Expression of Sumoylation Deficient Nkx2.5 Mutant in Nkx2.5 Haploinsufficient Mice Leads to Congenital Heart Defects

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Abstract

Nkx2.5 is a cardiac specific homeobox gene critical for normal heart development. We previously identified Nkx2.5 as a target of sumoylation, a posttranslational modification implicated in a variety of cellular activities. Sumoylation enhanced Nkx2.5 activity via covalent attachment to the lysine residue 51, the primary SUMO acceptor site. However, how sumoylation regulates the activity of Nkx2.5 in vivo remains unknown. We generated transgenic mice overexpressing sumoylation deficient mutant K51R (conversion of lysine 51 to arginine) specifically in mouse hearts under the control of cardiac α-myosin heavy chain (α-MHC) promoter (K51R-Tg). Expression of the Nkx2.5 mutant transgene in the wild type murine hearts did not result in any overt cardiac phenotype. However, in the presence of Nkx2.5 haploinsufficiency, cardiomyocyte-specific expression of the Nkx2.5 K51R mutant led to congenital heart diseases (CHDs), accompanied with decreased cardiomyocyte proliferation. Also, a number of human CHDs-associated Nkx2.5 mutants exhibited aberrant sumoylation. Our work demonstrates that altered sumoylation status may underlie the development of human CHDs associated with Nkx2.5 mutants.

Introduction

Nkx2.5, a cardiac specific homeobox gene, is one of the earliest markers for cardiac progenitor cells and is central for normal fetal cardiac structural morphogenesis. Nkx2.5 null mice died from cardiac malformations [1,2], while mice with Nkx2.5 haploinsufficiency caused atrial septal defects (ASDs), one of the most common forms of congenital heart diseases (CHDs) in humans, although the penetrance was low and was influenced by the mouse strain [3,4]. The critical role of Nkx2.5 in the conduction system was revealed by observations in both human patients with heterozygous Nkx2.5 mutants and in animal models [5,6]. Overexpressing wild type Nkx2.5 or one of its homeodomain (HD) mutants such as I183P in murine cardiomyocytes under the control of β-myosin heavy chain promoter resulted in embryonic lethality and/or heart failure, which was a consequence of dilated cardiomyopathy and/or a defective conduction system [7]. Transgene expression of the I183P mutant, driven by α-myosin heavy chain promoter (α-MHC), also led to cardiac dysfunction, but with later onset [8] and with phenotypic discrepancy [8,9]. Postnatal cardiac specific ablation of Nkx2.5 by tamoxifen-inducible Cre expression starting at 2 weeks of age led to a conduction defect and dilated cardiomyopathy [10]. In the clinic, a multitude of heterozygous Nkx2.5 mutants were shown to be etiologically implicated in nonsyndromic CHDs [11]. Thus, Nkx2.5 has an essential role in the formation of cardiac structure and in the maintenance of cardiac function.

Nkx2.5 is a moderate activator of its target genes. In addition to its physiological activity achieved via DNA binding through its HD, it functioned through associations with other transcription factors such as SRF, GATA4 and TBX proteins [12,13,14,15,16,17]. It is also noteworthy that the transcriptional activity of Nkx2.5 was modulated in part by posttranslational modification such as phosphorylation by casein kinase II (CKII) and sumoylation [18,19]. Sumoylation is a posttranslational modification in which the SUMO moiety is covalently and reversibly conjugated to the lysine residue(s) located in the target protein. Sumoylation on lysine 51 of the Nkx2.5 protein substantially increased its activity, and mutation of lysine 51 to arginine (K51R) suppressed SUMO linkage and the activity of Nkx2.5 [18]. However, how the sumoylation deficient Nkx2.5 mutant behaves in vivo is unknown.

Recent studies pointed to an important role of SUMO conjugation in cardiac structural morphogenesis and dysfunction [20,21]. SUMO-1 knockout (KO) mice exhibited cardiac structural defects, although the phenotypic penetrance may have been affected by genetic background [22]. Nkx2.5 played a role in the development of SUMO-1 KO-induced CHDs, because the compound Nkx2.5+/−/SUMO-1−/− mice exhibited more severe...
cardiac phenotypes than the mice with single heterozygous knockout of either Nkx2.5 or SUMO-1 [22]. Given the above findings and the effect of sumoylation on transcriptional activity of Nkx2.5 [18], we asked whether CHD-linked Nkx2.5 mutants exhibited abnormal sumoylation activity and how defective sumoylation of Nkx2.5 affected cardiogenesis in vivo. We uncovered that sumoylation was decreased in a number of naturally occurring Nkx2.5 mutants, including those HD mutants tested, and that cardiac restricted expression of the sumoylation deficient mutant, K51R, in the Nkx2.5 haploinsufficient (Nkx2.5<sup>−/−</sup>) mice led to CHDs.

