A TAT–Frataxin fusion protein increases lifespan and cardiac function in a conditional Friedreich’s ataxia mouse model

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Friedreich’s ataxia (FRDA) is the most common inherited human ataxia and results from a deficiency of the mitochondrial protein, frataxin (FXN), which is encoded in the nucleus. This deficiency is associated with an iron–sulfur (Fe–S) cluster enzyme deficit leading to progressive ataxia and a frequently fatal cardiomyopathy. There is no cure. To determine whether exogenous replacement of the missing FXN protein in mitochondria would repair the defect, we used the transactivator of transcription (TAT) protein transduction domain to deliver human FXN protein to mitochondria in both cultured patient cells and a severe mouse model of FRDA. A TAT–FXN fusion protein bound iron in vitro, transduced into mitochondria of FRDA deficient fibroblasts and reduced caspase-3 activation in response to an exogenous iron-oxidant stress. Injection of TAT–FXN protein into mice with a conditional loss of FXN increased their growth velocity and mean lifespan by 53% increased their mean heart rate and cardiac output, increased activity of aconitase and reversed abnormal mitochondrial proliferation and ultrastructure in heart. These results show that a cell-penetrant peptide is capable of delivering a functional mitochondrial protein in vivo to rescue a very severe disease phenotype, and present the possibility of TAT–FXN as a protein replacement therapy.

INTRODUCTION

Friedreich’s ataxia (FRDA; OMIM 229300) is a relentlessly progressive cardiac and neurodegenerative disease typically beginning in childhood that leads to loss of motor skills and, ultimately, inability to stand or walk within 10–15 years of onset (1). Virtually all patients develop a cardiomyopathy and heart failure is the most common cause of death (2,3). The prevalence of FRDA is ~1 in 50,000 people with equal frequency in males and females (4), and a carrier frequency of 1:60 to 1:120 (5–8). Inheritance is autosomal recessive and predominantly caused by a GAA triplet expansion in the frataxin (FXN) gene on chromosome 9q13–q21.11 (reviewed in 9,10). This triplet expansion, which often exceeds 800 repeats, is predicted to cause the formation of a triple-stranded DNA helix (11) leading to transcriptional inhibition and partial silencing of the FRDA locus with loss of FXN protein expression (12). Additionally, GAA triplet expansions may trigger chromatin condensation making the affected region of genomic DNA transcriptionally inactive (13). There is a correlation between the GAA repeat number and the onset and severity of clinical symptoms with higher repeat numbers being associated with earlier onset and more rapid rate of disease progression (14,15).

FXN is an essential and highly conserved protein expressed in most eukaryotic organisms that appears to function in mitochondrial iron homeostasis, notably the de novo biosynthesis of iron–sulfur (Fe–S) cluster proteins (16) and heme biosynthesis (17,18). FXN has been shown to bind iron along an acid ridge and the binding affinity can be significant (19). The exact function of FXN has not been defined but recent studies suggest that FXN functions as an allosteric activator with Fe2+ for Fe–S cluster biosynthesis by forming a four-protein complex that includes ISD11, ISCU, FXN and NFS1 (20–22). In this model, FXN induces a conformational change in the

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complex, enabling the direct sulfur transfer from cysteine for the Fe–S cluster assembly. The absence of FXN is associated with a loss of activity in Fe–S-containing proteins (23), such as aconitase, and a loss of energy production (24,25).

The 210 amino acid precursor FXN protein (23.1 kDa) contains an 80 amino acid mitochondrial targeting sequence (MTS) at the amino terminus. It is processed in two steps by the mitochondrial matrix processing peptidase (MPP) (26) as it is imported into the matrix (27). The intermediate form of FXN is cleaved at residue 42 by the MPP, and the mature form of FXN has been shown to be cleaved at amino acid 81 yielding a 130 amino acid with a predicted Mr of 14.2 kDa (28,29). Maturation of the precursor FXN occurs within the mitochondrial matrix and no other intra-mitochondrial posttranslational modifications have been identified.

Currently, there is no cure for FRDA. Treatment options at present logically include antioxidants and iron chelation (30,31). Although early clinical trials have shown modest biochemical improvement (32), these therapies have not shown substantial clinical improvement as they are designed to control downstream events resulting from the loss of FXN. Using a cell-penetrant peptide, transactivator of transcription (TAT), we tested the hypothesis that a TAT–FXN fusion protein could rescue the phenotype of FRDA using both patient fibroblasts, and extend the lifespan of the severe phenotype of the conditional FXN knockout mouse (33) as patient fibroblast cells, as well as the severe short-lived phenotype in vivo with mitochondrial protein defects.

RESULTS
TAT–FXN transduces into mitochondria of FXN-deficient human fibroblasts

The structure of the TAT–FXN fusion protein is shown in Figure 1A. TAT–FXN was expressed and purified from BL21 cells (see Supplementary Material, Fig. S1). To determine whether the TAT–FXN fusion protein would transduce across both cell and mitochondrial membranes, TAT–FXN was labeled with 5-idoacetamidofluorescein (5-IAF), incubated with FXN-deficient fibroblasts from FRDA patients for 3 h and then removed from the media. At 120 h after exposure to TAT–FXN, the cells were incubated with the mitochondrial-specific fluorescent dye, CMXRos (MitoTracker Red) (42,43), which localizes to mitochondria on the basis of the membrane potential, ΔΨm, and imaged as live cells by confocal microscopy. Figure 1B shows the green fluorescein from labeled TAT–FXN (panel 1), the red signal from mitochondrial uptake of MitoTracker Red (panel 2) and co-localization of both signals from mitochondria in panel 3. Previous work had shown that the TAT moiety must be removed after transduction into mitochondria or else it moves out of the mitochondrial matrix within 2 h (35,38). The continued presence of TAT–FXN in the mitochondria 120 h after treatment suggests that the TAT–FXN was processed in vivo by the mitochondrial MPP to remove the FXN MTS with its attached TAT peptide, thus leaving the processed FXN in the matrix.

TAT–FXN is processed by the MPP

To demonstrate that the TAT–FXN fusion protein would be appropriately recognized and cleaved by the MPP, we expressed and purified yeast MPP (44) to demonstrate processing of the precursor FXN. The fusion protein, TAT-mitochondrial malate dehydrogenase-enhanced green fluorescent protein (TAT-mMDH-eGFP), was used as a positive control because it has been shown to transduce into mitochondria in vivo and processed, and TAT–GFP was a negative control (38).

