Biallelic mutations in the ferredoxin reductase gene cause novel mitochondriopathy with optic atrophy

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Abstract
Iron–sulfur (Fe–S) clusters are ubiquitous cofactors essential to various cellular processes, including mitochondrial respiration, DNA repair, and iron homeostasis. A steadily increasing number of disorders are being associated with disrupted biogenesis of Fe–S clusters. Here, we conducted whole-exome sequencing of patients with optic atrophy and other neurological signs of mitochondriopathy and identified 17 individuals from 13 unrelated families with recessive mutations in FDXR, encoding the mitochondrial membrane-associated flavoprotein ferredoxin reductase required for electron transport from NADPH to cytochrome P450. In vitro enzymatic assays in patient fibroblast cells showed deficient ferredoxin NADP reductase activity and mitochondrial dysfunction evidenced by low oxygen consumption rates (OCRs), complex activities, ATP production and increased reactive oxygen species (ROS). Such defects were rescued by overexpression of wild-type FDXR. Moreover, we found that mice carrying a spontaneous mutation allelic to the most common mutation found in patients displayed progressive gait abnormalities and vision loss, in addition to biochemical defects consistent with the major clinical features of the disease. Taken together, these data provide the first demonstration that germline, hypomorphic mutations in FDXR cause a novel mitochondriopathy and optic atrophy in humans.

Introduction
Iron–sulfur (Fe–S) clusters act as inorganic cofactors in many protein activities and are essential for multiple important biological processes, including electron transport in mitochondrial respiration, cofactor synthesis, substrate binding/activation, Fe–S storage, DNA repair, and gene regulation (1,2). Fe–S cluster biosynthesis is a tightly regulated process dependent upon the coordinated delivery of iron and sulfide, principally within the mitochondria of eukaryotic cells. Formation of Fe–S clusters involves the use of electrons from sulfur. Following the assembly of a Fe–S cluster on a scaffold protein, the cluster is transferred to an acceptor protein. A variety of human disorders have been associated with impaired Fe–S cluster biosynthesis, including neurodegenerative disorders (e.g. Friedreich ataxia and myopathy with lactic acidosis) (1). Iron homeostasis, which requires precise synthesis and localization of Fe–S clusters in mitochondria, is critical to ensure that there are sufficient, but not toxic levels of iron for cellular functions. Excessive levels of iron favor the formation of excess oxygen free radicals and consequent mitochondrial dysfunction. The optic nerves are especially vulnerable to such oxidative stress.

Ferredoxin-1 and -2 [FDX1 and FDX2 (a.k.a. FDX1L)], which are encoded by two homologous genes in mammals and expressed in mitochondria, serve as electron donors in Fe–S cluster biosynthesis (3,4). Mitochondrial membrane-associated ferredoxin reductase (FDXR) is a flavoprotein that initiates the mitochondrial electron transport chain reaction by transferring electrons from NADPH to the mitochondrial cytochrome P450 system via FDX1 or FDX2 (5). This pathway is also critical to steroid hormone biosynthesis (6). FDXR has been shown to be essential for cell viability and contributes to p53-mediated apoptosis through the generation of oxidative stress in mitochondria (7). FDX2, in particular, has been shown to act as an electron source in the electron transfer chain with NADPH and FDXR (8). Moreover, biallelic mutations in FDX2 have been identified in patients with metabolic myopathy who have deficient mitochondrial complexes I, II, and III and deficient aconitase, suggestive of mitochondrial Fe–S cluster-related defects (9).

We performed clinical exome sequencing in families with suspected mitochondriopathy, with the aim of identifying disease-causing genes. We report 17 individuals—from 13 independent families—with biallelic FDXR mutations, exhibiting a wide range of signs of mitochondriopathy. The affected individuals presented with neurological abnormalities, with most showing some degree of optic atrophy and neuropathy. Patient fibroblast cells showed deficient ferredoxin NADP reductase activity and mitochondrial dysfunction. Such defects were rescued by overexpression of wild-type FDXR. In a parallel analysis in the mouse model, we observed that mice with the Fdxr mutation allelic to the most common mutation among our patients recapitulated major aspects of the human phenotype, including progressive peripheral nerve impairments, decreased Fdxr enzyme activity, and mitochondrial dysfunction. Hence, we report a convergence of clinical, mouse model, and experimental findings suggesting that a biallelic mode of inheritance of mutant FDXR causes mitochondriopathy with optic atrophy.