**Materials and Methods**

**Ethics Statement**

The animals were handled in accordance with institutional guidelines with appropriate approvals (protocol ID 09018) by IACUC at Texas A&M Health Science Center.

**Generation of transgenic and knockout mice**

The human influenza hemagglutinin (HA) tagged K51R mutant was generated as reported [18] and then was subcloned between the 5.4 kb mouse α-MHC promoter (provided by Dr. J. Robbins, University of Cincinnati) and the Simian virus 40 (SV40) polyadenylation sequence via Sall sites. The orientation of the inserted transgene was confirmed by DNA sequencing. The construct was microinjected into the pronucleus of fertilized eggs from FVB mice, and offspring were screened by polymerase chain reaction (PCR). One mouse line was identified to transmit detectable transgene. F0 founder was cross back to C57BL/6 and scored for positive transmission revealed by PCR using the 5.4 kb mouse α-MHC promoter driven luciferase reporter construct. The cardiac α-actin promoter driven luciferase reporter construct (Ca-actin-Luc), the expression vector encoding HA-tagged wild type Nkx2.5 and flag-tagged SUMO-1 was detailed previously [18,24]. For in vivo sumoylation assay, the expression vectors for wild type Nkx2.5 or Nkx2.5 mutants were transfected into Hela cells cultured on 6 cm plates in the presence or absence of encoding vector for flag-tagged SUMO-1. Whole cell lysates were purified 48 hours after transfection in the presence of 25 mM isopeptidase inhibitor N-ethylmaleimide (NEM), which prevents desumoylation of those SUMO-conjugated proteins.

**Antibodies**

Antibodies against Nkx2.5, HA and GAPDH were purchased from Santa Cruz Biotechnology, Inc., and anti-flag antibody (M2) was from Sigma.

**Plasmid construction, cell culture, transient transfection and western blot**

The cardiac α-actin promoter driven luciferase reporter construct (Ca-actin-Luc), the expression vector encoding HA-tagged wild type Nkx2.5 and flag-tagged SUMO-1 was detailed previously [13,24]. The human influenza hemagglutinin (HA) tagged K51R mutant was generated as reported [18] and then was subcloned between the 5.4 kb mouse α-MHC promoter and the Simian virus 40 (SV40) polyadenylation sequence via Sall sites. The orientation of the inserted transgene was confirmed by DNA sequencing. The construct was microinjected into the pronucleus of fertilized eggs from FVB mice, and offspring were screened by polymerase chain reaction (PCR). One mouse line was identified to transmit detectable transgene. F0 founder was cross back to C57BL/6 and scored for positive transmission revealed by PCR using the 5.4 kb mouse α-MHC promoter driven luciferase reporter construct. The cardiac α-actin promoter driven luciferase reporter construct (Ca-actin-Luc), the expression vector encoding HA-tagged wild type Nkx2.5 and flag-tagged SUMO-1 was detailed previously [18,24]. For in vivo sumoylation assay, the expression vectors for wild type Nkx2.5 or Nkx2.5 mutants were transfected into Hela cells cultured on 6 cm plates in the presence or absence of encoding vector for flag-tagged SUMO-1. Whole cell lysates were purified 48 hours after transfection in the presence of 25 mM isopeptidase inhibitor N-ethylmaleimide (NEM), which prevents desumoylation of those SUMO-conjugated proteins. 40 µg of prepared proteins from each group were subsequently analyzed in NuPage SDS gel, transferred to PVDF membrane, which was probed with desired antibodies and revealed by chemiluminescence.

**Histopathology**

Mouse neonatal hearts were dissected and fixed in 4% paraformaldehyde (PFA) at 4°C for overnight. Hearts were sectioned transversely at 10 µm thickness, which were stained with Hematoxylin and Eosin (H&E) according to the standard protocol.