Figure 2A shows that the control, TAT–mMDH–eGFP, is progressively processed to completion by an overnight incubation with MPP as predicted. The upper band (upper arrowhead) is the precursor protein of TAT–mMDH–eGFP as shown by the starting material in lane 0. A decrease in signal from the precursor band is seen with increasing incubation time (1, 3 h, overnight) with an increase in signal from the processed band (lower arrowhead). There are two cleavage
sites within the MTS of the rat mMDH protein (45,46) with the first cleavage site generating an intermediate size protein from the loss of the TAT peptide and part of the MTS (estimated protein mass of 3.2 kDa), and a second cleavage site to generate the mature mMDH by the loss of an additional 0.8 kDa protein mass. Figure 2C shows separation of these reaction products by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS–PAGE) and staining by Coomassie. Sequencing of the lower molecular weight band (boxed area) by Edman degradation confirms processing of the TAT–mMDH–eGFP at the first protease sensitive site beginning at amino acid 17 of the precursor mMDH protein. This is in agreement with earlier published data of mMDH processing in mouse (26) and rat (45) for which the cleavage site exactly matches these results (FSTSA). That the TAT–mMDH–eGFP is not processed to maturity at the second predicted protease sensitive site supports earlier findings that this site is cleaved by the mitochondrial intermediate protease in the matrix, which is not present in these reactions (26,47).

Figure 2B and D shows that TAT–FXN is also appropriately recognized and processed by the MPP. Figure 2B is a western blot probed with a monoclonal anti-human FXN antibody showing the TAT–FXN precursor band (upper arrowhead), and Figure 2D is a Coomassie-stained SDS–PAGE of these reaction products. With progressive incubation times, there is an increase in the intermediate band (middle arrowhead in Fig. 2B, lower arrowhead in Fig. 2D), and a smaller band in Figure 2B (lower arrowhead). The TAT–FXN is processed to completion in an overnight digest (lane o/n).

The boxed area of the gel in Figure 2D was sequenced by Edman degradation and yielded the peptide fragment, LRTDI, shown in the cartoon at the lower arrowhead in Figure 2D. This peptide fragment corresponds exactly to the first cleavage site of human FXN at position 42 of the precursor FXN protein leading to the intermediate form (27). Unlike TAT–mMDH–eGFP, it appears that TAT–FXN is fully processed at the second cleavage site (position 81) to the expected size of the mature form by yeast MPP as shown by the predicted lower signal (lower arrowhead in Fig. 2B). This finding is consistent with earlier studies, showing that MPP alone is responsible for generating the intermediate and mature forms of FXN (27). However, this lower band was too faint to positively identify from the stained SDS–PAGE gel and could not be cut out for sequencing by Edman degradation. This is also consistent with earlier studies showing that conversion of the intermediate to mature forms of FXN is rate limiting in vitro (27).

In these experiments, TAT–GFP (no MTS) was used as a negative control and did not demonstrate any processing by the MPP on either a western blot using anti-GFP antibody, or by SDS–PAGE and staining as expected (see Supplementary Material, Fig. S2). This confirmed that the MPP was specific for the MTS in these mitochondrial fusion proteins and did not generate a non-specific protease action.
FXN is predicted to have an iron-binding role that supports the formation of Fe–S clusters in proteins, such as aconitase, and in the biosynthesis of heme (48, 49). The ability of TAT–FXN to bind iron was determined in vitro using the redox cycle of hydroquinone (HQ) to quinone (50) to generate a signal. As shown in Figure 3A, HQ can auto-oxidize to quinone thus generating the superoxide free radicals (51), which can be assayed by oxidation of the non-fluorescent reactive dye 2′,7′-dichlorodihydrofluorescence diacetate (DCHF-DA) to the fluorescent dye 2′,7′-dichlorofluorescein (DCF). The oxidized quinone is then reduced to HQ by ferrous sulfate, which is present in the incubation. We predicted that if TAT–FXN bound free iron, there would be a reduction of free iron content capable of reducing quinone back to HQ and a corresponding decrease in the production of superoxide. This could be quantified by a decrease in fluorescent signal from DCF.

Figure 3B shows that the precursor FXN molecule (as TAT–FXN) can bind iron in vitro using the redox cycle of hydroquinone (HQ) to quinone (50) to generate a signal. As shown in Figure 3A, HQ can auto-oxidize to quinone thus generating the superoxide free radicals (51), which can be assayed by oxidation of the non-fluorescent reactive dye 2′,7′-dichlorodihydrofluorescence diacetate (DCHF-DA) to the fluorescent dye 2′,7′-dichlorofluorescein (DCF). The oxidized quinone is then reduced to HQ by ferrous sulfate, which is present in the incubation. We predicted that if TAT–FXN bound free iron, there would be a reduction of free iron content capable of reducing quinone back to HQ and a corresponding decrease in the production of superoxide. This could be quantified by a decrease in fluorescent signal from DCF.

**TAT–FXN binds iron in a cell-free system**

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**TAT–FXN can rescue FRDA fibroblasts from an iron-oxidant stress**

In the absence of FXN, it is widely accepted that deficient cells will have an increased sensitivity to oxidative stress, which most likely contributes to the cascade of events leading to cytotoxicity (53–55). Iron with HQ induces an oxidative stress to cells (56–58) because HQ also forms a lipophilic chelate with iron and rapidly transfers the metal across the normally impermeable plasma membrane (59). Neither HQ nor Fe alone in culture media is toxic to FXN deficient...
fibroblasts even after an extended exposure of 24 h (Supplementary Material, Figure S3). Thus, FXN-deficient cells were treated with TAT–FXN 24 h earlier followed by changing to media without TAT–FXN prior to treating with 5 μM Fe/HQ for 5 h (Fig. 3C). Treatment with TAT–FXN by itself had no effect on the cells (Fig. 3C, control panels 2 and 3) and they appear as untreated cells (panel 1). However, when these cells were treated with Fe/HQ, changes in the morphology (rounded and spiculated cells) and loss of adherence of these cells indicated that Fe/HQ was cytotoxic (Fig. 3C, panel 4). In contrast, cells that were treated with TAT–FXN prior to the addition of Fe/HQ (panels 5 and 6) were able to survive and had reduced evidence of cytotoxicity as shown by their morphologic appearance being identical to control cells (panel 1).

TAT–FXN reduces caspase-3 activation in FRDA fibroblasts exposed to iron-oxidant stress

To determine whether TAT–FXN is capable of protecting FRDA cells against apoptosis after exposure to Fe/HQ-oxidant stress, FXN-deficient patient fibroblasts, or fibroblasts from a control patient without FRDA, were assayed for caspase-3 activation under baseline conditions and again after exposure to an iron-oxidant stress.