Results
Clinical phenotypes associated with FDXR mutations
Among the 13 unrelated probands from diverse ethnic backgrounds with biallelic rare variants in FDXR (Fig. 1A), we identified 6 homozygotes and 7 compound heterozygotes (Table 1). As is common for mitochondrion-related diseases, patients presented with variable clinical phenotypes, consistent with mitochondriopathy. They were found to share the core clinical features of optic atrophy (Fig. 1C; Supplementary Material, Fig. S1A–C), ataxia, and hypotonia. Other common features in the cohort were neurological anomalies evidenced in MRI imaging, seizures, and global developmental delays. Other less common features included spasticity and respiratory failure. Several patients had abnormal movements and hearing loss. Six patients died in early age. The patients’ clinical features are summarized in Table 1 and additional detailed clinical information about the patients is provided in Supplementary Material, Note S1.

FDXR mutations are evolutionarily conserved
The FDXR protein contains two NAD(P)-binding domains and one FAD/NAD(P)-binding domain (Fig. 1B). In Family 1, we identified a homozygous rare variant, c.1174 C>T (p.R392W), an amino acid change in an evolutionarily conserved residue of the FAD/NAD(P)-binding domain of FDXR (Supplementary Material, Fig. S2A and B). Multiple sequence alignment was used to analyze amino acid conservation (Supplementary Material, Note S2). The FDXR mutations identified in other patients are also in the FAD/NAD(P)-binding domain with a hotspot at p.R392W, which accounts for 11 (42.3%) out of the 26 mutant alleles (Fig. 1B). This alteration is a relatively common Latino allele with 0.19%
minor allele frequency in this population (carrier rate 1/263; http://exac.broadinstitute.org/variant/17–72860036-G–A). Fourteen missense or nonsense FDXR mutations were identified in this study and eight of them (I143F, V158M, T211A, I213F, K280*, R315*, C359Y, D374N) clustered in the FAD binding site (Fig. 1B). Noteworthy, seven patients with homozygous R392W died at an early age, suggesting that this allele may have a more severe, detrimental impact on FDXR function. There were no biallelic truncating mutations identified in any patients and no homozygous loss-of-function alleles in gnomAD (http://gnomad.broadinstitute.org/), suggesting that complete loss of function of FDXR is likely incompatible with life.

Mutations of FDXR may cause loss of function due to protein instability

The FDX1/2 and FDXR proteins interact physically, and FDX1/2 mutations can affect binding to FDXR and/or cytochrome P450 (10). To investigate the possible functional consequences of FDXR mutations, we mapped all mutations to a crystal FDXR structure, based on the homologous FDXR structure (starting from blue at the N-terminal region, and ending at red in the C-terminus) for Bos taurus complexed with FAD (gold). The residue position of missense variants and starting point of truncations are indicated with red spheres.
### Table 1. Summary of clinical information