**Immunohistochemistry**

The mouse hearts at P7 were fixed in 10% formalin prepared in 1× phosphate-buffered saline (PBS), dehydrated and embedded in paraffin for histological sections. For antigen retrieval, 5 µM
sections were boiled in sodium citrate buffer (10 mM, PH6.0) for 10 minutes, and then blocked in 2% bovine serum albumin (Vector Laboratories, SP-3050/PBST (1XPBS/0.05% Tween20), followed by incubation with the primary antibodies monoclonal anti-cardiac troponin T (cTnT, 1:200, Lab Vision, MS-295-P) and rabbit anti-Ki67 (1:200, Abcam, ab15580) in blocking solution at 4°C overnight. Sections were then washed, followed by incubation with 1:500 Alexa fluor® 594 goat anti-rabbit secondary antibody (Invitrogen, A31632) and Alexa fluor® 488 donkey anti-mouse secondary antibody (Invitrogen, A21202) in blocking solution at RT for 30 minutes. Sections were mounted with Vectashield® with DAPI (Vector Laboratory) and photographed under a Leica fluorescence microscope. TUNEL staining was conducted according to manufacturer’s protocol (Promega). The number of double Ki67 positive or TUNEL positive and cTnT positive cells (Ki67+/cTnT+ or TUNEL/cTnT+) was scored in at least three randomly selected fields in the left ventricle or atrioventricular septum and expressed as mean ± SEM.

**Echocardiography**

Two dimensional M-mode of Vevo 770 (Visual Sonics, Toronto, Canada) was used to measure the cardiac function on mice of interest with various ages as designed. Briefly, mice were anesthetized by inhalation of 1% isofluorane and fixed to a warm pad. Chest hair was removed and contacted with the probe to record cardiac function indices. Evaluations of cardiac function and heart dimensions were performed by an expert analyst blinded to the genotype of animals.

**Reverse transcription-quantitative PCR (RT-qPCR)**

RNAs were extracted from K51R-Tg and age-matched control mouse hearts using Trizol based on manufacturer’s protocol. 1 or 5 μg total RNAs were used for reverse transcription reaction using cloned reverse transciptase (Invitrogen), followed by quantitative PCR on machine MX3000 (Strategene) using specifically designed cloned reverse transcriptase (Invitrogen), followed by quantitative PCR on machine MX3000 (Strategene) using specifically designed primers for the particular genes of our interest. The sequences (5’ to 3’) of oligos used in this study were shown below: ANF: forward, ggctcaacagactgtgatgtgt; reverse, gctgtagcctgtgctgtcattc; cardiac α-actin [26] : forward, ctcctctccaggtgaatgt; reverse, gcctatcctgctgtcattc; cardiac α-MHC: forward, ggcacaacgcgcagagag; reverse, tgtgtaggagatctctgcgt; cardiac β-MHC: forward, ggcaaanacatctgctgtc; reverse, tcggagctgacagagag; skeletal α-actin [29] : forward, agagacatgtgccacaaggaag; reverse, cgttcctgccaatcctcg; SERCA forward,aatggcacttctgctgc; reverse, tagtattactactcctc; connexin 40: forward, atctcccaactgctgct; reverse, aggagaagctcctgaggg; GAPDH: forward, atgttcaggtgtaatctg; reverse, gaagacacaggtagacac.

**Statistical analysis**

The unpaired Student’s t test, Chi-square test was applied to determine statistical significance between groups when applicable and shown in each Figure legend.

**Results**

CHDs-linked Nkx2.5 mutants displayed aberrant SUMO modification

As an initial step to investigate whether altered sumoylation status underpinned those Nkx2.5 mutants that were associated with CHDs, sumoylation assay using Ni²⁺-NTA chromatography was performed on Hela cell lysates expressing His₆-tagged SUMO-1 and Nkx2.5 wild type (wt) or one of seven point mutants created on murine Nkx2.5 protein, R25C, T177M, N187K, R188G, Y190C, C265Y and V309L (Fig. 1A). These point mutants corresponded to the seven missense human Nkx2.5 mutants respectively that were previously described and were involved in human CHDs [5,21]: R25C: T178M, N188K, R189G, Y191C, C270Y and V315L. As shown in Fig. 1B, four Nkx2.5 HD mutants tested, T177M, N187K, R189G, Y190C, showed significant decrease in sumoylation, while R25C still exhibited robust sumoylation. To better evaluate the changes in sumoylation activity of Nkx2.5 mutants vs. wt, western blots, in which loading controls were better monitored, were carried out on cell lysates purified from transfected Hela cells expressing Nkx2.5 wt or one of its mutants in the absence or presence of flag-epitoped SUMO-1, as indicated in Fig. 1C. Also, the relative level of sumoylated Nkx2.5 wt or mutants after normalization to the level of corresponding free Nkx2.5 protein was objectively determined using software ImageJ (Fig. 1D). While Nkx2.5 wt and those mutants tested were expressed at comparable levels, N187K and R188G abolished over 80% sumoylation compared with Nkx2.5 wt, and T177M, Y190C and V309L blocked ~50% sumoylation (Fig. 1C&D). Also, C265Y showed ~30% decrease in SUMO linkage. Among these mutants, R25C appeared to have a normal sumoylation activity, however, this “normal” sumoylation status of R25C was achieved with much lower expression level of SUMO-1 (Fig. 1C, right panel), indicative of R25C’s high sumoylation affinity.

