR repression domains to function in a heterologous context and (ii) the functional equivalence of the two domains. Using the L8G5-Luc reporter, Gal4-MyoR potently inhibited transcription mediated by LexA-VP16, while Gal4-Hand2 augmented LexA-VP16 transcriptional activity (Fig. 5C), a finding consistent with the notion that Hand2 functions as a transcriptional activator (37). However, when the N terminus of MyoR (domain 1-100) was fused to the bHLH domain and C terminus of Hand2 (domain 97-214), this Gal4 fusion protein repressed LexA-VP16-dependent activation. Conversely, when the N terminus of Hand2

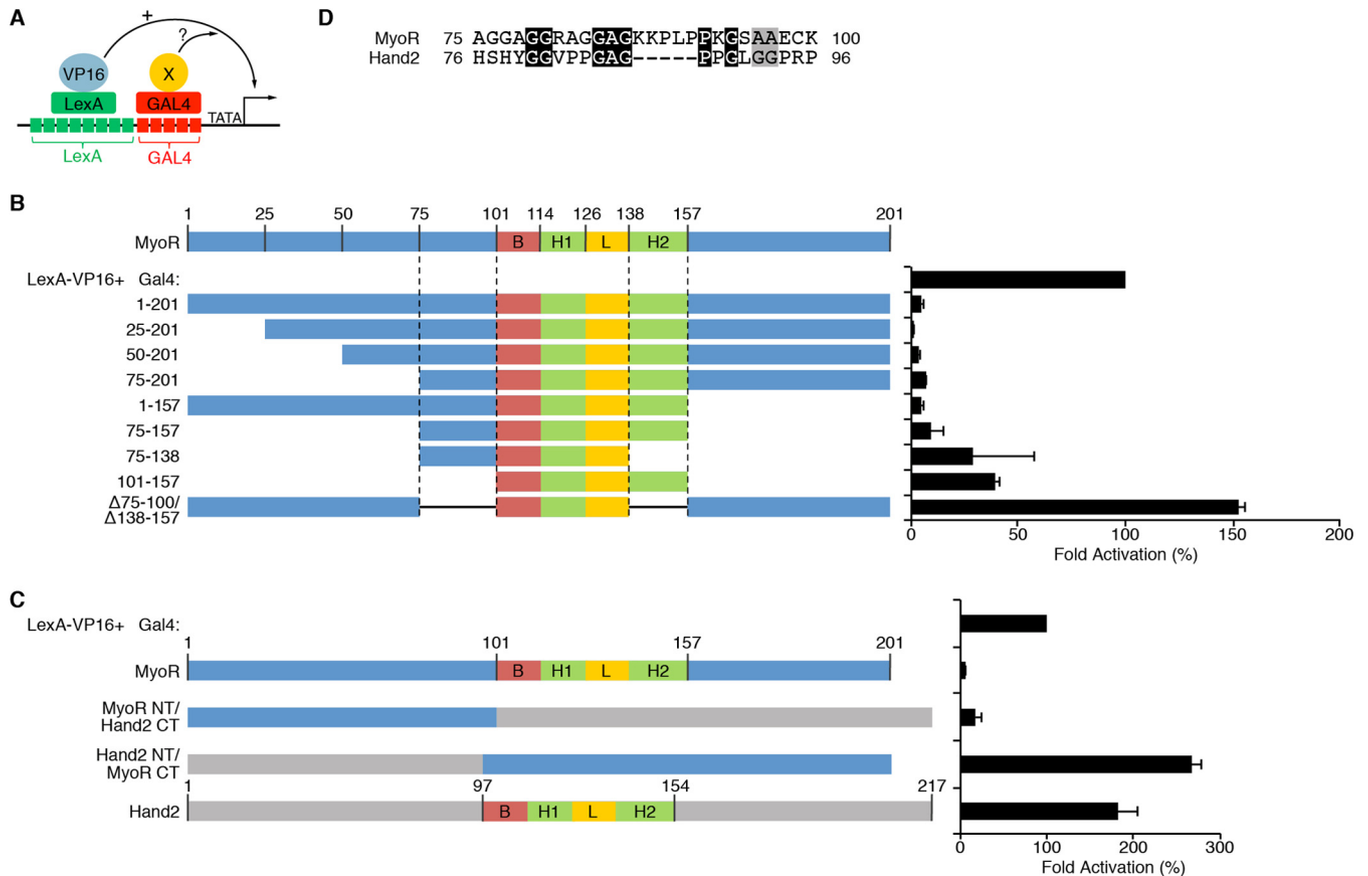


FIG 5 MyoR requires two nonequivalent repression domains to inhibit transcription. (A) Diagram depiction of the *trans*-repression assay system used for these experiments. The system utilizes a synthetic reporter construct containing eight LexA binding sites adjacent to five Gal4 binding sites. The LexA DNA-binding domain is fused with the HSV VP16 transactivation domain to mediate potent transcriptional activation. Candidate repressors are fused to the Gal4 DNA-binding domain and assessed for inhibitory activity in transient-transfection experiments. (B) A total of 100 ng of Lex8-Gal5-Luc was cotransfected with 100 ng of LexA-VP16 and 200 ng of the indicated Gal4-MyoR deletion construct. Also, 10 ng of CMV-lacZ was transfected for normalization purposes. The data are represented as the fold activation compared to LexA-VP16 alone (100%). N-terminal deletion to amino acid 75 or C-terminal deletion to residue 157 does not impair the ability of Gal4-MyoR to repress transcription by LexA-VP16. C-terminal deletion to residue 138 or N-terminal deletion to residue 100 partially impairs Gal4-MyoR repression. However, deletion of both putative repression domains (75-100 and 138-157) completely eliminates the ability of Gal4-MyoR to inhibit LexA-VP16-dependent transcriptional activation. (C) Assessment of repression activity by various Hand2-MyoR chimeric Gal4 constructs. As expected, Gal4-MyoR represses and Gal4-Hand2 activates LexA-VP16 mediated transcriptional activity. A Gal4 fusion protein containing MyoR 1-100, which includes the N-terminal repression domain (75-100), and the C terminus of Hand2, potentially repressed transcriptional activation mediated by LexA-VP16. In contrast, a Gal4 fusion protein containing the