| Family | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 |
|--------|---|---|---|---|---|---|---|---|---|----|----|----|----|
| Gender | F | M | M | M | M | F | F | M | M | F | M | M | M |
| Genotype | p.R392W hom | p.R392W | p.F531V/p.G443S | p.R392W hom | p.R392W | p.P74L/p.R392W | p.R392W hom | p.T211A/p.G443C | p.R392W | p.V158M/p.P409L | p.R392W hom | p.R392W | p.V158M/p.K280* |
| Age examined | 1.6 yrs; d. 6 yrs | 4 yrs; d. 2.3 yrs | 5.4 yrs; d. 6 yrs | 1.3 yrs; d. 2 yrs | 1.8 yrs | 3 yrs; d. 1.6 yrs | 4 yrs | 1.8 yrs | 1.8 yrs | 15.5 yrs | 4 yrs | 2.5 yrs | 0.3 yrs |
| Microcephaly | No | No | No | Yes | Yes | Yes | Yes | Yes | Yes | No | No | No | Yes |
| Optic atrophy | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | No | Yes |
| Brain MRI | Normal | Normal | Normal | dCC e | abnl FL | Normal | Normal | dCC e | abnl FL | Normal | Normal | Normal | Normal |
| Seizures | No | No | No | No | Yes | Yes | Yes | Yes | No | No | No | Yes | Yes |
| EEG | Normal | Normal | Abnormal; dCF | Abnormal; severe encephalopathy | Normal | Abnormal; B&Sp | Normal | Abnormal; B&Sp | Normal | Abnormal; slow wave | Abnormal; clay wave | Normal |
| GDDq | Yes | Yes | No | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes |
| Regression | No | Yes | Yes | No | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes |
| Hypotonia | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes | Yes |
| Spasticity | Yes | Yes | Yes | Yes | No | No | No | No | No | No | No | No | No |
| Ataxia | Yes | MD | No | MD | No | No | No | MD | Likely | No | Yes | Yes |
| Mitochondrial Morphology | abnl shape, loss cristae | abnl shape | Normal | I-IV | abnl shape | Normal | I-IV | abnl shape | Normal | I-IV | abnl shape | I-IV 28% | abnl shape | I-IV 28% |
| Muscle ETC Results | deFG | deFG | deFG | deFG | deFG | deFG | deFG | deFG | deFG | deFG | deFG | deFG | deFG |

*a* abnormal corpus callosum, abnl CC  
*b* diffusion of the globi pallidi, dGP  
*c* diffusion of the midbrain, dMB  
*d* diffusion of the central and cortical atrophy, dCCA  
*e* decreased corpus callosum, dCC  
*f* chronic extra-axial collections in bilateral frontal convexities abnormal frontal lobe, abnl FL  
*g* increased cerebellar atrophy, inCA  
*h* decreased cerebral white matter, deCWM  
*i* decreased basal ganglia, deBG  
*j* decreased signals globus pallidus and substantia nigra, deGP & SN  
*k* abnormal basal ganglia and mesencephalic peduncles, abnl BG & MP signal  
*l* cerebellar atrophy, CA  
*m* abnormal white matter periventricular signal, abnl PV signal  
*n* infantile spasms and seizures, IS & Seizure  
*o* diffuse cerebral function, dCF  
*p* chronic background with splash, abnl B & S  
*q* globe development delay, GDD  
*r* abnormal movements, abnl MO  
*s* Movement disorder, MD  
*t* abnormal mitochondrial morphology, abnl shape  
*u* abnormal shape, abnl shape  
*v* elongated and abnormally shaped mitochondria, abnl shape  
*w* abnormal morphology, abnl shape  
*x* increased mitochondrial number, In#
of the mutations, I143F and D374N, appear to lie at a putative interface between FDXR monomers (12). The remaining mutations, namely T211A, I213F, C359Y, R392W, G443C, and G443S, were buried, at least partially; these mutations could disrupt functionality by causing a destabilization of the protein structure (Fig. 1D). Examination of the energies of these mutations (Supplementary Material, Table S2) indicated that C359Y, G443C, and G443S are likely to disrupt the folding of FDXR. However, the modes of disruption of T211A, I213F, and R392W are unclear. There is no indication that these positions would be expected to disrupt a posttranslational modification site or protein interaction site. Previous results suggested that the mutation in the N-terminus could have significant effects on FDXR biochemistry by increasing the K_d of FDXR with cytochrome P450. Importantly, these mutations could also decrease cholesterol side chain cleavage activity (10). Together, the mutations in FDXR could impair protein functions through multiple mechanisms, including protein-protein interactions, protein stability and binding affinity to the substrates.