In Figure 3D, caspase-3 activation was significantly elevated (P < 0.001) in fibroblasts from a patient with FRDA when compared with fibroblasts from a healthy control patient. After exposure to an iron-oxidant stress, the same FRDA fibroblasts expressed significantly greater amounts of activated caspase-3 compared with baseline FRDA fibroblasts (P < 0.001). Treatment of FRDA fibroblasts with TAT–FXN prior to the introduction of an iron-oxidant stress was significantly protective against activation of caspase-3 when compared with Fe/HQ-treated FRDA fibroblasts (P < 0.001). As expected, there was no significant difference in caspase-3 activation between the control fibroblasts and FRDA fibroblasts exposed to Fe/HQ and treated with TAT–FXN (P > 0.05). This protection was dose related and was not due to the presence of TAT–FXN in the culture media because the fusion protein had been removed 24 h prior to introduction of the iron-oxidant stress. At 40 μg/ml, the amount of caspase-3 activation was below the baseline of the FRDA fibroblasts (P < 0.001).

TAT–FXN increases the survival rate and lifespan in Fxn-KO mice

The ability of TAT–FXN to protect FXN-deficient patient fibroblasts against Fe/HQ stress in vitro provided a strong rationale for determining whether a TAT–FXN fusion protein could rescue the Fxn knock out (Fxn-KO) mouse. Because the homozygous deletion of the Fxn gene is embryonic lethal (60), mice conditional for deletion of exon 4 of the Fxn gene (33) were studied using the NSE-Cre transgene (61–63) to generate progeny with the loss of FXN in heart and neural crest-derived tissues. Two groups of mice were examined for rescue with a TAT–FXN fusion protein: Fxn-KO mice receiving either TAT–FXN protein or PBS beginning at Day 3 of life (designated as 3d) until study end at 60 days of life, and Fxn-KO mice receiving either TAT–FXN protein or no PBS beginning at Day 12 of life ± 2 days (designated as 12d) and continuing until death (end of life study). All mice had to reach an age of 10 days to be included in the study, and all mice were weaned at 28 days of age. Control animals consisted of littermates heterozygous for the conditional allele (L3+/+) carrying or not carrying the recombinase, and had no clinical or biochemical phenotype (33). Fxn-KO mice in the 3d group received intraperitoneal (IP) injections three times per week, with either 15 mg/kg body weight of TAT–FXN, or an equivalent volume of sterile PBS. Dosing in the Fxn-KO 3d group was stopped at 60 days of life (end of study). The 3d control heterozygous littermates received equivalent volume injections of PBS. Fxn-KO mice in the 12d group received either TAT–FXN 20 mg/kg body weight (100 μl injections) given by the IP route beginning at 12 days of life ± 2 days twice weekly until death, or no injections of PBS. The 12d control heterozygous littermates did not receive injections.

The lengths of survival for both 3d and 12d groups were analyzed using the Kaplan–Meier estimator with log-rank sums to test the null hypothesis that there was no difference between populations in the probability of a death at any time point (64). The results are shown in Figure 4A and B. Pairwise multiple comparisons between all combinations of pairings (Holm–Sidak method) were performed to determine significance between groups and are shown in Table 1. By log-rank sums’ treatment with TAT–FXN beginning either at 3 days of age (3d group), or 12 days of age (12d group), resulted in a highly significant increase in lifespan of the Fxn-KO mouse. In the 3d group, treatment with TAT–FXN extended the mean survival of the Fxn-KO mouse by ~49% to 41.1 ± 4.3 days and was significant (P = 0.008) when compared with the mean age at death of 27.5 ± 2.3 days in the Fxn-KO mouse receiving only PBS (Table 1). It is noteworthy that five mice (31%) reached the end of the study at 60 days and died when TAT–FXN was discontinued, whereas no mouse in the PBS group reached this age and only one mouse survived beyond 35 days. In the 12d group, treatment with TAT–FXN extended the mean survival of the Fxn-KO mouse by ~53% to 43.1 ± 4.2 days and was significant (P = 0.003) when compared with the mean age at death of 28.1 ± 1.4 days in the Fxn-KO mouse not receiving PBS (Table 1). It is noteworthy that five mice (16%) exceeded 60 days of life with two mice expiring at 75 and 88 days. No mouse in the untreated group exceeded 40 days of life. There was no significant difference in the mean age at death between the 3d and 12d Fxn-KO mice receiving TAT–FXN, and in agreement with the original characterization of the model (33), there was no significant difference in the mean ages of death between the Fxn-KO mice receiving PBS (3d) versus no PBS (12d). In the 12d KO no PBS group, 6 of 29 animals (21%) were removed (censored) prior to the end point of death for biochemical and histological studies, whereas 8 of 32 (25%) in the 12d KO TAT–FXN group were censored for the same reasons and are accounted for in the Kaplan–Meier analysis.

The survival rate of animals was examined by determining the number of animals surviving 10% longer beyond the mean age of death of the untreated 12d Fxn-KO mouse of 28.1 ± 1.4...
days. Figure 4C shows that the TAT–FXN-treated animals in both the 3d and 12d groups had significantly higher survival rates to 31 days of age and beyond. Of the 3d group, 69% of the TAT–FXN group survived to 31 days and beyond compared with only 27% of the PBS-treated group (P = 0.022). Of the 12d group, 76% of the TAT–FXN-treated animals survived to 31 days and beyond compared with 33% of the untreated Fxn-KO animals (P = 0.002). The 12d groups were adjusted for censoring prior to 31 days, which lowers their total numbers when compared with (B). (D) Two-month-old heterozygous female mice were injected with PBS (100 μl/injection), TAT–FXN or TAT-mMDH-eGFP, each at 2 mg/kg/week (total protein) for 2 months per IP route. Tissues were fixed in 10% formalin and stained by hematoxylin and eosin.

control heterozygous mice with PBS injections predictably had no deaths and were significantly different from all Fxn-KO mice groups.

Finally, histology of organs was performed as a control to evaluate possible tissue inflammatory response to chronic IP injections of TAT-fusion proteins. Two different TAT-fusion proteins, TAT–FXN and TAT-mMDH-eGFP (38), and the PBS carrier fluid were injected into adult 2-month heterozygous control female mice twice weekly for 2 months. There was no evidence of inflammation in the liver, brain or heart as shown by the lack of inflammatory cells (Fig. 4D).
Survival curves of groups were analyzed with pairwise multiple comparison procedures (Holm–Sidak method) using an overall significance of $P = 0.05$. Growth velocity as a percentage of body weight was calculated for the three groups (12d) using the two-point average weight model (reviewed in 65) across the time period of 7–10 days after injections with TAT–FXN were started in the treated group. We tested the null hypothesis that the growth between treated and untreated Fxn-KO mice would be the same. Growth velocity is most commonly expressed as g/kg body weight/day, but was modified for mice to reflect their smaller weight and is expressed as g/g starting weight/day × 100%.