Since SUMO conjugation promoted the capacity of Nkx2.5 to activate target gene promoters [18], we next compared the transcriptional activities of Nkx2.5 wt and those mutants in the presence of SUMO-1. Luciferase activity assays were performed on Hela cell lysates expressing Ca-actin-Luc along with Nkx2.5 wt or one of those above-mentioned mutants in the absence or presence of increasing dosages of SUMO-1 expression vector as indicated (Fig. 1E). In line with the observation from sumoylation assays (Fig. 1C&D), SUMO-1 substantially increased the capacity of Nkx2.5 wt to activate target gene promoter up to ~120 fold, but this increase was impaired by the HD mutations. C309L mutant also failed to fully respond to SUMO-1. SUMO-1 enhanced the transcriptional activity of Nkx2.5 wt and those mutants in the presence of increasing dosages of SUMO-1 expression vector as indicated (Fig. 1C&D). While Nkx2.5 wt and those mutants tested were expressed at comparable levels, N187K and R188G abolished over 80% sumoylation compared with Nkx2.5 wt, and T177M, Y190C and V309L blocked ~50% sumoylation (Fig. 1C&D).

**Transgene expression of sumoylation deficient Nkx2.5 K51R mutant in the murine heart**

Since a number of CHDs-associated Nkx2.5 mutants showed defective sumoylation, we asked whether sumoylation deficient Nkx2.5 mutant K51R [18] could be causally linked to CHDs once expressed in vivo. We generated cardiomyocyte specific expression of HA-tagged K51R transgene (K51R-Tg) under the control of cardiac α-MHC promoter (Fig. 2A). One founder of K51R-Tg mouse transmitted detectable HA-epitoped K51R transgene in murine hearts (Fig. 2B). The expression level of HA-tagged Nkx2.5 was approximately 1.5~2 folds compared with that of Nkx2.5 protein in the wild type mouse heart, as determined by protein blot using a specific antibody against Nkx2.5 (Fig. 2C). The offspring from the cross between K51R-Tg and wild type C57BL/6 mice were born at the expected Mendelian ratio (~50%), indicating no
embryonic loss. Except that two unusual premature deaths occurred at postnatal day (P) 2 and 9 (P2 and P9), which carried atrial septal defect (ASD) (data not shown), all other K51R-Tg mice appeared normal and fertile.

To gain an insight into cardiac functions in the mice bearing the K51R mutant, echocardiography was performed on K51R-Tg and wild type mice at ages of P50 and P300, respectively, using Vevo 770. Surprisingly, overexpression of the sumoylation...
showed enlarged hearts from K51R-Tg/Nkx2.5/−/− mice (Fig. 4, compare E with B, C and D), with both atria gluted with blood, indicating heart failure. The subsequent histological analysis revealed the presence of compound ASD/VSD in the K51R-Tg/Nkx2.5/−/− mouse hearts (compare E’ with B’, C’ and D’) in 10 out of 11 demised K51R-Tg/Nkx2.5/−/− mice examined, and a single ASD in 1 mouse. Dead pups from other genotype groups showed either normal structure (Nkx2.5/−/− and K1R-Tg, respectively) or a single ASD (wt) (Fig. 4F). Thus, expression of the transgene K51R in the existence of Nkx2.5 haploinsufficiency caused CHDs, which was likely the primary cause of premature death observed in the compound K51R-Tg/Nkx2.5/−/− mice.

