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LQTS Mutation N1325S in Cardiac Sodium Channel Gene SCN5A Causes Cardiomyocyte Apoptosis, Cardiac Fibrosis and Contractile Dysfunction in Mice

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Abstract

Objective—Mutations in the cardiac sodium channel gene *SCN5A* cause long QT syndrome (LQTS). We previously generated a LQTS mouse model (TG-NS) that overexpresses LQTS mutation N1325S in *SCN5A*. The TG-NS mice manifested the clinical features of LQTS including spontaneous VT, syncope and sudden death. However, the long-term prognosis of LQTS on the structure of the heart has not been investigated in this or any other LQTS models and human patients.

Methods and Results—Impaired systolic function and reduced left ventricular fractional shortening were detected by echocardiography, morphological and histological examination in two lines of adult mutant transgenic mice. Histological and TUNEL analyses of heart sections revealed fibrosis lesions and increased apoptosis in an age-dependent manner. Cardiomyocyte apoptosis was associated with increased activation of caspases 3 and 9 in TG-NS hearts. Western blot analysis showed significantly increased expression of key Ca²⁺ handling proteins L-type Ca²⁺ channel, RYR2 and NCX in TG-NS hearts. Increased apoptosis and altered expression of Ca²⁺ handling proteins could be detected as young as 3 months of age when echocardiographic showed little or no alterations in TG-NS mice.

Conclusions—Our findings revealed for the first time that LQTS mutation N1325S in *SCN5A* causes cardiac fibrosis and contractile dysfunction in mice, possibly through cellular mechanisms involving aberrant cardiomyocyte apoptosis. Therefore, we provide the experimental evidence

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supporting the notion that some LQTS patients have an increased risk of structural and functional cardiac damage in a prolonged disease course.

Keywords

Long QT syndrome; Dilated cardiomyopathy; Heart Failure; Cardiac sodium channel gene SCN5A; Mutation; Cardiomyocyte apoptosis

1. Introduction

The SCN5A gene, encoding the voltage-gated cardiac sodium channel Na_v1.5, is one of the major genes responsible for long QT syndrome (LQTS).[1,2] The N1325S mutation in SCN5A was identified in a 96-member family in which 21 members died suddenly.[3] A follow-up inquiry on the N1325S family discovered that one of the three family members with echocardiographic data showed structural abnormalities consistent with dilated cardiomyopathy (DCM).[3] This observation prompted us to investigate the long-term prognosis of LQTS in a transgenic mouse model that overexpresses the N1325S mutation (referred to herein as TG-NS mice).

Two lines of TG-NS mice (TG-NSL3 and TG-NSL12 with 1–2 and 10 copies, respectively, of the mutant *SCN5A* with N1325S) and two lines of mice expressing the wild-type *SCN5A* (TG-WTL5 and TG-WTL10 with 1–2 and 10 copies, respectively, of the wild-type *SCN5A*) were generated.[4–6] TG-NSL3 and TG-NSL12 mice showed increases of QTc by 23 and 91%, and increases of action potential duration (APD) by 53 and 128%, respectively, relative to control mice.[4] APD became unstable and markedly prolonged with increasing frequency of stimulation in TG-NS mice.[4] AP alternans, early and delayed after-depolarizations (EADs and DADs, respectively) were detected at decreasing diastolic intervals in TG-NS but not in control mice.[3,4] Compared to TG-WT and non-transgenic (NT) control mice, TG-NS mice showed increased frequencies of ventricular tachycardia (VT), ventricular fibrillation (VF), syncope or sudden death.[4,5] In the current study, we carried out a long-term follow-up investigation of TG-NS mice, using TG-WT mice as agematched controls.[6] We characterized TG-NS mice for cardiac morphology and contractile function, cardiomyocyte apoptosis, fibrosis, activation of caspases, and remodeled expression of key Ca²⁺ handling proteins.

2. Materials and methods

2.1 Experimental animals

Generation and genotyping of TG-NS and TG-WT transgenic mice have been described previously.[4,6] All mice are on a CBA/B6 background. Experimental procedures performed in this study were approved by the Cleveland Clinic Institutional Animal Care and Use Committee, and conform with the NIH Guidelines.