The results were analyzed by ANOVA for significance with Dunn’s method for pairwise multiple comparisons (Fig. 5). The control littermate animals experienced a mean growth velocity of 4.67 ± 1.97% per day. The median for this group was 4.38% with a range of 3.39% (25%) to 5.96% (75%). By comparison, the KO TAT–FXN (12d) animals grew significantly slower than controls with a mean growth velocity of 1.97 ± 3.36% per day ($P < 0.05$). The median of this group was 1.55% with a range of 0.21% (25%) to 4.178% (75%). Notably, however, the growth velocity of the KO TAT–FXN (12d) animals was significantly higher than the KO no PBS (12d) ($P < 0.05$) animals. In agreement with the previous characterization of the model (33), the KO no PBS (12d) animals had negative growth velocities across this time frame with a mean of $-1.04 ± 1.92%$, indicating that they were losing weight rather than gaining as seen in the other two groups. The median growth velocity of the KO no PBS (12d) group was $-0.465%$ with a range of $-2.368%$ (25%) to 0.455% (75%). Finally, the growth velocity of the KO no PBS (12d) was significantly lower than that of the control group ($P < 0.05$).

Table 2 shows the characteristics of these three groups. There were no differences in the starting or ending ages during which these measurements were made, indicating that the animals were at equivalent periods of growth in their lives. As expected, the starting weight of the control animals was significantly higher ($P < 0.05$) than either of the KO TAT–FXN (12d) or of the KO no PBS (12d) animals, but there was no significant difference between the starting weights of the two groups of Fxn-KO animals. All of the control animals gained weight (% positive weight gain) and 79% of the KO TAT–FXN (12d) animals gained weight ($P = NS$). However, only 30% of the KO no PBS (12d) animals showed any weight gain across this period, which was significantly lower ($P < 0.05$) than either the KO TAT–FXN (12d) group (indicated by the ‘*’)) or the control group (indicated by the ‘**’). At the end of the study, the control group animals had a mean weight of 11.3 ± 1.78 g, which was significantly higher than either KO TAT–FXN (12d) (6.8 ± 1.90 g) or KO no PBS (12d) (5.7 ± 1.68 g) animals ($P < 0.001$ for both). The KO no PBS (12d) animals were significantly smaller than the KO TAT–FXN (12d) animals at the end of the measurement period ($P < 0.016$).

**TAT–FXN increases the activity of aconitase in the heart of the Fxn-KO mouse**

Aconitase is a mitochondrial enzyme with Fe–S clusters. The loss of FXN has been shown to decrease its activity in both cell culture models (66), and in vivo (33,67). We tested the hypothesis that Fxn-KO animals treated with TAT–FXN would
have a higher activity of aconitase in their hearts when compared with untreated KO animals.

Using three age-matched animals in each group: heterozygous littermates given PBS as controls (Ctl), KO TAT–FXN (3d) and KO + PBS (3d), total aconitase activity was determined from heart homogenates by monitoring the formation of NADPH by isocitric dehydrogenase following the aconitase-dependent conversion of citrate to isocitrate (Cayman Chemical). In Figure 6A, the specific activity of aconitase from each group of animals shows that TAT–FXN is able to reconstitute activity of aconitase in the KO TAT–FXN group. The specific activity of aconitase from the heterozygous littermate control group, Ctl PBS (3d), was 37.52 ± 8.54 nmol/min/mg protein, and that of the KO TAT–FXN (3d) group was 31.36 ± 5.19 nmol/min/mg protein (P = NS). In contrast, the KO PBS (3d) group was significantly lower than both the Ctl and KO TAT–FXN (3d) groups at 18.91 ± 5.55 nmol/min/mg protein.

Western blotting was performed where the same samples were assayed for mitochondrial aconitase (Fig. 6B). This shows that mitochondrial aconitase protein mass is increased in the samples from the KO PBS (3d) group when compared with the Ctl and KO TAT–FXN groups. Densitometry of the Ctl PBS group averaged 7428 ± 358 pixels, the KO TAT–FXN group averaged 6787 ± 358 pixels, and the KO PBS group was 9068 ± 1292 pixels. There was significantly greater mitochondrial aconitase protein mass in the KO PBS group than in the KO TAT–FXN group (P = 0.04), and there was no significant difference between the Ctl PBS and KO TAT–FXN groups. Taking Figure 6A and B together, mitochondrial aconitase protein mass is increased in the KO PBS mice while total aconitase activity is decreased. In contrast, in the KO TAT–FXN group, aconitase activity and protein mass are restored to near Ctl levels. These data are consistent with recent findings showing that mitochondrial assembly of Fe–S clusters is necessary for cytosolic Fe–S-dependent proteins (68–70), as well as mitochondrial Fe–S-dependent enzymes.

To show that human FXN was present and processed in the hearts from the TAT–FXN-treated Fxn-KO mouse (Fig. 6C), western blotting was performed using whole heart homogenates from KO TAT–FXN (12d) mice (n = 3, lanes 1–3), KO no PBS (12d) mice with no injections (n = 3, lanes 4–6), and Ctl no PBS (12d) littermates (n = 3, lanes 7–9) with no injections. After separation on 12% SDS–PAGE and blotting, the membranes were probed using monoclonal antibody to human FXN (MitoSciences), which recognizes mouse FXN poorly. Lane 0 was loaded with the purified, precursor TAT–FXN that was injected and serves as a positive control. The Fxn-KO mice injected with TAT–FXN (lanes 1–3) demonstrate a strong signal at the estimated Mₘ of processed FXN of ~15 kDa. In contrast, there is essentially no signal in the Fxn-KO animals who did not receive

Table 2. Growth characteristics of mice

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Start Wt</th>
<th>End Wt</th>
<th>Start Age</th>
<th>End Age</th>
<th>% Positive Wt gain</th>
</tr>
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<tbody>
<tr>
<td>Controls</td>
<td>25</td>
<td>8.20 ± 1.88 g</td>
<td>11.30 ± 1.78 g</td>
<td>13.1 ± 3.1d</td>
<td>21.6 ± 2.5d</td>
<td>25/25 (100%)</td>
</tr>
<tr>
<td>Treated</td>
<td>24</td>
<td>5.92 ± 1.64 g</td>
<td>8.84 ± 1.90 g</td>
<td>12.7 ± 3.3d</td>
<td>21.4 ± 2.9d</td>
<td>19/24 (79%)</td>
</tr>
<tr>
<td>Untreated</td>
<td>20</td>
<td>6.11 ± 1.49 g</td>
<td>5.71 ± 1.68 g</td>
<td>14.3 ± 2.7d</td>
<td>23.2 ± 2.9d</td>
<td>6/20 (30%)</td>
</tr>
</tbody>
</table>

Values are the group means ± standard deviation.