Hand2 N terminus fused with the MyoR C terminus, including the C-terminal repression domain, was unable to inhibit LexA-VP16 activation of reporter gene expression. (D) Protein alignment between amino acids 75 to 100 of MyoR and the corresponding region of Hand2 with conserved residues shaded black and similar residues shaded gray. Based on the alignment, MyoR contains a short peptide sequence (KKPLP) that is not present in Hand2.

(domain 1-97) was fused to the bHLH domain and C terminus of MyoR (domain 101-201), this Gal4 fusion protein augmented transcriptional activity mediated by LexA-VP16. These experiments suggest that the ability of MyoR and Hand2 to function as a repressor and activator, respectively, is determined by unique sequences contained within the N terminus of each protein. Given that residues 75 to 100 constitute the minimal repression domain within the MyoR N terminus, we searched for amino acids in this region that diverge from Hand2. Sequence alignment of this region between MyoR and Hand2 revealed several unique residues, including a Proline-rich motif (KKPLP) that is only present in MyoR (Fig. 5D). Collectively, these studies demonstrate that the MyoR N-terminal repression domain (domain 75-100), but not the C-terminal repression domain (domain 138-157), can function in a heterologous context and is sufficient to convert Hand2 into a repressor.

MyoR regulates Cx30.2 expression in vivo. Given that MyoR is enriched in the AVN (Fig. 1) and directly inhibits Gata4-dependent Cx30.2 expression in cell culture (Fig. 2B), we wished to evaluate whether MyoR regulates Cx30.2 expression *in vivo*. To address this question, we bred Cx30.2-lacZ transgenic mice with MyoR^{-/-} animals (41) to obtain MyoR^{+/-}; Cx30.2-lacZ/+ double-transgenic mice. Then, double-transgenic mice were bred with MyoR^{+/-} animals to generate MyoR^{+/+}; Cx30.2-lacZ/+ and MyoR^{-/-}; Cx30.2-lacZ/+ littermate mice to compare lacZ expression in a wild-type versus MyoR-null background. When we performed whole-mount X-Gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) staining on E16.5 hearts from these mice, we did not detect any obvious changes in the pattern of Cx30.2-lacZ expression (data not shown), demonstrating that MyoR does not alter the spatial distribution of Cx30.2-lacZ transcripts. However, this experiment did not address whether MyoR influences