To investigate if the mutations of FDXR affect the protein’s stability, we tested the FDXR protein level in patient fibroblast cells. As shown in Figure 2A and B, the protein levels of FDXR in two of the patients’ fibroblast cell lines that were tested decreased dramatically. To confirm this finding, we examined the expression and localization of FDXR in control and patients’ fibroblasts by confocal microscopy. In this experiment, mitochondrial localization was demonstrated by a signal overlap between FDXR (Fig. 2C and D, green) and mitochondria (Fig. 2C and D, red). As shown in Figure 2C and D, we found that FDXR levels are significantly lower in the patient fibroblast cells with the homozygous mutations R392W. While we found that FDXR localization was mainly in mitochondria, FDXR was also detectable in nuclei and in the cytoplasm, in both control and patient fibroblasts. 3D image analysis confirmed these
observations (Fig. 2C and D, Supplementary Material, Movie S1). Previous electron microscopy and immunogold labeling studies report that FDXR is sublocalized to the mitochondrial matrix (13) where it initiates the electron-transport chain and ultimately leads to the conversion of cholesterol to pregnenolone in the first rate-limiting step of steroid hormone biosynthesis (13,14). The significance of the presence of FDXR in nuclei and the cytoplasm is yet to be elucidated. Mitochondrial Fe–S cluster biogenesis machinery proteins, namely NFS1 and IDS11, were also observed in the cytoplasm, though the functional relevance of their presence also remains to be elucidated (15,16).

FDXR mutations cause mitochondrial dysfunction in patient fibroblast cells

To investigate the functional consequences of patient mutations and to demonstrate causation, we assessed FDXR enzyme activity and mitochondrial functions in cultured fibroblasts from individual II-2 in Family 1 (p.R392W/R392W) who is homozygous for the p.R392W mutation. To confirm causation, wild-type FDXR was overexpressed in the patient fibroblasts for complementation experiments (rescue) and expression was confirmed by immunofluorescence microscopy of tdTomato protein (Supplementary Material, Fig. S3A–C). As shown in Figure 3A, Ferredoxin NADP reductase activity in patient fibroblasts, non-transduced patient fibroblasts, and patient fibroblasts overexpressing FDXR (rescued) were 48%, 53%, and 94% respectively of the mean value measured for age and gender matched control fibroblasts. Basal OCR, a measure of cellular bioenergetics, in patient fibroblasts was 24% of that in control fibroblasts (100%, reference) (Fig. 3B and C). ATP production, an index of mitochondrial oxidative phosphorylation capacity, was significantly decreased in patient fibroblasts compared with that in control fibroblasts and rescued in patient fibroblasts transduced with FDXR (As shown in Fig. 3D). FDXR transduction of patient fibroblasts resulted in a partial rescue of basal OCR. Likewise, ATP-linked OCR, maximal OCR and reserve capacity OCR in patient fibroblasts were significantly lower than the rates observed for control fibroblasts and these reductions were also reversed in FDXR-transduced patient fibroblasts.
family 2 (P G443S/F51L), a compound heterozygote, showed the p.R392W mutation, showed major deficiency in complexes II-III and mild reduction of complex I and IV activities in the muscle fibroblasts were 95% and 98% of control levels, respectively. However, complex I and IV activities did not differ significantly among the various groups. Consistent with this observation, this patient’s deceased brother who is also homozygous for the FDXR mutations cause increased ROS production in patient fibroblasts

Fe–S clusters are components of the mitochondrial complexes I and III. Mutations of FDXR could impair Fe–S cluster biosynthesis and result in decreased complex I and complex III activity. However, our results suggest that mutations of FDXR cause a broader impairment of mitochondrial function. Accumulating evidence suggests that FDXR plays an important role in ROS production (17). Therefore, we tested ROS production in patient cell lines. Our results showed that although superoxide production was altered in patient fibroblasts, reactive oxygen species (ROS) levels did not differ (Fig. 4A). However, when ROS production was induced by introduction of pyocyanin ROS, superoxide production levels increased significantly in patient fibroblasts compared with the control fibroblasts (Fig. 4B). This effect was blocked in the presence of a ROS production inhibitor (Supplementary Material, Fig. S5). This result is consistent with previously published data, which demonstrate that FDXR can influence ROS-mediated apoptosis in epithelial cancer cells (13,17). Altogether, these findings indicate that mutations in FDXR could have a broader effect on mitochondrial functions through the ROS pathway, contributing to more general mitochondrial dysfunction.