Decreased cardiomyocyte proliferation in the compound K51R-Tg/Nkx2.5/−/− mouse hearts

Cardiac structural phenotypes such as ASDs and VSDs can be associated with decreased proliferation and/or increased apoptosis of cardiomyocytes [33,34,35]. To explore the molecular basis underlying the CHDs present in the compound K51R-Tg/Nkx2.5/−/− mice, cardiomyocyte proliferation in the heart sections from age-matched wt, K1R-Tg, Nkx2.5/−/− and compound K51R-Tg/Nkx2.5/−/− mice (n = 3 for each group) at P7 was first assessed using antibodies against Ki67, a cell proliferating marker [36], and cardiac tropinin-T (cTnT), a cardiomyocyte marker [37]. As shown in Figure 5, A-D, the number of proliferating cardiomyocytes (double Ki67+/cTnT+) in both left ventricle and atrioventricular septum of the compound K51R-Tg/Nkx2.5/−/− mice was substantially lower than those in other three genotype animals (number/field: K51R-Tg/Nkx2.5/−/− 2.0±0.67 vs. wt 10.2±1.03, Nkx2.5/−/− 9.3±0.95, and K1R-Tg 9.5±0.85, p<0.0001, and data not shown). Next, the same tissue sections mentioned above were stained with TUNEL reagent. The rates of double TUNEL+/cTnT+ positive cardiomyocytes were similar between these four genotypes (Fig. 5 F-G, 1–2 cells/field, p>0.05). Collectively, these findings suggested that defective cardiomyocyte proliferation but not altered apoptosis underpinned CHDs present in the compound K51R-Tg/Nkx2.5/−/− mice.

Two out of four surviving compound K51R-Tg/Nkx2.5/−/− mice examined exhibited dilated cardiomyopathy

Although ~65% of the compound K51R-Tg/Nkx2.5/−/− mice died prematurely, ~35% of them survived to adulthood. We then asked whether these mice exhibited cardiac dysfunctions. Non-invasive cardiac function analysis was carried out on surviving progenies at approximately 4 months age obtained from the cross between K51R-Tg and Nkx2.5/−/− mice. As shown in Table 1, the compound K51R-Tg/Nkx2.5/−/− mice exhibited an increase in LV volume during systole (33.08±8.42 μl vs. mice of wt 15.50±9.08 μl, Nkx2.5/−/− 13.78±5.67 μl, and K1R-Tg 12.32±0.83 μl, respectively), and during diastole (64.95±10.93 μl vs. mice of wt 47.75±12.68 μl, Nkx2.5/−/− 34.50±7.25 μl, and K1R-Tg 48.47±1.38 μl, respectively). Also, compound K51R-Tg and Nkx2.5/−/− mice showed a decrease in cardiac contraction capacity as indicated by both percent ejection fraction (K51R-Tg/Nkx2.5/−/− 51.68±7.53 vs. wt 72.32±9.33, Nkx2.5/−/− 65.97±9.59, and K1R-Tg 74.45±2.3, respectively) and percent fractional shortening (K51R-Tg/Nkx2.5/−/− 26.59±4.96 vs. wt 41.75±7.26, Nkx2.5/−/− 36.84±7.13, and K1R-Tg 42.45±2.14, respectively). However, the difference of the above-mentioned indexes between these groups was not statistically significant due to variations among individual K51R-Tg/Nkx2.5/−/− mice examined. Further elaborated interrogation revealed enlarged left ventricular chamber with impaired cardiac functions in two of four compound K51R-Tg/
Figure 3. Dysregulation of cardiac genes in K51R-Tg mice. RT-qPCR revealed altered transcription of some cardiac genes examined such as BNP and myosin light chain (MLC)-2v in K51R-Tg hearts (n = 6) compared with those of control hearts (n = 4) at P50. RT-qPCR was performed as described in Materials and Methods. Unpaired student’s t test was used for statistically significant analysis. doi:10.1371/journal.pone.0020803.g003

Figure 4. Compound K51R-Tg/Nkx2.5+/– mice exhibited ASD and/or VSD. A. Mortality rate of compound K51R-Tg/Nkx2.5+/– mice was significantly higher than those of other genotype mice. The data were collected from 65 animals of 7 litters. The number shown above each bar indicated the mortality rate of that corresponding group. B–E, Representative gross view of heart morphologies in littermate wt, Nkx2.5+/–, K51R-Tg and compound K51R-Tg/Nkx2.5+/– mice at P4 was shown. Note that the atria of the compound K51R-Tg/Nkx2.5+/– mouse heart were filled with blood. A’–D’, H&E staining on 10 μm sections of hearts corresponding to A–D was shown. Arrows in D’ indicated ASD and VSD, respectively. Scale bar, 100 μm. F. Incidence of cardiac defects in the demised mice from four genotype groups. Chi square test was used for statistical analysis. **, p<0.01, vs. other groups. RA, right atrium; LA, left atrium; RV, right ventricle; LV, left ventricle; ASD, atrial septal defect; VSD, ventricular septal defect. doi:10.1371/journal.pone.0020803.g004
Nkx2.5+/− mice examined (Fig. 6), suggesting dilated cardiomyopathy, although the size of the sample needs to be expanded to reach a statistical significance.