2.2 Morphological and histological analysis

Mice were euthanized by CO_2 , and lungs and hearts were isolated immediately. The lungbody weight ratio and heart-body weight ratio were determined by immediate weighing. The whole heart morphology was examined under a dissection microscope (LEICA MZFLIII) at the same magnification. Then, the hearts were washed by PBS and fixed in 4% paraformaldehyde and paraffin-embedded. Serial sections of $10~\mu m$ were cut and stained with haematoxylin/eosin (H&E) or Masson's trichrome. All sections were prepared in an identical manner. The presence of interstitial collagen fiber accumulation was used as a marker of cardiac fibrosis. The percentage of interstitial fibrosis to the total heart section

area was calculated from 10 randomly selected microscopic fields in four individual sections per heart using ImagePro Plus software (Media Cybernetics, Inc., Bethesda MD).

2.3 TUNEL assays

Frozen sections of heart tissues were used for the TUNEL assay with an ApopTag Peroxidase In Situ Apoptosis Detection Kit (CHEMICON) according to the manufacturer's instructions. Twenty microscopic fields were randomly selected from each section and six individual sections per heart were examined at a high magnification (400X). For each line of mice, 6–17 mice were studied (n=6–17). For each mouse, 5000 to 8,000 cells were counted. The number of TUNEL-positive cells and the number of total nuclei were counted, and the percentage was calculated as described. [7]

Adjacent heart sections were also examined by double staining of H&E and TUNEL. TUNEL staining was performed as described above except that H&E staining, instead of methyl green above, was used for counterstaining.

2.4 Immunohistochemistry

Frozen heart sections were first stained with TUNEL as described above, and then subjected to a second immunohistochemistry with VECTASTAIN ABC-AP system (Vector Laboratories) with a monoclonal anti-α-actinin antibody (diluted at 1:500, Sigma). The sections were then stained with a biotinylated secondary antibody, and visualized with Vector Red alkaline phosphatase substrate. The sections were counterstained by methyl green.

2.5 Echocardiography

High-frame rate (>200 fps) echocardiography was preformed as described [8] with a 14-MHz transducer coupled to Applio ultrasound machine (Toshiba, Japan).

2.6 Assays for activation of caspases

Caspase-3 activation activity in myocardial tissues was measured with the Caspase-3 Colorimetric Activity Assay Kit according to the instructions by the manufacturer (Chemicon). Caspase-8 and caspase-9 activation activity in myocardial tissues was measured using the Caspase-8 and -9 Colorimetric Assay Kit according to the instructions by the manufacturer (R&D Systems). The assay was performed in triplicate in 96-well plates.

2.7. Western blot analysis

Total protein extracts were extracted from fresh, isolated hearts by using the lysis buffer containing 20 mM Tris-HCl (pH 8.0), 100 mM NaCl, 1 mM EDTA, 0.5% NP-40 and 1 \times protease inhibitor cocktail solution. Western blot analysis was performed with 30–75 μg of total protein extracts using primary antibodies against Na⁺/Ca²⁺ exchanger NCX-1 (Chemicon), SERCA2 (Santa Cruz Biotechnology), dihydropyridine receptor alpha 2 (Bioreagents) and ryanodine receptor RyR2 (Sigma) as well as GAPDH as the loading control.

2.8 Statistical analysis

All values are presented as mean \pm SEM. Statistical analysis for group data was performed using one way ANOVA. The paired Student's *t*-test was used to determine the significant level of a difference between two variables. The significance level was set at a P value of less than 0.05 or wherever indicated in the figure legends.