**Significantly different from the control group (P < 0.05).*

**Significantly different from the treated group (P < 0.05).**
Table 3. Cardiac function in controls, treated KO and untreated KO mice

<table>
<thead>
<tr>
<th>Group</th>
<th>Age, days</th>
<th>EF, %</th>
<th>FS, %</th>
<th>SV, µl</th>
<th>CO, ml/min</th>
<th>LVIDs, mm</th>
<th>LVIDd, mm</th>
<th>LVPWs, mm</th>
<th>Ao peak, mm/s</th>
<th>E–A ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ctl no PBS ± STDEV</td>
<td>33 ± 13.7</td>
<td>69.36 ± 8.65</td>
<td>38.62 ± 6.87</td>
<td>30.81 ± 7.47</td>
<td>10.96 ± 3.58</td>
<td>2.03 ± 0.35</td>
<td>3.30 ± 0.35</td>
<td>0.97 ± 0.18</td>
<td>0.58 ± 0.12</td>
<td>875.84 ± 172.75</td>
</tr>
<tr>
<td>KO TAT–FXN ± STDEV</td>
<td>32.2 ± 12.6</td>
<td>38.69 ± 14.47</td>
<td>18.50 ± 7.91</td>
<td>14.64 ± 3.30</td>
<td>5.06 ± 1.17</td>
<td>2.59 ± 0.49</td>
<td>3.16 ± 0.32</td>
<td>0.75 ± 0.14</td>
<td>0.62 ± 0.12</td>
<td>536.95 ± 114.11</td>
</tr>
<tr>
<td>KO no PBS ± STDEV</td>
<td>26 ± 4.1</td>
<td>40.99 ± 17.75</td>
<td>20.70 ± 9.56</td>
<td>14.15 ± 6.51</td>
<td>3.19 ± 1.69</td>
<td>2.47 ± 0.69</td>
<td>3.06 ± 0.57</td>
<td>0.78 ± 0.18</td>
<td>0.59 ± 0.17</td>
<td>374.11 ± 146.96</td>
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</tbody>
</table>

Number in each group is in parenthesis, and values are the group means ± standard deviation. Mitral valve inflow 'E'-wave to 'A'-wave ratio (E–A ratio) were measured. EF, ejection fraction; FS, fractional shortening; SV, stroke volume; CO, cardiac output; LVIDs, left ventricular internal diameter in systole; LVIDd, left ventricular internal diameter in diastole; LVPWs, LV posterior wall; Ao peak, aortic peak velocity.

*Significantly different from the control group (P < 0.05).

**Significantly different from the treated group (P < 0.05).
Finally, the mitral Doppler column (Fig. 7A) shows the velocity of blood flow into the left ventricle during diastole from the left atrium across the mitral valve. A normal transmitral flow pattern and velocity was present in the control animal where the ratio of the height of the E-wave to the A-wave (E–A ratio) was 1.5, which is associated with normal inflow into the left ventricle. A similar mitral inflow pattern was seen in the KO TAT–FXN animal. In contrast, the mitral inflow pattern in the KO no PBS animal showed absent E-waves with only A-waves remaining indicating impaired diastolic function of the left ventricle.

The E–A ratios were measured for 6 control animals (2.1 ± 0.51), 5 KO TAT–FXN animals (2.6 ± 1.07) and 11 KO no PBS animals (1.1 ± 0.86), and compared using ANOVA (Table 3). Notably, the E–A ratio was significantly higher in the KO TAT–FXN animals when compared with the KO no PBS animals (P = 0.003), higher in the control animals when compared with the KO no PBS animals (P = 0.039), and there was no significant difference between the controls and KO TAT–FXN animals (P = 0.275). The E–A ratio was >1.5 in all of the control animals (100%), 80% of the KO TAT–FXN animals and only 36% of the KO no PBS animals. The lower E–A ratios in the KO no PBS animals strongly suggest left ventricular diastolic dysfunction and a greater reliance on atrial systole to fill the left ventricle (72). These findings of decreased transmitral inflow and aortic Doppler velocities are similar to patients with FRDA in which myocardial velocity gradients during both systole and early diastole have been shown to be reduced (73).

The heart rates of the KO TAT–FXN animals were also significantly higher than the KO no PBS animals. A representative electrocardiogram (EKG) from Ctl no PBS (12d), KO TAT–FXN (12d) and KO no PBS (12d) animals is shown in Figure 7B, and quantitative data are shown in Figure 7C. The heart rates of the control and KO TAT–FXN animals were both 329 bpm, whereas the KO no PBS animals was 197 bpm. Figure 7C shows that the mean heart rates between controls

Figure 7. Echocardiography of FRDA animals. (A) Ctl = heterozygous littermates, no PBS (12d). KO TAT–FXN = FXN-KO-treated with TAT–FXN (12d). KO no PBS = FXN-KO without PBS (12d). The M-mode column shows the internal dimension of the left ventricle between the interventricular septum (IVS) and left ventricular posterior walls (LVPW). Scale in millimeters is on the right, and EKG is on the bottom. The aorta column shows the Doppler velocity of blood flow in the ascending aorta for all three groups. Note that the velocity scale for the Ctl animal is 1 m/s, whereas it is cm/s for the other two groups. The mitral column shows the Doppler mitral inflow pattern and velocity for all three groups. E-wave and A-wave are marked. The velocity scale on the right is cm/s for the Ctl and KO no PBS animals, and 1 m/s for the KO TAT–FXN animal. (B) The EKG from a representative animal in each of the three groups is shown. The heart rate was determined from the EKG of each mouse at the same time point during echocardiography. Note that the time scale (bottom) is in milliseconds. (C) The mean heart rate of all three groups analyzed by one-way ANOVA with pairwise multiple comparison procedures (Holm–Sidak method).
and KO TAT–FXN (349 ± 41) animals were not significantly different, but the KO no PBS mice (226 ± 59) were significantly lower than KO TAT–FXN or control mice (P < 0.001 for both comparisons). Although the measures of systolic function, such as EF, FS and SV, were not significantly different between KO TAT–FXN and KO no PBS animals, the higher heart rate and improved diastolic filling of the KO TAT–FXN animals translated into a significantly higher cardiac output (CO) when compared with the KO no PBS animals (P = 0.02).

In summary, these data show that the Fxn-KO animals treated with TAT–FXN had a higher CO due to a higher heart rate than the KO no PBS animals. Additionally, the KO TAT–FXN animals had significantly better diastolic function (higher E–A ratios) than the KO no PBS animals although their systolic function was not significantly different.