Mitochondrial dysfunction is recapitulated in FdxrR389Q/R389Q mice

Taking advantage of a newly discovered mouse strain carrying a spontaneous mutation in Fdxr, we further examined the role of FDXR in peripheral neuropathy, mitochondrial function, and optic atrophy. These mice were previously reported to have a recessive gait phenotype and were shown by exome sequencing to harbor a homoygous missense mutation in Fdxr that results in an R389Q change in the encoded protein (corresponding to human FDXR p.R392Q and allelic to the common patient mutation p.R392W) (18). Using comprehensive gait analyses, we found that this mutation causes progressive impairment of posterior gait dynamics (Supplementary Material, Movie S2). To test FDXR and mitochondrial function in this mouse model, we conducted FDXR enzyme activity assays using tissues from homozygous FdxrR389Q/R389Q mice and WT littermate controls (Fig. 5A). Consistent with the human enzyme assays, we observed significantly reduced FDXR activity levels, relative to WT levels in both muscle (33 and 49%, respectively) and brain (39 and 45%, respectively). Similar results were observed in other tissues, including the liver and heart (Supplementary Material, Fig. S7A) which showed impaired complex I and complex III activities (Fig. 5C; Supplementary Material, Fig. S7C), and decreased ATP production (Fig. 5B; Supplementary Material, Fig. S7B).

Visual acuity defects are associated with loss of retinal ganglion cells (RGCs) and abnormal myelination of optic nerve in FdxrR389Q/R389Q mice

In addition to ataxia, the majority of the patients with FDXR mutations in this study had vision impairment or loss and optic atrophy (Table 1 and Supplementary Material, Note S1). To test if FdxrR389Q/R389Q mice recapitulate these clinical features, we tested visual acuity using functional optokinetics in adult mice (15–17 weeks of age). The optokinetic response scores from FdxrR389Q/R389Q mice were significantly reduced as compared with WT controls (Fig. 6A). We then performed a histopathological assessment of the retina. We counted the retinal ganglion cell (RGC) layer nuclei in H&E stained, paraffin sections of eyes of 2-month-old and 6-month-old WT and FdxrR389Q/R389Q mice.
No significant difference in the number of RGC layer nuclei between WT and mutant mice at the age of 2 months old was found (Supplementary Material, Fig. S8A). However, a significant reduction in the RGC layer nuclear number was observed in 6-month-old \(Fdxr^{R389Q/R389Q}\) mice, with a 54% loss of the RGC layer nuclei (Fig. 6B) as compared with WT mice. Transmission electron microscopy (TEM) of retina from 6-month-old mice confirmed RGC loss in \(Fdxr^{R389Q/R389Q}\) mice, and revealed abnormal cytoplasm, shrunken nuclei, and abnormal chromatin aggregates in remaining RGCs (Fig. 6F).

RGC axons in the optic nerve were also studied with TEM. No difference was found in the 2-month-old WT and \(Fdxr^{R389Q/R389Q}\) mouse RGC axons (Supplementary Material, Fig. S8B). However, a significant decrease in the thickness of the myelin membrane of RGC axons was found in 6-month-old \(Fdxr^{R389Q/R389Q}\) mice (Fig. 6C), indicating that the \(Fdxr^{R389Q/R389Q}\) mice had compromised myelination of RGC axons, probably related to defects in oligodendrocyte function. Also, mitochondria inside RGC axons of the \(Fdxr^{R389Q/R389Q}\) mice had structural damage, including cristae loss and the appearance of “ghost” mitochondria (Fig. 6C), which were consistent with the findings in the patient fibroblasts indicating that mutations of \(FDXR\) had broader effects on mitochondrial structure and functions.

**Discussion**

Synthesis of iron-sulfur (Fe–S) cluster scaffold proteins requires a coordinated delivery of iron, sulfur, and electrons. Although the iron donor for Fe–S clusters remains elusive, two Fe–S cluster assembly proteins, IscA (19) and frataxin (20), have been proposed as possible iron chaperones delivering iron for Fe–S cluster assembly in IscU (21). The electrons required for Fe–S cluster assembly in IscU are delivered by ferredoxin, which is reduced by ferredoxin reductase using NADPH as the electron donor (8,22). In humans, there are two ferredoxins: FDX1 and FDX2 (8). Both FDX1 and FDX2 appear to be important for Fe–S cluster biogenesis (4,8). Nevertheless, deletion of FDX1 in Hela cells did not elicit any defects in Fe–S cluster biogenesis (3), and mutation of FDX2 has been associated with a mitochondrial muscle myopathy, which is characterized by severely impaired activities of Fe–S proteins in mitochondria (9). Furthermore, while FDX1 is exclusively expressed in adrenal gland, FDX2 is ubiquitously found in virtually all tissues (3). Thus, FDX1 may participate in the synthesis of bile acid, vitamin D, and steroid hormones (8,22–24), while FDX2 is likely involved in Fe–S cluster biogenesis (3,22). Humans have one ferredoxin reductase (FDXR) (14), which is a FAD-dependent enzyme that reduces FDX1 and FDX2 using NADPH as the electron donor (8,22). Biallelic mutations in the \(FDXR\) gene (\(FDX1L\)) have been identified in patients with metabolic myopathy with deficient mitochondrial complexes I, II, and III and aconitase, suggesting mitochondrial Fe–S-related defects (9). Depletion of FDXR diminishes Fe–S cluster assembly and causes mitochondrial iron overload (8), indicating that FDXR has a crucial role in Fe–S cluster biogenesis in human cells.