3. Results

3.1 Morphological and histological evidence of abnormal cardiac structure in TG-NS mice

Gross morphological examinations of whole hearts and transverse heart sections from TG-NS mice detected myocardial fibrosis and increased chamber dimensions compared to TG-WTL10 control mice. Enlarged and dilated hearts were seen consistently in both lines of TG-NS mice (TG-NSL3 and TG-NSL12) at 6–10 months of age, but not in comparable control TG-WTL10 and NT mice (Fig. 1). Typical features of DCM were detected in 3 of 6 TG-NSL12 mice and 3 of 9 TG-NSL3 mice by histological examinations and echocardiography described below. Dilatation or enlargement of the heart was rarely detected in TG-NS mice at the age of 2–3 months (data not shown). There were no significant differences in values of the heart weight/body weight ratio among TG-NS, TG-WTL10 and NT mice (Fig. 2A, left panel). However, the lung weight/body weight ratio, an index of pulmonary congestion and left ventricular systolic dysfunction, was significantly increased in TG-NS mice compared with NT mice or TG-WTL10 mice (Fig. 2A, right panel). Histological analysis showed pulmonary congestion and edema in TG-NSL3 and TG-NSL12 mice at the age of 6–10 months (Fig. 2B). No significant alteration in a liver/body weight ratio and hepatic congestion was detected in the two lines of TG-NS mice.

Masson's trichrome staining of heart sections from 4 groups of mice at 6–10 months of age showed a significant increase of development of cardiac fibrosis in both lines of TG-NS mice compared with NT and TG-WTL10 mice, respectively (Fig. 3). For mice at the age of 2–3 months, fibrosis was barely detectable and no significant differences were found among the TG-NSL12, TG-NSL3, TG-WTL10, and NT mice (data not shown). Therefore, the morphological and histological alterations in TG-NS mice are indicative of an age-dependent development of contractile dysfunction and myocardial fibrosis.

3.2 Abnormal cardiac structure and functional performance in TG-NS mice by echocardiography

To further explore the phenotype of TG-NS mice, echocardiography was carried out to examine the structural and functional performance of the heart. A significant increase in systolic left ventricular end dimension (LVEDS), and decreases in systolic intraventricular septal thickness (IVSS) and systolic posterior wall thickness (PWS) in 6–10 month TG-NS mice relative to NT or TG-WTL10 mice was noted (Fig. 1 bottom panels, and Table 1). Left ventricular fractional shortening (LVFS) was significantly less in TG-NS mice than in NT mice: values of LVFS in TG-NSL12, TG-NSL3, and NT mice were 24.85 \pm 5.64, 42.78 \pm 1.95, and $60.0 \pm 5.7\%$, respectively (P < 0.005 and P < 0.05 for TG lines 12 and 3 vs. NT mice; Table 1). TG-WTL10L10 mice displayed normal values of structural parameters (Fig. 1 and Table 1) and a normal LVFS value of $58.44 \pm 1.80\%$ (P > 0.05 compared to NT mice), and there were no significant differences between the two TG-WTL10L10 lines with different copy numbers of the transgene (data not shown). Reduced LVFS in TG-NS hearts indicates left ventricular systolic dysfunction. It is interesting to note that the value of LVFS in hearts from TG-NSL12 mice decreased from $42.31 \pm 0.8\%$ at 3 months to $24.85 \pm 5.64\%$ at 8 months (Table 1). Similarly, age-dependent alternations in heart structure, as reflected by larger values of LVEDS and LVEDD in 8-month than in 3-month old TG-NSL12 mice (Table 1), were also observed. These data suggest that the development of contractile dysfunction in TG-NSL12 mice was progressive.

The heart rate did not vary significantly in young mice (3 months old) among different groups (Table 1). In 8-month old mice, the heart rate in TG-NSL3 mice did not vary significantly from control mice, however, the rate in TG-NSL12 mice was increased significantly compared to control mice.

3.3 Progressive loss of cardiomyocytes by apoptosis in TG-NS mice

Contractile impairment observed in TG-NS mice prompted us to conduct further studies to investigate the underlying mechanisms. Apoptosis has been recognized as playing an important role in HF.[9] We performed TUNEL assays on cross-sections of TG-NS hearts to detect apoptosis. H&E counterstaining was carried out in conjunction with the TUNEL assay (Fig. 4Aiii, 4A-iv) to identify the apoptotic cells by histology and morphology. Moreover, a typical morphological feature of apoptosis, condensation of nuclear chromatin, was captured in sections from TG-NS heart (Fig. 4A-v), which further supports apoptotic development. [10] A higher percentage of TUNEL-positive nuclei was found in TG-NS hearts than in control hearts (Fig. 4A-i, 4A-ii, 4B). The percentage of TUNEL-positive nuclei was $2.83 \pm 0.28\%$ in TG-NSL12, $2.13 \pm 0.53\%$ in TG-NSL3, $0.06 \pm 0.02\%$ in TG-WTL10 and $0.07 \pm 0.01\%$ in NT hearts at the age of 6–10 months (Fig. 4B). Immunohistochemical analysis using a cardiomyocyte-specific anti-α-actinin antibody in conjunction with the TUNEL assay verified that the TUNEL-positive cells were indeed cardiomyocytes (Fig. 4A-vi). Further analyses showed that cardiomyocyte apoptosis (i.e., the percentage of TUNEL-positive cells) increased with age in TG-NSL12 hearts from 0.21% in 3-month-old mice to 3.90% at the age of more than 10 months (Fig. 4C).