TAT–FXN maintains cardiac ultrastructure and reduces cardiomyocyte apoptosis in the Fxn-KO mice

Previous data had shown that cardiomyocyte ultrastructure was severely disturbed in the Fxn-KO mouse heart (33). In particular, there was marked proliferation of mitochondria, loss of myofibrils and disruption of the normal mitochondria–sarcomere relationship. We tested the hypothesis that hearts from animals treated with TAT–FXN would have normal ultrastructure as measured by mitochondrial number and appearance using electron microscopy (EM), and normal
myofibril content and relation to mitochondria. Three groups of mice, Ctl PBS (3d), KO TAT–FXN (3d) and KO PBS (3d), were studied at 28 days of age and the results are shown in Figure 8.

In panels A.1–3 are electron micrographs from three representative animals which include controls (A.1), TAT–FXN-treated Fxn-KO mice (A.2) and PBS-treated Fxn-KO mice (A.3). The control animal demonstrates normal mitochondrial morphology and numbers. The relationship of mitochondria to myofibrils is well ordered with virtually all mitochondria touching a myofibril and roughly one mitochondrion per sarcomere (74). In panel A.2, the TAT–FXN-treated Fxn-KO mouse heart shows essentially the same morphology as a normal control. Mitochondria are evenly distributed among the myofibrils and most of them measure one sarcomere in length. The number of sarcomeres per field is approximately the same as the controls. In contrast, the heart from a PBS-treated Fxn-KO mouse (A.3) shows markedly disrupted myofibril structure with very few sarcomeres per field, extensive dysmorphic mitochondrial proliferation and a loss of mitochondria to myofibril relationship. The mitochondria have widely varying sizes.

To quantify the results of EM imaging, planimetry of mitochondria and myofibrils was performed to calculate their respective cumulative areas on EM micrographs from multiple hearts (74,75). The results are expressed as a mitochondria-to-sarcomere ratio in Figure 8B. Both the controls (n = 5) and TAT–FXN-treated Fxn-KO mouse hearts (n = 4) have low ratios (<1) that are not statistically different from each other. In contrast, the PBS-treated Fxn-KO hearts (n = 4) have a much higher ratio (>3) of mitochondria to sarcomeres that is statistically significant when compared with both the controls and TAT–FXN-treated mice. This was confirmed by performing western blotting of heart tissue from all three groups to compare expression of a mitochondrial Complex I protein, NDUFA9, with expression of a contractile protein in heart, α-actinin. The results are shown in Figure 8C, and demonstrate that there is no significant difference in the ratio of mitochondria to sarcomeric protein mass in the control and KO TAT–FXN (3d) mouse heart. In contrast, the KO PBS (3d) mouse had significantly higher mitochondria to sarcomeric protein ratio when compared with the control or KO TAT–FXN (3d) mice.

To evaluate the heart for programmed cell death in the Fxn-KO mouse, hearts from Ctl PBS (3d) (n = 5), KO TAT–FXN (3d) (n = 4) and KO PBS (3d) (n = 4) were matched for age (26 days of life ± 2 days) and stained for caspase-3 activation as a marker of apoptosis. Cardiomyocytes positive for activated caspase-3 were identified based on rod-shaped morphology and staining, and quantified as described previously (76) (Fig. 8E). For each heart, five transverse sections across the left ventricle were counted completely and are plotted in Figure 8D. As expected, control mice had very low numbers of caspase-3-positive cardiomyocytes (1.05 × 10⁻⁵/μm² ± 1.26 × 10⁻⁵). In contrast, the KO PBS (3d) mouse hearts had significantly higher numbers of caspase-3-positive cardiomyocytes (1.37 × 10⁻⁴/μm² ± 5.11 × 10⁻⁵) when compared with control hearts (P < 0.001). Strikingly, the KO TAT–FXN (3d) hearts were not different from the controls (8.48 × 10⁻⁶/μm² ± 9.9 × 10⁻⁶, P = NS), and were significantly less than the untreated KO PBS (3) hearts (P < 0.001).

Thus, the mitochondrial relationship to myofibrils is severely disrupted in the untreated Fxn-KO mouse heart, but is normalized by treatment with TAT–FXN. In concert with this finding, programmed cell death in the Fxn-KO mice treated with TAT–FXN is indistinguishable from the control hearts, and both are significantly lower than untreated Fxn-KO mouse hearts. These findings strongly support the conclusion that TAT–FXN fusion protein has rescued the cardiac function resulting in an increased lifespan and survival rate in the treated Fxn-KO mice.

**DISCUSSION**

The key finding from these experiments is that a cell-penetrant peptide is capable of delivering a replacement protein to mitochondria in vivo in amounts sufficient to rescue a very severe (fatal) disease phenotype. This has not been accomplished before and, in conjunction with other studies showing TAT-fusion proteins can restore mitochondrial enzyme activity in heterozygous mice (40), strongly supports the use of cell-penetrant peptides as a platform for developing novel therapeutic interventions for mitochondrial diseases. Mice homozygous for a conditional deletion of the Fxn gene lived substantially longer and had an increased rate of survival when treated with TAT–FXN when compared with the untreated Fxn-KO group. Mature processed human FXN was recovered from Fxn-KO animals injected with TAT–FXN, indicating that the TAT–FXN reached the mitochondrial matrix. Analysis of cardiac function in these mice revealed that the treated animals had better diastolic function and a higher heart rate, which resulted in a higher CO even though both had evidence of impaired systolic function. This is important because diastolic dysfunction is a component of the heart disease of FRDA patients (73,77–79) and improvements in this parameter would be expected to improve heart failure and survival. The growth velocity was also improved in the treated animals. This was true even when dosing began later in life, i.e. 12 days of age, which has analogy to the typical age at diagnosis in humans (10). Histologic analysis of heart showed that treated KO animals maintained normal ultrastructure and had less apoptotic events than the untreated KO animals. Additionally, treated KO animals had higher levels of total aconitase activity in the heart than in untreated animals, showing that protein replacement using TAT–FXN is capable of reconstituting Fe–S cluster-dependent enzymes. This also demonstrates that even though protein replacement with TAT–FXN was started later, it was still capable of rescuing the KO phenotype to extend the lifespan and survival.

There are substantial hurdles to delivering replacement proteins to mitochondria in vivo which are addressed by the use of cell-penetrant peptides. Mitochondria contain their own genome, ~16 500 bp in humans, which encodes 13 proteins and the transfer and ribosomal RNAs needed to translate them (80). The remaining hundreds of proteins needed for efficient mitochondrial function are encoded by the nuclear genome and imported in a multi-step energetic process from the cytosol (81). Proteins are essentially denatured as they
pass through the import apparatus and also do not contain post-translational modifications, such as glycosylation. As a result, mitochondria are well adapted to re-folding and processing proteins to generate active enzyme complexes. Thus, mitochondria present unique challenges to the development of therapies that address deficient or aberrant protein function, but also have unique features, such as the processing peptidases to remove a fusion protein tag like TAT, which can be leveraged to deliver an exogenous protein in designing a potential therapy.