Here, we identified biallelic rare variants in \(FDXR\) by clinical exome sequencing in 17 affected individuals from 13 independent families who share a wide range of presentations consistent with a mitochondrial disorder, and optic atrophy was observed in the vast majority of these patients. We performed genetic complementation experiments by transfecting the wildtype \(FDXR\) gene in patient-derived fibroblast cells to support the pathogenesis. Mechanistically, mutations of FDXR may disrupt the function domains of the enzyme based on 3-D crystal model. Our data also suggest some mutations may affect the protein conformation and/or the protein stability. Abnormal
ROS production has been identified in this study and in many mitochondrial diseases, which could be secondary to mitochondrial respiratory chain defects and may play very important roles in the pathogenesis of FDXR mutations. A mouse model carrying a mutation allelic to a common mutation found in our patients showed progressive impairments in enzyme activity and mitochondrial function and recapitulated the major human phenotypes. Mouse model provided a great opportunity to study the pathogenesis of Fdxr mutations. In this study, we were also able to examine the pathogenesis of vision loss. The vision loss seen in Fdxr<sup>R389Q/R389Q</sup> mice is progress- and aging-dependent, suggesting degenerative process. This is further supported by aging-dependent RGC loss and dysmorphic mitochondria in optic nerve. The clinical presentations, mouse models, and experimental findings suggest a biallelic mode of inheritance of FDXR mutations causing mitochondrialopathy with optic atrophy.

Mitochondrial diseases often present with myopathy, optic atrophy, and pigmentary retinopathy in addition to common central nervous system symptoms such as encephalopathy, global developmental delays, and seizure (25). Eye abnormalities, ataxia, and hypotonia exhibited by our patients (Table 1) are consistent with Fe–S disease clinical features (1). Our initial phenotypic analysis of Fdxr<sup>R389Q/R389Q</sup> mice revealed that they share many congruent neurological features with human patients, but there were some discrepancies, including differences in complex activities that could be explained by species-related differences. Alternatively, the most common mutation in humans is p.R392W, rather than p.R392Q. Since tryptophan (W) is a much more radical change from arginine than glutamine (Q), this could explain differences between the mouse and human phenotypes. To further refine the mouse model and provide more complete recapitulation of clinical features, future efforts will be directed towards the generation and characterization of mouse models carrying patient specific mutations. CRISPR/Cas9 genome editing technology to generate knock-in and knockout mouse lines with the same amino acid changes as the hot spot mutation (p.R392W) in humans would provide additional value since the phenotypes of the “stiff” mouse are slightly different from human subjects.

This study is the very first to investigate the clear fact that biallelic FDXR mutations cause human disease, and provides us a unique opportunity to understand the biology of the FDXR pathway. This is a unique opportunity to characterize the molecular mechanisms behind this new disorder. Since FDXR occupies a crucial upstream position at the top of the iron-sulfur cluster biogenesis pathway, with the potential to influence numerous downstream proteins in an epistatic manner, this study provides us the opportunity to understand the major biochemical consequences of mutations in this pathway. Some FDXR variants, such as p.R392W, appear quite frequently in our patients, suggesting that these particular mutations could be the tip of the iceberg for a large, previously unidentified population of patients afflicted by FDXR-mediated diseases. Furthermore, given the fact that FDXR functions as both a...
mitochondrial protein and a protein involved in Fe-S cluster biogenesis, this study permitted us to compare and contrast the distinct yet overlapping phenotypes of these two major classes of disorders (mitochondrial disorders versus Fe-S cluster biogenesis disorders), which will be critical in the development of sound medical management related to these pathways. The combination of human subjects and mouse model allowed us to clearly elucidate these biochemical pathways and will help in the development of precise, targeted therapies for FDXR-related disorders, as well as Fe-S cluster disorders in general.