Discussion

Nkx2.5 is a key transcription factor for embryonic cardiogenesis and its naturally occurring mutations are causally linked to human cardiac anomalies. Based on our previous findings that Nkx2.5 was a sumoylation substrate [18], in the present study we investigated the effects of CHDs-associated Nkx2.5 missense mutations on Nkx2.5 sumoylation and the impact of accumulated unsumoylatable Nkx2.5 mutant K51R on murine heart development and function.

CHDs-associated Nkx2.5 mutants and sumoylation activity

The present study revealed that all four Nkx2.5 HD mutants and one C-terminal domain mutant C309L tested rendered either substantial or intermediate level of decrease in SUMO conjugation, respectively, while the intact principle SUMO site was existent, indicating the involvement of these sites in the regulation of Nkx2.5 sumoylation. The similar phenomena of modulation of sumoylation by non-sumoylation site(s) have been observed in other SUMO targets [38]. The diminished capability of these Nkx2.5 mutants to be sumoylated was also reflected in the target promoter activity assays, in which the presence of SUMO-1 failed to fully potentiate the activity of any of those sumoylation-deficient mutants compared with that of wild type Nkx2.5. On the other hand, the R25C mutant showed no significant changes in SUMO modification. However, it is noteworthy that the mutant R25C exhibited higher sensitivity to SUMO conjugation compared with wild type Nkx2.5, because the lower expression of SUMO-1 protein induced its sumoylation to the level equivalent to that of wild type Nkx2.5 with higher expression of SUMO protein. This increased sensitivity of R25C to sumoylation probably accounted for the higher activation of target gene promoter by R25C, possibly due to active endogenous sumoylation machinery in Hela cells. And, if this is the case, then the addition of exogenous SUMO-1 would do little to further increase its activity. However, this hypothesis needs further investigation. The higher transcriptional activity of R25C mutant observed in the present study was not observed in a previous report [5]. This discrepancy was likely attributable to the different systems in which the experiments were conducted. Increased sumoylation sensitivity was also observed in other SUMO substrates once a mutation occurred. For instance, DJ-1, a protein involved in the onset of Parkinson’s disease (PD), was sumoylated on lysine residue 130 (K130) [39]. The DJ-1 mutant L166P, which was identified in human PD patients, promoted sumoylation by forming poly- or multi-SUMO-1, and/ or by forming abnormal aggregates [39]. However, the altered sumoylation of L166P decreased the activity of DJ-1.

How these mutations affected Nkx2.5 sumoylation in the presence of entire principle SUMO site remains unclear. It appears that the relative distance from a mutation site to the SUMO site had no any predictable impact on Nkx2.5 sumoylation. However, it is noteworthy that the mutant R25C exhibited higher sensitivity to SUMO conjugation compared with wild type Nkx2.5, because the lower expression of SUMO-1 protein induced its sumoylation to the level equivalent to that of wild type Nkx2.5 with higher expression of SUMO protein. This increased sensitivity of R25C to sumoylation probably accounted for the higher activation of target gene promoter by R25C, possibly due to active endogenous sumoylation machinery in Hela cells. And, if this is the case, then the addition of exogenous SUMO-1 would do little to further increase its activity. However, this hypothesis needs further investigation. The higher transcriptional activity of R25C mutant observed in the present study was not observed in a previous report [5]. This discrepancy was likely attributable to the different systems in which the experiments were conducted. Increased sumoylation sensitivity was also observed in other SUMO substrates once a mutation occurred. For instance, DJ-1, a protein involved in the onset of Parkinson’s disease (PD), was sumoylated on lysine residue 130 (K130) [39]. The DJ-1 mutant L166P, which was identified in human PD patients, promoted sumoylation by forming poly- or multi-SUMO-1, and/ or by forming abnormal aggregates [39]. However, the altered sumoylation of L166P decreased the activity of DJ-1.