Our approach took advantage of the fact that the MPP will proteolytically remove the MTS and any peptide sequence upstream of the MTS. This is important because if the TAT moiety remains attached, it is equally capable of transducing the fusion protein out of the mitochondria and cell (38). With processing by the MPP of the TAT leader sequence, the mitochondrial matrix ultimately sees the native mature FXN protein. The current data show that the precursor is ultimately cleaved at position FXN80-81 to yield the start of the mature peptide as 81SGTLGH (29). Our results identified the intermediate cleavage site at 82LRTDI when cleaved in vitro consistent with the description by Cavadini et al. (27). We were unable to generate in vitro sequence information by Edman degradation for the mature fragment consistent with earlier findings that the rate of the second cleavage in vitro is much slower. However, a mature fragment is generated and detected by western blotting, as shown in Figure 2B. This is in good agreement with the mature peptide predicted by Schmucker et al., as being FXN81-210 (29). Based on these data, it is logical to conclude that TAT–FXN is being properly recognized and cleaved by the native MPP.

The finding that the TAT–FXN-treated Fxn-KO animals had higher heart rates is significant. Neuron-specific enolase (NSE) is expressed in multiple regions of the heart including paraganglionic-like structures at high density near the upper third of the atrial septum (82,83). In addition, NSE has also been identified in multiple cell types, including nerve fibers, atrial adipose tissues, cardiomyocytes, intramyocardial paraganglia, which are closely related to myocardial fibers (82,84) and the conducting system of heart (85). Finally, the expression of NSE is developmentally regulated and is present in heart early during embryogenesis (62,63). Thus, NSE-Cre expression would be predicted to disrupt Fxn gene expression very early in heart and in multiple cell types, including innervations of the heart. Replacement of FXN would logically be expected to allow greater sympathetic response to cardiac innervation, which, when combined with improved diastolic filling of the ventricle, would result in higher CO.

It was interesting that the TAT–FXN-treated animals did not demonstrate complete rescue, even when treatment was begun on the third day of life. The TAT–FXN-treated Fxn-KO animals did not reach the same growth rate as the controls, nor live as long. Given that the promoter (NSE) driving the Cre expression is on in mid-embryogenesis (E10.5) in the brain (86,87) and heart (62,63), this also means that these animals had a congenital absence of FXN protein in selected tissues, and their disease phenotype had a substantial time to develop prior to birth. Earlier studies had shown that constitutive ablation of the Fxn gene in the mouse was embryonic lethal emphasizing the importance of FXN protein early in development (60). Thus, one possibility for why complete rescue was not achieved is that substantial organ damage may already have occurred by the time treatment was initiated. This conclusion is supported by reports of children with the FRDA genotype who died very young from other causes prior to onset of symptoms and suggests that organ damage may be present from birth (88). Initiation of TAT–FXN prenatally in the mouse may, therefore, achieve greater rescue. Alternatively, there may be an inadequate FXN protein mass in critical tissues, such as the heart and brain, to achieve complete rescue, or else human FXN cannot substitute completely for mouse FXN. Finally, although no tissue immune response was identified, it is possible that humoral immune response eventually develops as has been reported for TAT-purine nucleoside phosphorylase (41), and may decrease effectiveness of injected TAT–FXN. As reported by Toro and Grunebaum, however, it was notable that even though the mouse mounted an immune response to TAT-purine nucleoside phosphorylase, the fusion protein was still quite effective at restoring enzyme activity.

It is also important to note that the animal model used in these studies is not an exact duplicate for either the phenotype or the genotype of patients with FRDA. Typically, patients will have <20% of normal FXN levels in affected tissues but not complete loss of FXN (89), and do not manifest overt symptoms until adolescence. In contrast, the mouse model used in these studies has a complete loss of FXN in those tissues expressing NSE and presents a very severe phenotype beginning at birth. These include tissues such as the brain, dorsal root ganglia, and other parts of the nervous and neuroendocrine systems, as well as the heart (85). However, using this model is a stringent test of the hypothesis that it is not necessary to replace FXN to normal levels to achieve a rescue of the phenotype. We were able to achieve an increased lifespan and restoration of enzymatic function in these animals with minimal dosing of TAT–FXN. Because enzymatically active TAT-fusion proteins have been shown to cross the blood-brain barrier (90–93) and our preliminary data have shown the presence of TAT–FXN in the brain of these treated animals (unpublished data), we would predict that future studies will show an improved neurologic function in treated animals. Studies of human FXN levels suggest that partial replacement of FXN may be enough to restore an adequate cellular function (94). If true, then partial replacement of FXN in FRDA patients with low, but measurable amounts of FXN may be adequate to restore normal cellular function. This would be especially important if tissue damage from the loss of FXN is cumulative and would justify early screening to initiate treatment(s).

METHODS AND MATERIALS

Detailed Methods and Materials are contained within the Supplementary Material.

TAT–FXN cDNA construction, protein expression and purification

The cDNA for human precursor FXN (GenBank accession NM_000144) was obtained from Invitrogen (clone ID
mMDH–eGFP and TAT–GFP fusion proteins were expressed exchanged into PBS for injection into animals. The TAT–FXN was purified using nickel affinity chromatography, and cells for expression. The soluble fraction from expression TAT–FXN (a gift from the Steve Dowdy lab, Washington University).

The complete His–TAT–FXN cDNA construct (termed containing the TAT sequence at the N terminus along with a tryptophan to a glycine, and was cloned in-frame into the NcoI–EcoRI sites of a bacterial expression vector containing the TAT sequence at the N terminus along with a 6× His tag at the N terminus to allow affinity purification (a gift from the Steve Dowdy lab, Washington University). The complete His–TAT–FXN cDNA construct (termed TAT–FXN) was transformed into BL21(DE3)pLysE cells for expression. The soluble fraction from expression was purified using nickel affinity chromatography, and exchanged into PBS for injection into animals. The TAT–mMDH–eGFP and TAT–GFP fusion proteins were expressed and purified as described previously (38) with minor modifications.

Expression and purification of mitochondrial processing peptidase
Plasmids containing the coding sequences of the mature α-MPP and β-MPP for the yeast were a kind gift from Drs. Jiri Adamec and Henry Weiner (both of Purdue University) and have been described as pETYA and pETYB, respectively (44). Both plasmids were subcloned into the pET-19b vector for expression and purification using a His affinity tag, and transformed into BL21(DE3) cells for expression and purification by nickel affinity chromatography.

Sequencing of TAT-fusion proteins
TAT-fusion proteins were incubated with MPP in a 1:1 ratio (e.g. 1 μg MPP total protein with 1 μg TAT–FXN total protein) at 37°C in a modified Factor XA cleavage buffer (20 mM Tris, pH 7.7, 1 mM CaCl2, 50 mM NaCl, and 1 mM β-mercaptoethanol). Reaction products were separated by SDS–PAGE, transferred to Immobilon membranes and sequenced at the Iowa State University Protein Facility. Sequences obtained by Edman Degradation were compared with published sequences for intermediate and mature forms of human FXN (27) and rat mMDH (95,96).