3.4 Increased activation of caspase-3 and 9 associated with cardiomyocyte apoptosis

As apoptosis is usually mediated by activation of caspases,[7] we next examined the state of activation of caspases 3, 8, and 9 in TG-NS hearts. Caspase-3 activity was nearly 60% greater in TG-NSL12 than in TG-WTL10 or NT hearts (Fig. 5A). Further analysis revealed a significant 2-fold increase in caspase-9 (Fig. 5C), but no change in caspase-8 activity (Fig. 5B) in TG-NSL12 hearts compared with controls. These data suggest that apoptosis in TG-NS hearts may involve the caspase-9-dependent mitochondrial apoptotic pathway.

3.5 Alterations of Ca²⁺ handling proteins in TG-NS mice

Remodeling of expression of some key Ca^{2+} handling proteins was evaluated in TG-NS and control mice (Figure 6). Significantly increased expression of NCX-1, RyR2 and L-type Ca^{2+} channel was detected in both lines of TG-NS mice compared to TG-WTL10 and NT mice (Figure 6A, 6B, 6D). However, the expression of SERCA2 was not significantly different amongst the 4 groups of mice (Figure 6C).

4. Discussion

The N1325S mutation in the cardiac sodium channel gene *SCN5A* has been previously shown to cause type-3 LQTS (LQT3) in both humans and mice. The results from this study clearly demonstrate that mutation N1325S also causes myocardial damage in a progressive, age-dependent manner. Echocardiographic and histological examinations of hearts from the TG-NS mice with N1325S revealed structural and functional impairment of the left ventricle, and findings reminiscent of abnormalities seen in human heart failure patients, which were not detected in age-matched control TG-WTL10 mice overexpressing wild-type *SCN5A*. The lung weight/body weight ratio in TG-NS mice was greater than in control mice. Pulmonary congestion and edema were observed in both lines of TG-NS mice. The increased ratio may reflect the presence of lung edema secondary to left ventricular dysfunction.[11]

The observed phenotype in TG-NS mice is unlikely to be caused by excessive overexpression of *SCN5A*. Transgenic TG-NSL12 mice expressing the mutant N1325S *SCN5A* gene and transgenic TG-WTL10 mice expressing the wild-type gene have the same copy number and expression level of the transgene, but only the former developed the heart failure phenotype. Secondly, TG-NSL3 mice, which express 1–2 copies of the mutant

transgene, also developed the same DCM and heart failure phenotype as TG-NSL12, albeit with less severity. Similarly, it is unlikely that the insertion location of the transgene is the cause of heart failure in TG-NS mice. First, two mutant TG-NS lines developed the same cardiac phenotype as TG-NSL12, but two TG-WT lines (L5 and L10) did not show signs of heart failure. Furthermore, two transgenic lines (unrelated to this study) that overexpress another *SCN5A* mutation (R1193Q) did not show the heart failure phenotype (data not shown). Taken together, the results on these multiple transgenic lines support the interpretation that it is the N1325S mutation, rather than the overexpression of *SCN5A* or the insertion location of the transgene in the genome, which causes contractile dysfunction.