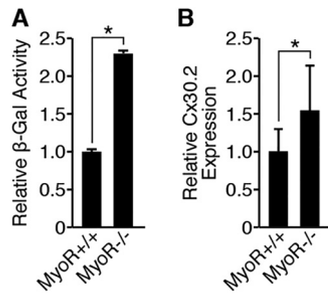


FIG 6 MyoR represses Cx30.2 expression *in vivo*. (A) The AVC region of MyoR^{+/+}; Cx30.2-lacZ/+ ($n = 3$) and MyoR^{-/-}; Cx30.2-lacZ/+ ($n = 2$) littermates was dissected at E16.5, and lysates were assessed for β -galactosidase activity as a quantitative measure of Cx30.2-lacZ expression. Enzyme activity is shown relative to MyoR^{+/+}; Cx30.2-lacZ/+ AVC levels. (B) Endogenous Cx30.2 expression was measured by qRT-PCR in the E16.5 AVC of MyoR^{+/+} ($n = 10$) and MyoR^{-/-} ($n = 10$) animals. Cx30.2 gene expression is shown relative to the AVC of MyoR^{+/+} mice. *, $P < 0.05$.

the absolute levels of Cx30.2-lacZ expression, since whole-mount X-Gal staining is qualitative when performed under saturating conditions. Therefore, we conducted colorimetric determination of β -galactosidase activity in the AVC region under linear assay conditions to obtain a more quantitative assessment of lacZ expression (23). Using this approach, we found that the AVC of MyoR^{-/-}; Cx30.2-lacZ/+ mice contained ~ 2 -fold more X-Gal activity compared to the AVC of MyoR^{+/+}; Cx30.2-lacZ/+ mice at E16.5 (Fig. 6A), an observation consistent with the notion that MyoR represses Cx30.2 enhancer activity *in vivo*. Although this experiment suggests that MyoR functions as an AVN-specific transcriptional repressor *in vivo*, it does not address whether MyoR directly inhibits endogenous Cx30.2 gene expression. Thus, we compared Cx30.2 expression levels in the AVC of MyoR-null mice compared to wild-type littermates (Fig. 6B). This experiment demonstrated that Cx30.2 expression was elevated in MyoR-null mice, although the results were not as dramatic as our findings with the Cx30.2-lacZ transgenic line. Taken together, these studies demonstrate that MyoR functions *in vivo* to repress Cx30.2 enhancer activity and endogenous Cx30.2 expression.

MyoR modulates cardiac electrical activity. We previously demonstrated that Gata4 heterozygous mice have shortened PR intervals as a result of reduced Cx30.2 expression, suggesting that a Gata4-dependent regulatory circuit regulates AV conduction delay (23). Given our results implicating MyoR in Gata4-dependent regulation of Cx30.2, we wished to determine whether MyoR also impacts AV conduction delay. Since MyoR represses Gata4, we hypothesized that MyoR-null mice would have a prolonged PR interval due to increased Cx30.2 expression; this would contrast with the shortened PR interval observed in mice with reduced Cx30.2 expression due to Gata4 haploinsufficiency (23) or genetic ablation (21). To test this hypothesis, we interbred MyoR^{+/-} mice to generate MyoR^{+/+} and MyoR^{-/-} mice for EKG analysis (Fig. 7A). From these experiments, we observed an increase in the PR interval of MyoR-null mice compared to wild-type littermates (39.6 ± 1.1 ms versus 44.6 ± 2.0 ms; $P < 0.05$) (Fig. 7B), thus indicating prolonged AV conduction. Importantly, none of the other EKG parameters, including heart rate (HR), QRS interval, and QT interval, differed significantly between the two groups of mice. Interestingly, one MyoR-null mouse exhibited more severe AV conduction defects, such as Mobitz I and Mobitz II second