Fluorescent labeling of TAT–FXN and localization in mitochondria
TAT–FXN was labeled with 5-IAF. Labeled protein was separated from unreacted salts on a PD-10 column with buffer exchange into PBS. FRDA fibroblasts were treated with 10 μg/ml of fluorescein-labeled TAT–FXN for 3 h, and then incubated with fresh media without labeled TAT–FXN for 120 h. At the time of microscopy, the cells were incubated with 200 nM of the mitochondrial dye CMXRos for 30 min at 37°C and imaged live on a Bio-Rad MRC-1024 laser scanning confocal inverted microscope.

Iron-binding activity of TAT–FXN protein in cell-free system
An aliquot of 5 μM each of ferrous sulfate and HQ (Fe/HQ) was incubated with PBS in the presence and in the absence of TAT–FXN (20, 40 μg/ml), or BSA (20 μg/ml) (negative control). An aliquot of 5 μM of DCHF-DA was hydrolyzed to DCHF by adding 200 mM NaOH and incubating the mixture in the dark for 30 min on ice. After 30 min, 250 μM of Na2HPO4 was added to neutralize the excess NaOH. An aliquot of 10 μl of this DCHF mixture was added to 200 μl of the reaction mixture to obtain 0.25 μM DCHF in the final assay. The fluorescence of DCF was measured at excitation wavelength of 485 nm and emission wavelength of 530 nm on a SpectraMax 340pc microplate spectrophotometer. As a positive control, an incubation containing 100 μM of EDTA as an iron chelator was performed to evaluate the role of iron in the oxidative mechanism of HQ.

Rescue of FRDA fibroblasts from oxidative stress by TAT–FXN
FRDA fibroblasts, and fibroblasts from a healthy age- and sex-matched control, were treated with and without TAT–FXN (20 and 40 μg/ml), or an equal volume of carrier fluid (PBS) as a negative control for 5 h, washed with PBS and then cultured overnight in culture media. The cells were then washed twice with PBS followed by the addition of Fe/HQ (5 μM of each component) in culture media for 5 h as an oxidant stress. Control cells were not treated with Fe/HQ. Following the 5 h treatment, cells were photographed under light microscopy for evidence of cytotoxicity.

Caspase-3 determination in TAT–FXN-treated FRDA fibroblasts
FRDA and age-/sex-matched control fibroblast cells were treated with and without TAT–FXN (20 and 40 μg/ml) in culture media for 5 h as an oxidant stress. After incubation, the cells were washed twice with PBS and then treated with and without Fe/HQ (5 μM of each component) in culture media for 3.5 h. After the plates were washed with PBS, the cells were scraped from the plate in 1 ml PBS, centrifuged and the cell pellet was mixed in cell lysis buffer per the manufacturer’s instructions. The protein content of each sample was estimated and 750 μg of protein from each sample condition was loaded in a 96 black well, flat bottom, polystyrene assay plate and caspase-3 levels in each condition were quantified using the fluorescent substrate, 7-amino-4-methylcoumarin, at an excitation wavelength of 342 nm and emission wavelength of 441 nm.

FXN conditional KO animals, dosing and survival analysis
All animal protocols were approved by the Institutional Animal Care and Use Committee at Indiana University School of Medicine. Mice were bred for conditional deletion of the Fxn gene as described with minor modifications (33,87). Briefly, NSE-Cre mice were crossed with mice homozygous for a conditional allele of Frdx (Frdx1/L3) to generate mice heterozygous for the conditional allele.
carrying the NSE-Cre transgene (Frda<sup>1L3+/−;NSE-Cre</sup>). These Frda<sup>1L3+/−;NSE-Cre</sup> mice were then crossed with Frda<sup>1L3/L3</sup> mice to generate the final genotype with deletion of the Fxn gene in tissues expressing NSE. Genotyping of these animals was performed using oligonucleotide primers as described by Puccio et al. (33). Mice were dosed according to body weight with a total volume of ~20 μl/g of weight given IP. The dose interval was based on the published T1/2 of 50 h for FXN (97). Survival curves for the mice were calculated using the Kaplan–Meier curve as described previously (64).

Electron microscopy

Tissue sections (~1–2 mm<sup>3</sup> volume) were fixed in modified Karnovsky’s solution with 2% paraformaldehyde/2% glutaraldehyde in 0.1 M phosphate buffer, with post fixation in 1% osmium tetroxide in phosphate buffer for 1 h. The tissues were embedded in resin for sectioning. Sections were imaged on a Tecnai G2 12 Bio Twin transmission electron microscope at 80 kV at the Electron Microscopy Center of Indiana University School of Medicine.

Histology

Hearts were cryoprotected in 30% sucrose, embedded and sectioned at 6 μm thickness using standard techniques. Five transverse sections from each heart, sampled from the mid-point between the apex and base, were post-fixed in 4% paraformaldehyde and screened for anti-activated caspase-3 immunoreactivity, followed by a horseradish peroxidase-conjugated secondary antibody. The signal was visualized with a diaminobenzidine reaction as described previously (98).

Echocardiography

Control littersmates, and Fxn-KO animals treated with, or without TAT–FXN, underwent echocardiography at 10–14d after initiation of injections with TAT–FXN in the KO-treated group. A 40 MHz hand-held mechanical transducer containing both imaging (B-mode) and Doppler transducers with a frame rate of 34 Hz was used to image the heart in multiple views as described previously (99). VisualSonics software (Version 2.3.2) was used to calculate ejection and shortening fractions, stroke volume, ventricular dimensions, and interpret the Doppler interrogation of mitral and aortic valve flow.

Aconitase activity

Aconitase-specific activity was measured in whole heart homogenates of ventricular tissues based on the downstream generation of NADPH. Briefly, whole heart was homogenized on ice and the NADPH change was measured via the absorbance at 340 nm using the end-point method on the Spectramax M5. Rates of conversion were normalized to total protein to generate specific activity.

Statistical analysis

All calculations, analyses and graphs were performed using SigmaPlot version 12.0 (Systat Software, Inc.). Data are presented as mean (± SD) unless otherwise indicated. Statistical comparisons between the two groups were made using Student’s t-test with Mann–Whitney rank-sum test if the groups failed the normality test (Shapiro–Wilk), or equal variance test for normalized data. For comparisons between more than two groups, ANOVA and the Holm–Sidak method for multiple pairwise comparisons were used unless otherwise noted in the text. A P-value of <0.05 was considered to be significant.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

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