Previously, we reported that in an LQTS family with the N1325S mutation, one of the three patients with echocardiographic data available clearly developed DCM.[3] Furthermore, in an LQTS family with the *SCN5A* mutation E1784K, the autopsy report on the proband (who died suddenly at age 31) revealed DCM.[12] Recently, Shi et al. reported that two mutation carriers in an LQT3 family with mutation delQKP 1507–1509 in *SCN5A* showed features of DCM [13]. The results with transgenic mice in this study substantiate the human studies and clearly indicate that there may be a potential risk of DCM associated with LQT3. However, it is important to note that the severity of contractile dysfunction seems to vary from a mouse to another. In humans, three patients in the LQTS family with the N1325S mutation had echocardiographic examinations, and one of them showed DCM. In mice, 50% of TG-NSL12 mice (3 of 6 mice) and 30% of TG-NSL3 mice (3 of 9 mice) showed typical features of DCM by both echocardiographic and histological examinations.

Several mutations in SCN5A have been reported to be associated with DCM, for example, heterozygous missense mutations D1275N, T220I, R814W and D1595H and one truncation mutation (2550–2551 ins TG).[14,15] Compound heterozygous mutations W156X and R225W in SCN5A were associated with DCM, severe cardiac conduction disturbances and degenerative changes in the conduction system, and irregular wide complex tachycardia.[16] When expressed in Xenopus oocytes, mutant channels with W156X did not produce any current and those with R225W showed severely reduced sodium currents and abnormal gating. Three other mutations, D1275N, R814W and D1595H, were also functionally characterized. When expressed in Xenopus oocytes, mutation D1275N shifted the voltage dependence of activation to more depolarized voltages.[17] Mutations R814W and D1595H were studied in HEK293 cells.[18] D1595H impaired fast inactivation by slowing entry into the inactivated state, whereas R814W increased channel opening at hyperpolarized voltages. Interestingly, mutant R814W channels showed an increased late window current, a similar characteristic shared by mutation N1325S under the current study. These studies suggest that SCN5A mutations associated with DCM act by a diverse array of mechanisms. Curiously, different from the patients with mutation N1325S, the single patient with mutation R814W exhibited dilated cardiomyopathy, atrial fibrillation, and non-sustained ventricular tachycardia, but not LQTS associated with late sodium currents.

Multiple studies examined the link between prolonged QTc and heart failure in human patients. Davey et al. showed that the resting QTc was significantly longer in heart failure patients (470 +/- 10 ms) than age-matched controls with normal left ventricular structure and function (421 +/- 6 ms).[19] Among HF patients, 43%–72% of them showed prolonged QTc of >440 ms. VT, VF, and sudden death are also commonly documented in more than 50% patients with HF.[20–22] These important observations associated with heart failure were all captured in the TG-NS mice under this study. Therefore, TG-NS mice may serve as an animal model for investigating the pathogenic mechanisms of QT prolongation and ventricular arrhythmias associated with HF.

Our results suggest that cardiomyocyte apoptosis may be a cause of DCM in TG-NS mice. Consistent with this, cardiomyocyte apoptosis was shown to be the cause of contractile dysfunction in a transgenic mouse model, and its inhibition by caspase inhibitors improved LV function.[7] Cardiomyocyte apoptosis may also be a cause of ventricular arrhythmias in TG-NS mice. Apoptosis may lead to cardiac fibrosis, which then serves as a substrate for reentry arrhythmias detected in TG-NS mice. Although the detailed mechanism for apoptosis in TG-NS hearts is not known, activation of caspases 3 and 9, but not caspase-8, suggests that cardiomyocyte apoptosis in these mice is via the mitochondrial apoptotic pathway. An extensive microarray study revealed that expression of STAT1 was markedly increased in TG-NS hearts[23] and such an increase may contribute to increased cardiomyocyte apoptosis[24]. Furthermore, we showed that compared to TG-WTL10 mice, expression of L-type Ca²⁺ channel, RYR2, and NCX was increased in TG-NS mice (Fig. 5), which may be associated with abnormal Ca²⁺ homeostasis. Increased expression of L-type Ca²⁺ channel on the cell surface and calcium release channel RYR2 on sarcoplasmic reticulum membranes may increase the intracellular calcium concentration. Increased NCX expression in the context of increased late sodium current generated by mutation N1325S may increase the reverse mode activity of NCX, resulting in the increased intracellular calcium concentration. Aberrant Ca²⁺ homeostasis was linked to apoptosis and activation of caspases.[25] Future studies are needed to distinguish whether STAT1 overexpression or Ca²⁺ abnormalities is a major cause of apoptosis in TG-NS hearts. Although we can not exclude the potential contribution of incessant tachyarrhythmias to development of contractile dysfunction in TG-NS mice, we are not able to establish any association between development of these two disorders based on our data. We could detect both arrhythmias and contractile dysfunction in as early as 3 month old TG-NS mice and we have not observed any age preference to the onset of arrhythmias over contractile dysfunction. Therefore, the possibility exists that mutation N1325S could lead to the development of both incessant tachyarrhythmias and contractile dysfunction in parallel. However, tachycardia in TG-NSL12 mice could significantly contribute to the further development of contractile dysfunction, which can be investigated in the future by potential prevention of the phenotype by long-term application of beta blockers.