How these mutations affected Nkx2.5 sumoylation in the presence of entire principle SUMO site remains unclear. It appears that the relative distance from a mutation site to the SUMO site had no any predictable impact on Nkx2.5 sumoylation. For instance, four HD mutations and V309L, the former closer to K51 than C265Y, and the latter farther away from K51 than C265, all showed repressive effects on Nkx2.5 sumoylation. R25C, on the other hand, which is the most adjacent to K51, increased the sensitivity of Nkx2.5 to sumoylation. Possibly, these mutations caused changes of three dimensional structure of Nkx2.5, which mask the SUMO site and/or make it less accessible to SUMO pathway components (HD and C309L mutations), or expose the SUMO site better for sumoylation (R25C mutation). Alternatively, these sites are required for a direct physical contact of Nkx2.5 with SUMO pathway factors, and mutations either disrupt or enhance this contact. The above findings indicate that all HD mutations of Nkx2.5, which negatively affect its DNA binding activity, also severely affect its sumoylation. These observations point to the likelihood that various cardiac phenotypes associated with different Nkx2.5 mutants are in part...
Table 1. Cardiac functional analysis in the progenies obtained from cross-breeding between K51R-Tg and Nkx2.5<sup>+/−</sup> mouse lines.

| Genotypes                  | wt (n = 3) | Nkx2.5<sup>+/−</sup> (n = 4) | KS1R-Tg (n = 3) | KS1R-Tg/Nkx2.5<sup>+/−</sup> (n = 4) |
|----------------------------|------------|-------------------------------|-----------------|-------------------------------------|
| Diameters; s (mm)          | 2.01 ± 0.47| 1.91 ± 0.37                   | 1.97 ± 0.05     | 2.85 ± 0.35                         |
| Diameters; d (mm)          | 3.36 ± 0.36| 2.94 ± 0.28                   | 3.43 ± 0.04     | 3.84 ± 0.28                         |
| Volume; s (μl)             | 15.50 ± 0.08| 13.78 ± 5.67            | 12.32 ± 0.83    | 33.08 ± 8.42                         |
| Volume; d (μl)             | 47.75 ± 12.68| 34.50 ± 7.25            | 48.47 ± 1.38    | 64.95 ± 10.93                        |
| stroke volume (μl)         | 32.25 ± 3.95| 21.22 ± 3.01               | 36.14 ± 2.17    | 31.88 ± 3.84                         |
| %EF                       | 72.32 ± 9.33| 65.97 ± 9.59               | 74.45 ± 2.37    | 51.68 ± 7.53                         |
| %FS                       | 41.75 ± 7.26| 36.84 ± 7.13               | 42.45 ± 2.14    | 26.59 ± 4.96                         |
| cardiac output (ml/min)    | 12.87 ± 2.24| 9.39 ± 1.39                | 14.41 ± 0.41    | 13.42 ± 1.83                         |
| LV mass (mg)               | 123.17 ± 12.72| 111.88 ± 18.32          | 113.53 ± 11.84  | 128.80 ± 16.49                       |
| BW (g)                     | 25.25 ± 2.80| 25.25 ± 2.33               | 25.73 ± 1.95    | 24.65 ± 2.95                         |
| LV mass/BW                 | 4.60 ± 0.42| 4.60 ± 0.93                 | 4.40 ± 0.16     | 5.30 ± 0.51                          |

Echocardiography was performed on animals of 4 months age as described in Materials and Methods. Note that although there was no statistically significant difference reached in all parameters shown between compound K51R-Tg/Nkx2.5<sup>+/−</sup> group and the other three groups due to small size of the sample examined, there was a trend of increase in diastolic and systolic volumes, and a trend of decrease in cardiac function (%EF and %FS) in the compound K51R-Tg/Nkx2.5<sup>+/−</sup> mice compared with the other group animals. EF, ejection fraction; FS, fractional shortening.

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Figure 6. Two out of four surviving compound K51R-Tg/Nkx2.5<sup>+/−</sup> mice exhibited dilated cardiac chamber and cardiac dysfunction. Representative echocardiography images from age-matched four genotype animals were shown. Note the dilated left ventricular chamber in K51R-Tg/Nkx2.5<sup>+/−</sup> mouse. Red broken line: left ventricular internal diameter (LVID, diastolic); green broken line: left ventricular internal diameter (LVID, systolic). EF, ejection fraction; FS, fractional shortening.

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attributable to the different degrees of altered sumoylation activity of Nkx2.5 mutants.