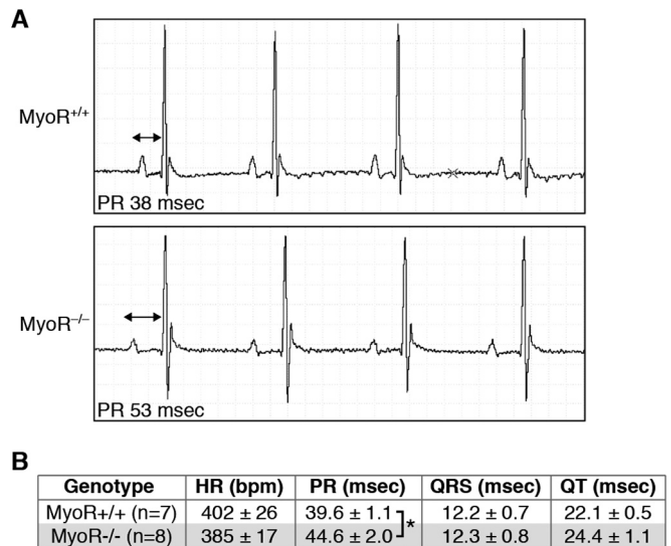


FIG 7 MyoR modulates AV delay. (A) EKG analysis was performed on 4-week-old MyoR^{+/+} and MyoR^{-/-} mice, and representative tracings from heart rate-matched littermates are shown. Note the mild prolongation in the PR interval (double-headed arrow) of MyoR^{-/-} mice compared to MyoR^{+/+} mice. (B) Summary of EKG analysis on MyoR^{+/+} ($n = 7$) and MyoR^{-/-} ($n = 8$) mice. The PR interval is prolonged in MyoR^{-/-} mice compared to MyoR^{+/+} mice, while the HR, QRS interval, and QT interval are similar between groups. *, $P < 0.05$.

degree AVB (data not shown), which were never observed in wild-type littermate mice. Collectively, these studies demonstrate that MyoR modulates AV conduction delay.

DISCUSSION

Here we show that MyoR is expressed in the developing AVN and represses Gata4-dependent activation of the Cx30.2 enhancer. We demonstrate that MyoR directly interacts with Gata4 via a protein domain that also binds to Tbx5, but each transcription factor utilizes a unique binding surface. We also show that MyoR contains two discrete repression domains, and the N-terminal repression domain can function in a heterologous context to convert Hand2 into a repressor. Consistent with these results, we find that genetic ablation of MyoR increases Cx30.2 expression and delays AV conduction, thus implicating MyoR in a Gata4-dependent transcriptional circuit that establishes normal cardiac rhythm.

Pharyngeal mesoderm links MyoR to heart development. Previous studies have demonstrated that deletion of MyoR and capsulin results in complete absence of specific muscles of mastication (41). Consistent with this phenotype, MyoR is developmentally expressed in splanchnic pharyngeal mesoderm (SPM), which contains cranial skeletal muscle precursor cells (47–49). Interestingly, the SPM also contains the second heart field (SHF), which contributes cells to the inflow tract (IFT), SAN, atria, right ventricle, outflow tract, and AVN (50). In fact, mounting experimental evidence supports the novel idea that a common subset of precursor cells within the SPM contributes to both the muscles of mastication and SHF-derived structures within the heart (47). Interestingly, MyoR transcripts have been previously detected in the SHF-derived sinus venosus/IFT region of the developing chick heart (51). In the present study, we provide evidence that MyoR functions within the mouse AVN, a structure that is also derived in