In summary, our results provide convincing evidence that cardiac structural abnormalities and contractile dysfunction can arise from transgenic overexpression of a congenital LQT3 mutation, N1325S, in *SCN5A*, which validates our earlier finding of association of DCM with one N1325S carrier in humans. Overexpression of N1325S was associated with cardiomyocyte apoptosis and cardiac fibrosis, which could be contributing factors to contractile dysfunction. We postulate that expression remodeling of key cardiac proteins, including Ca²⁺ handling proteins or STAT1, leads to activation of caspases 3 and 9, which promotes cardiomyocyte apoptosis, resulting in contractile dysfunction. Furthermore, our results suggest that overexpression of an *SCN5A* mutation not only increases the occurrence of arrhythmias and sudden death, but also increases the risk for development of cardiac pump failure. Although heart failure can arise from a myriad of etiologies, the TG-NS mice with mutation N1325S should serve as an interesting model for studying the molecular mechanisms that underlie arrhythmias and progressive cardiac dysfunction and the development of heart failure.

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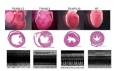


Figure 1.

Abnormal cardiac structure in TG-NS mice. The gross morphology of the whole heart (first row) and H&E staining of transverse heart sections (second row; horizontal bars indicate 200 μm) revealed cardiac enlargement and ventricular dilatation in 6–10-month old TG-NSL12 and TG-NSL3, but not in TG-WTL10 or NT mice. Representative echocardiographic M-mode images for each of the corresponding hearts are illustrated in the third row.

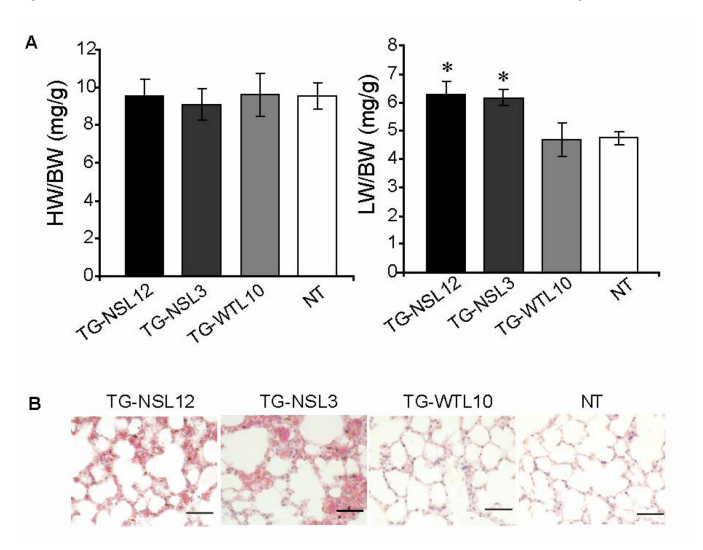


Figure 2. Histological analysis of the lung. (A) Significant increases in the lung weight/body weight ratio were observed in both lines of TG-NS mice (TG-NS-L12, n=12; TG-NS-L3, n=11), compared with TG-WTL10 (n=10) or NT (n=19) mice (*P<0.01). The heart/body weight ratios of 6–10-month old TG-NSL12, TG-NSL3, TG-WTL10, and NT mice were not different. (B) Pulmonary congestion and edema were observed in both lines of TG-NS mice at 6–10 months of age. The alveolar capillaries of the lung were distended and filled with erythrocytes. In some areas, the alveolar space was filled with fluids. Normal alveolar structures were shown as in TG-WTL10 or NT mice. n=3; Scale bar (bottom right panel) = $20 \, \mu m$.