**Sumoylation-deficient Nkx2.5 K51R mutant and CHDs**

Previous studies demonstrated that expressing wild type Nkx2.5 or one HD mutant I183P caused embryonic lethality or cardiac anomalies [7,8]. In contrast to the above observations, cardiac specific expression of K51R mutant in the wild type mice was not sufficient to generate significant cardiac defects. Although K51R and those HD mutants shared certain properties such as attenuated transcriptional activity and reduced sumoylation [18], it is possible that K51R and HD mutants could behave differently in vivo. For example, K51R and HD mutants may disrupt physical association with different co-factor(s). Alternatively, it may also involve variable expression levels of the transgenes. However, in striking contrast to the lack of overt cardiac malformation in single K51R-Tg mice, a large portion of Nkx2.5 haploinsufficient mice with overexpressed K51R mutant in heart developed lethal congenital heart defects – ASD/VSD. It should be pointed out that the ASD/VSD observed in the compound K51R-Tg/Nkx2.5/-/- mice might not be attributable only to the defective sumoylation of K51R; the certain level of dominant negative activity of K51R mutant should also be taken into consideration [18]. However, the current findings support the notion that K51R mutant could mimic those sumoylation deficient CHDs-associated Nkx2.5 mutants in vivo to generate cardiac structural defects in the condition of Nkx2.5 haploinsufficiency.

The exact mechanism(s) by which K51R mutant caused CHDs in the presence of one wild type Nkx2.5 allele remains unclear. Apparently, apoptosis was not involved, despite it is a critical process for cardiac development and function. Although normal cardiomyocyte proliferation was observed in the hearts of Nkx2.5 null embryos at embryonic day (E) 10.5 [1], overexpression of wild type Nkx2.5 did increase cardiomyocyte division [40]. Given the finding that K51R mutant inhibited the transcriptional activity of wild type Nkx2.5 partially via competing with Nkx2.5 dimerization [18], the observation of substantially suppressed cardiomyocyte proliferation in the compound K51R-Tg/Nkx2.5/-/- mice indicated a positive role of Nkx2.5 in cardiomyocyte division. Whether this diminished cardiomyocyte proliferation was solely responsible for the development of CHDs present in the compound K51R-Tg/Nkx2.5/-/- mice requires further interrogation.

One of the limitations of the approach used in the present study is that cardiac α-MHC controlled K51R expression was restricted to myocardium, therefore the anomalies involving great vessels such as double outlet right ventricle (DORV) and Tetralogy of Fallot (TOF) that were seen in human patients associated with heterozygous Nkx2.5 mutants were not expected/observed in our model. Thus, it will be interesting to see if K51R knockin in murine model would lead to the development of great vessel defects that resemble those observed in human patients.

**SUMO and cardiovascular disorders**

Involvement of the SUMO pathway in a number of pathogenic processes such as the development of craniofacial defects, neurodegenerative disease progression and cancer development has been considerably investigated [41,42,43], however, evidence for the role of SUMO in the development of cardiovascular disorders is newly emerging. SUMO targets a number of transcription factors, such as serum response factor (SRF) [44], GATA4 [43,46], and myocardin [25], which are essential for normal cardiac morphogenesis and function [20]. It is important to note that the covalent attachment of SUMO to the above-mentioned factors positively regulated their functions, counter to the general belief that sumoylation serves as a negative regulator of the activity of substrates. Recently, two naturally occurring sumoylation-defective mutants of the nuclear structural protein, lamin A/C, were proposed to be potentially linked to human familial dilated cardiomyopathy [47]. Similarly, the present study demonstrated altered sumoylation status in a number of human Nkx2.5 mutants that are implicated in CHDs. Moreover, expression of sumoylation deficient Nkx2.5 mutant in the mice with one wild type Nkx2.5 allele caused CHDs. Given the findings of CHDs in the SUMO-1 mutant mice [22], and identification of a unique mutation in the cis-regulatory sequence of the SUMO-1 gene in a patient with both clef lip/palate and ASD [22], our studies further indicate a significant influence of the SUMO pathway in the cardiovascular diseases. With the advances in the phenotypic analysis of additional animal models with genetic modifications that specifically target the SUMO conjugation machinery, we believe that the essentiality of an intact SUMO conjugation pathway for normal cardiac structural morphogenesis and function will be clarified.

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**Author Contributions**

Conceived and designed the experiments: JW. Performed the experiments: JW IPM. Analyzed the data: JW EYK LC YM WY. Contributed reagents/materials/analysis tools: JC. Wrote the paper: JW.

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