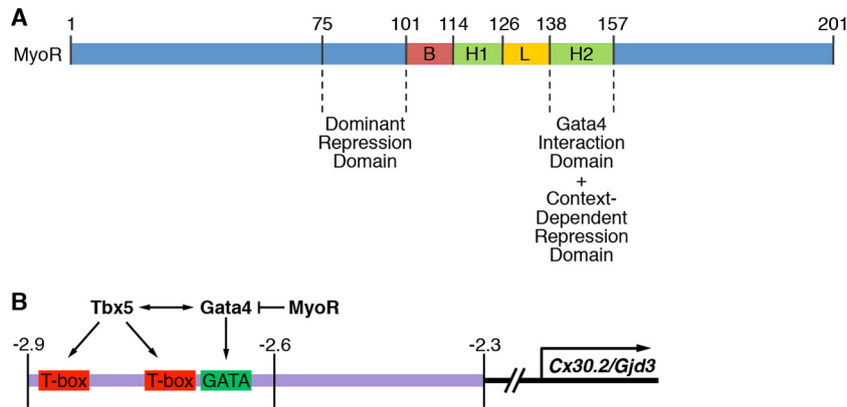


FIG 8 Model for MyoR-dependent regulation of Cx30.2 expression. (A) Summary of MyoR functional domains identified in the present study. Residues 138 to 157 serve as a Gata4 interaction and context-dependent repression domain. The domain encompassing amino acids 75 to 100 functions as a *trans*-repression domain even in a heterologous context. (B) Tbx5 and Gata4 regulate Cx30.2 expression via a 0.6-kb minimal regulatory element that directs expression to the AVN. Based on the present study, we propose that MyoR functions within AV nodal cells as a Gata4-specific transcriptional repressor to decrease Cx30.2 gene expression and thus modulate AV delay. Given the results of our protein-protein interaction experiments, we conclude that MyoR does not inhibit transcriptional activation by competing with Tbx5 for Gata4 interaction. Instead, we speculate that MyoR possesses a *trans*-repression domain that recruits as-yet unidentified corepressors to inhibit Gata4- and Tbx5-dependent activation of Cx30.2 expression.

part from the SHF. Perhaps not unexpectedly, MyoR thus functions within two distinct derivatives of the SPM, the muscles of mastication and the AVN. The implication for such functional duality is that MyoR must be capable of executing unique tissue-specific transcriptional programs depending upon the developmental context.

Individual transcription factors can potentially activate multiple gene expression programs by cooperating with tissue-specific cofactors. Although Gata4 is abundant within the heart, for example, it is not known to be expressed in the developing facial musculature. Therefore, we postulate that Gata4 serves as a heart-specific cofactor that directs MyoR-dependent gene regulation within the heart. Likewise, MyoR may serve as a Gata4 cofactor to distinguish the gene expression signature of the AVN from other regions of the heart. Previous studies have established that cardiomyocyte specification and organogenesis rely upon a densely interconnected regulatory network of cardiac transcription factors (i.e., Gata4, Mef2c, and Tbx5) (52). Given that these transcription factors are widely expressed in the heart, however, it remains unclear how individual transcription factors modulate gene expression programs within specific cardiac cell types. Our findings provide support for a model in which Gata4 serves as a critical node during heart development upon which additional cofactors act to impact lineage-specific gene expression (24). Indeed, Gata4 interacts with several transcription factors (Hand2 and MyoR) and coactivators (Fog1, Fog2, and CBP/p300) that directly influence various aspects of cardiac morphogenesis and gene expression. Based on the results of our experiments, we postulate that MyoR and Hand2 form an antagonistic pair of Gata4 cofactors that modulate cell type-specific gene expression depending upon their relative abundance. Similarly, we speculate that additional antagonistic pairs of transcriptional cofactors may fine-tune Gata4-dependent transcription in various subdomains within the developing heart.

Transcriptional repression and cardiac conduction. Multiple transcriptional repressors have been shown to influence formation and function of the conduction system. For example, Tbx2 and Tbx3 are expressed within the central conduction system to re-

press working cardiomyocyte-specific genes (11, 12). To accomplish this function, they interact with Nkx2.5 and prevent transcriptional activation of the chamber-specific genes *Nppa* and *Cx43* to reinforce conduction cell identity. In a similar fashion, the SAN-specific repressor *Shox2* inhibits Nkx2.5-dependent activation of atrial gene expression (53). The AVC repressors *Msx1* and *Msx2* repress chamber-specific *Cx43* expression (54); similarly, *Irx3* is expressed in the AV bundle and represses *Cx43* expression (55). Collectively, these examples highlight a common theme by which tissue-specific transcriptional repressors prevent chamber-specific gene expression within the conduction system. In contrast, the transcriptional repressor *Id2* functions within AV bundle cells to regulate cell cycle exit rather than to repress chamber-specific gene expression (56). Thus, cell cycle regulation is a second method by which transcriptional repressors influence conduction system formation. Here we demonstrate that the transcriptional repressor MyoR fine-tunes *Cx30.2* expression within the AVN. Thus, MyoR-dependent modulation of a gap junction protein defines a third strategy for transcriptional repressors to influence conduction system function. These repressive strategies are unlikely to be mutually exclusive, however, so we cannot exclude the possibility that MyoR also affects cardiac conduction by additional mechanisms. Taken together, we propose that transcriptional repressors can modulate cardiac conduction by three distinct mechanisms: (i) repression of chamber-specific genes, (ii) regulation of cell cycle exit, and (iii) modulation of gap junction protein levels.