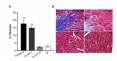


Figure 3.

Significant increase of development of cardiac fibrosis in both lines of TG-NS mice. (A) Representative stained sections from 4 groups of mice (n=5 mice/group) indicates fibrosis (light blue color) in both lines of TG-NS mice, but not in TG-WTL10 and NT mice. (B) Fibrotic tissue was greater in 6–10-month old TG-L12 and TG-L3 mice than in TG-WTL10 or NT mice (bar graph), by Masson's trichrome staining of heart sections. Scale bar (bottom right panel) = $20~\mu m$. "**" Denotes statistically significant difference from TG-WTL10 (P<0.0001).

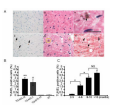


Figure 4.

Evaluation of cardiomyocyte apoptosis. (A) Representative images of heart sections using the TUNEL assay. Brown nuclei are TUNEL positive and apoptotic cells (arrows). i, NT; ii, TG-NSL12; iii and iv, TUNEL stained images counterstained with H&E for NT and TG-NSL12, respectively; v, a representative image of a TUNEL-positive nucleus undergoing condensation of nuclear chromatin (arrow); vi, image of TG-NSL12 with staining for both TUNEL and a monoclonal anti- α -actinin antibody, a probe specific for cardiomyocytes. Scale bar = 20 μ m. (B) The fraction of TUNEL-positive nuclei in hearts of 6–10-month old TG-NSL12 (n=10) and TG-NSL3 (n=6) mice was significantly greater than in hearts of TG-WTL10 (n=10), and NT (n=17) mice. (** P<0.0001 vs TG-WT; * P<0.05 vs NT). (C) The number of TUNEL-positive nuclei in hearts of TG-NSL12 mice increased with age. (*P<0.05; NS means no significant difference; n=4 for 2–3 months, n=6 for 4–6 months, n=6 for 8–10 months, n=5 for more than 10 months group).

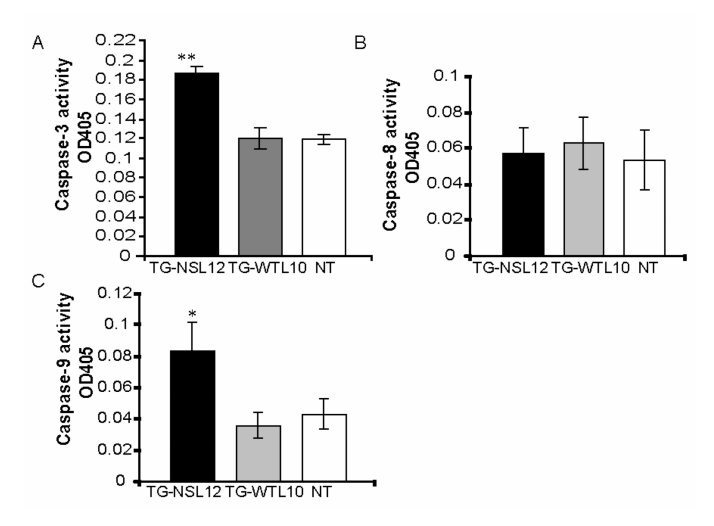


Figure 5. Assessment of caspase activation in heart tissue. (A) Increased caspase-3 activation was detected in hearts from TG-NSL12 (n=17) compared to those from TG-WTL10 (n=9) and NT (n=17) mice. **P=2.6 × 10⁻⁹ vs NT. (B) No significant difference was detected in caspase-8 activation amongst the three heart groups. (C) Increased caspase-9 activation was detected in TG-NSL12 hearts (n=10) compared to TG-WTL10 (n=12) and NT (n=11).age, 3–5 months old; *P<0.05.vs NT.

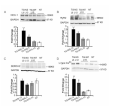


Figure 6.