In conclusion, we identified a novel disease-causing gene FDXR associated with mitochondrial diseases. The biallelic FDXR mutations cause optic atrophy and neuropathy. Because of its upstream role in the Fe–S cluster biogenesis pathway, FDXR mutations can also affect many pathways. In addition, FDXR is required for the cholesterol side-chain cleavage to pregnenolone, a metabolic intermediate of many steroid hormones, raising the possibility that cholesterol pathway could be a potential therapeutic target for the treatment of this condition. Further characterization of the existing models will provide insights for the function of FDXR and the molecular pathology associated with disruption of this protein function. This knowledge will contribute to the existing framework for understanding mitochondrialopathy and improve clinical intervention of these diseases.

Materials and Methods

Clinical information

Patients with optic atrophy, encephalopathy and/or peripheral neuropathy were recruited for clinical whole exome sequencing. They were evaluated by a clinical geneticist, and the clinical features are summarized in Table 1 and Supplementary Material, Note S1. Patients have diverse ethnic backgrounds (Hispanic, Asian, Iranian, and Caucasian ethnic background). For patients who underwent additional procedures, informed consent was obtained in accordance with the Human Subjects Committee, Cincinnati Children’s Hospital Medical Center (CHCM).

Whole-exome sequencing

For patients 1, 2, and 5, whole exome sequencing was performed on the parent–proband trio at Ambry Genetics (Aliso Viejo, CA). Exome library preparation, sequencing, bioinformatics, and data analyses were performed as previously described (26). In brief, exome enrichment was performed using the IDT xGen Exome Research Panel V1.0 (Integrated DNA Technologies; Coralville, IA) and sequenced using paired-end, 100- or 150-cycle chemistry on the HiSeq or NextSeq system (Illumina; San Diego, CA). Data were annotated with the Ambry Variant Analyzer tool, and stepwise filtering included the removal of common alterations, intergenic and 3'/5' untranslated region variants, intronic variants outside ±2, and synonymous variants unlikely to have splicing effects. Variants were then filtered based on family history and possible inheritance models.

For patients 4 and 6–9, whole-exome sequencing was performed at Baylor College of Medicine. The sequencing was performed as described by Yang and colleagues (27). A minimum of 20 x coverage was required for variant calling. About 95% of single nucleotide variants and 88.2–95.0% of indels could be identified.

For patients 10–14, sequencing was performed on exon targets isolated by capture with the Clinical Research Exome kit (Agilent Technologies; Santa Clara, CA) at GeneDx. Genomic DNA was extracted from whole blood samples obtained from affected children and their parents. The supporting sequencing technology and variant interpretation protocol employed are described elsewhere (28).

Generation of stable FDXR overexpressing cell line

Fibroblasts derived from the proband in Family 1 and a control were grown in Dulbecco’s modified Eagle’s medium (DMEM) (Gibco, Thermo Fisher Scientific; Grand Island, NY) supplemented with 10% fetal bovine serum at 37 °C with 5% CO2. WT human FDXR cDNA was obtained from Addgene (pTRE-FDXR WT, transcript variant 3, NM_001258012.3). The full-length cDNA was cloned into the PLVX-ires-tomato lentiviral expression vector (Clontech, Takara Bio Inc.; Shiga, Japan) with an In-Fusion Cloning kit (Clontech) as described previously (25). Briefly, the FDXR PCR fragment and 15-bp vector sequence were fused with linearized vector by In-Fusion Enzyme. The construct was packaged into viral particles in HEK293T cells and lentiviruses were concentrated by ultracentrifugation (CCHMC Vector Core). Transduction of the patient’s fibroblast cell line was performed according to established methods (25). Fibroblasts were seeded in 6-well plates; 24 h later, concentrated lentivirus (titered at 5.44 x 10^6) was added to the cells with 4 μg/ml polybrene. Transduced cells were sorted by flow cytometry with a fluorescein isocyanate (FITC) filter, and tdTomato-positive cells were selected for FDXR overexpression. Phase contrast and immunofluorescence was evaluated by conventional fluorescence microscopy (Zeiss Axiophot; Oberkochen, Germany).