Previous studies have established that some bHLH repressors inhibit transcription by interacting with histone deacetylase-containing corepressor complexes (57), such as N-Cor/SMRT, NuRD, and Co-REST, while others function in a dominant-negative fashion (58). We demonstrate here that MyoR interacts with Gata4 and inhibits transcription via two separable, but non-equivalent, repression domains (Fig. 8). Interestingly, the N-terminal repression domain can function in a heterologous context, while the C-terminal repression domain cannot. Moreover, the N terminus of MyoR converts Hand2 into a repressor, while the N terminus of Hand2 converts MyoR into an activator. Thus, these

findings support the idea that the ability of MyoR and Hand2 to function as a repressor and activator, respectively, is entirely dictated by the impact of their N terminus on transcription. Consistent with this idea, we identified a proline-rich sequence (KKPLP) within the N-terminal repression domain of MyoR that is absent from Hand2 (Fig. 5D). Given that SH3 domains can bind to proline-rich motifs (59, 60), we searched the mouse genome for histone-modifying proteins that contain an SH3 domain and identified a single protein, PRMT2. However, we were unable to demonstrate that PRMT2 potentiates MyoR-dependent repression (J. P. Harris and N. V. Munshi, unpublished data), and the precise mechanism by which MyoR inhibits transcription remains unknown. Since the C-terminal repression (H2) domain is required when deleted from MyoR but is unable to inhibit transcription when transferred to Hand2, our studies suggest that this domain is necessary for repression only in a specific context. To reconcile these findings, we speculate that the MyoR H2 domain indirectly influences transcription by mediating a structural role that can be fulfilled by the H2 domain of another bHLH family member such as Hand2.

MyoR as a candidate quantitative trait locus (QTL). Previous twin studies have provided strong evidence that cardiac conduction is a heritable trait (61). Based on the finding that EKG parameters are more correlated in monozygotic versus dizygotic twins, it has been estimated that 34% of the PR interval is heritable. Although AV delay has been correlated with specific mutations in transcription factors, ion channels, and gap junction proteins, it remains unclear whether such mutations account for a significant proportion of PR interval variability in the general population (2). More recently, genomewide association studies have identified numerous single nucleotide polymorphisms (SNPs) that associate with PR interval length in the general population (62, 63). Although the influence of each individual SNP is modest, the overall impact across the entire population is predicted to be significant. In other words, PR interval is a continuous variable that is likely to be shaped by the contribution of multiple QTLs with modest individual effects. From our experimental results, it is clear that MyoR is not required for AVN formation or function *per se*; indeed, the PR interval of MyoR knockout mice is only ~13% longer than wild-type mice. Rather, we suggest that MyoR contributes to the PR interval in mice by modulating AV conduction in concert with a collection of additional QTLs such as Tnni3k (64). Since mutations in MyoR would not be predicted to significantly impact fitness, SNPs within the MyoR locus could contribute to PR interval variability in the general human population. Especially given the results of recent large-scale whole-exome sequencing efforts in diverse human populations (65, 66), we anticipate that individuals with homozygous loss-of-function MyoR mutations are likely to exist. Since Cx31.9, the human ortholog of mouse Cx30.2, has not been detected in the human cardiac conduction system (67), we predict that MyoR operates through additional conserved pathways to impact PR interval variability in humans. Although the validation of these hypotheses must await the accumulation of large sequencing data sets, the search for additional QTLs that contribute to PR interval variability in the general population will remain an area of active investigation.

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