Expression of NCX-1 (A), RyR2 (B), SERCA2 (C) and L-type Ca^{2+} channel (D) in hearts from TG-NS and NT mice at 3–5 months of age. Representative Western blots are shown at left panels and summarized data for the Western blot densitometrical analysis are at right panels. Cardiac expression of NCX-1 (A), RyR2 (B) and L-type Ca^{2+} (D) was significantly increased in TG-NSL3 and TG-NSL12 mice, but not in TG-WTL10 mice or NT mice. Expression of SERCA2 (C) was not significantly different amongst the four groups of mice. (TG-NSL12 and TG-NSL3, n = 7, respectively; TG-WTL10 and NT, n = 8, respectively). **P<0.001; *P<0.005.

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Table 1

Echocardiographic assessment of cardiac structure and function in the mice

		8 month	8 month old mice		31	3 month old mice	
Parameters	TG-NSL12 (6)	TG-NSL3 (9)	TG-NSL12 (6) TG-NSL3 (9) TG-WTL10 (6) NT (9)	(6) LN	TG-NSL12 (7)	TG-NSL12 (7) TG-NSL3 (5) NT (10)	(0I) LN
LV FS (%)	$24.9 \pm 5.6^{**#}$	$42.78 \pm 1.95^*$	58.44 ± 1.80	59.56 ± 3.88	42.3 ± 1.3**	45 ± 5.52*	63.2 ± 3.62
IVSD(mm)	0.94 ± 0.08	1.08 ± 0.08	0.91 ± 0.10	1.08 ± 0.07	0.98 ± 0.03	1.03 ± 0.06	0.95 ± 0.04
IVSS(mm)	$1.25 \pm 0.18^*$	1.67 ± 0.10	1.73 ± 0.10	1.83 ± 0.08	1.57 ± 0.07	1.64 ± 0.04	1.7 ± 0.09
LVEDD(mm)	$4.14\pm0.31\#$	3.33 ± 0.18	3.76 ± 0.17	3.32 ± 0.15	3.07 ± 0.07	3.35 ± 0.32	2.84 ± 0.14
LVEDS(mm)	3.14 ± 0.37 *#†	$2.04 \pm 0.10^*$	1.56 ± 0.11	1.41 ± 0.18	1.8 ± 0.04 **	1.91 ± 0.35	1.07 ± 0.13
LVPWD(mm)	0.96 ± 0.06	1.02 ± 0.05	0.99 ± 0.10	1.03 ± 0.09	0.90 ± 0.07	0.86 ± 0.04	0.91 ± 0.07
LVPWS(mm)	$1.28 \pm 0.11^*$	$1.34 \pm 0.05^*$	1.62 ± 0.13	1.61 ± 0.08	$1.32 \pm 0.08^*$	1.28 ± 0.12	1.53 ± 0.05
HR (bpm)	$812 \pm 37.7^{*7}$	645 ± 37.3	536 ± 36.5	643 ± 46.3	758 ± 57.1	745 ± 53.7	709 ± 28.2

ventricular cavity size in diastole; LVEDS-left ventricular cavity size in systole; LVPWD-left ventricular posterior wall thickness in diastole; LVPWS- left ventricular posterior wall thickness in systole; HRheart rate(bpm-beats per minute),TG-NSL12-mutant transgenic line L12; TG-NSL3- mutant transgenic line L3; TG-WTL10-transgenic mice with wide type SCN5A (line 10); NT-non-transgenic mice; The significant difference between TG-WT and NT in the 3 month old mice, thus the data are shown for the NT group only. LVFS-left ventricular fractional shortening, percent change in left ventricular cavity There were no significant differences between the two TG-WT lines with different copy numbers of the transgene, thus the data are shown for TG-WT L10 only. As in the 8 month old mice, there was no dimensions with systolic contraction([(LVEDD-LVESD)/LVEDD]×100); IVSD - left ventricular septal wall thickness in diastole; IVSS-left ventricular septal wall thickness in systole; LVEDD-left number of mice used for each group is in the bracket after each mouse line; Compared with NT, Page 16

 $^{^{**}}_{P < 0.005}$,

 $^{^{\}ast}$ P < 0.05; Compared with 3 month-old group,

 $^{^{\#}}$ P < 0.05; Compared with TG-NSL3,

 $^{^{7}\}mathrm{P}<0.05$