Mouse strain

All procedures involving mice were approved by The Jackson Laboratory's Institutional Animal Care and Use Committee and performed in accordance with the National Institutes of Health guidelines for the care and use of animals in research. The strain used for this study was B6; 129S-Fdxr<sup>m1J</sup>Otop3<sup>m1J/GrsrJ</sup> (The Jackson Laboratory, Stock #026096). This strain was previously shown to exhibit an abnormal, recessive gait phenotype, and harbors a missense (autosomal recessive) mutation in the Fdxr gene resulting in a p.R389Q change in the Fdxr protein (18). Protein alignment also shows that this amino acid change is located at the same location as mutation R392W in the human ortholog. Although these mice also carry a linked homozygous variant of uncertain clinical significance in the 5’ untranslated region of Otop3, it is likely not contributing to the phenotype because mice with a targeted deletion of the Otop3 gene show increased startle reflex and oligodactyly (MGI: 5450617).

Enzymatic assay of ferredoxin NADP reductase

Ferredoxin NADP reductase activity was measured as described previously (29). Briefly, a reaction buffer containing 50 mM Tris-HCl (pH 7.8 at 25 °C), 200 μM NADPH, 75 μM cytochrome C-oxidized, and 10 μM ferredoxin was prepared and incubated at room temperature (RT) for 30 min. Cells were grown to stationary phase in DMEM supplemented with 10% fetal bovine serum, harvested, and lysed in standard RIPA lysis buffer. Similarly, mouse tissues (muscle, brain, liver, heart, and kidney)
from 4-week-old mice were minced into 1–2-mm³ pieces in ice-cold isolation buffer [10mM EDTA/0.05% trypsin in phosphate buffered saline (PBS)], incubated for 30 min on ice, and centrifuged at 300 × g for 5 min. The harvested pellets were homogenized in ice-cold PBS containing 10mM EDTA and protease inhibitor using a GentleMACS dissociator. The homogenized samples were filtered with 70 μm filters and centrifuged (4 °C, 5 min at 800 × g). Finally, human cell or mouse tissue lysates were added to the reaction buffer, mixed, and spectrophotometric measurements were taken at A550nm. Relative enzyme activity was calculated after normalization relative to control samples.

**Measurements of mitochondrial respiration**

OCRs in fibroblast cell lines were measured with a XF-96 extracellular flux analyzer (Seahorse Biosciences; North Billerica, MA) (30,31). The XF96 analyzer creates a transient, 7 μl-chamber in specialized microplates that allows determination of oxygen and proton concentrations in real time. The cultured cells were transferred to an XF96 analyzer assay plate at a density of 1 × 10⁴ cells/well and allowed to grow overnight. The measured parameters included basal respiration, ATP production, and maximal respiratory capacity. To allow comparison across experiments, OCR and acidification rate data are expressed in pmol/min and mH/min, respectively, normalized to cell protein in individual wells determined by Bradford protein assays (Bio-Rad; Hercules, CA). OCRs were determined under normal conditions and after the addition of oligomycin (1.5 μM), carbonyl cyanide p-(trifluoromethoxy) phenylhydrazone (FCCP) (0.5 μM), rotenone (1 μM), and antimycin A (1 μM).

**ATP levels**

ATP was measured using the ATP Bioluminescence Assay Kit CLS II, according to the manufacturer’s instructions (Roche Diagnostics; Basel, Switzerland) (32). Briefly, the assay buffer and substrate were equilibrated to RT, and the buffer was mixed with the substrate (assay reagent) to obtain a homogeneous solution. After a 30-min equilibration of the cell plates to RT, 100 μl of assay reagent was added to each well with 2 × 10⁴ fibroblasts, and the content was mixed for 2 min on an orbital shaker to induce cell lysis. After a 10-min incubation at RT, the luminescence was measured on a Synergy H1 microplate reader (BioTek Instruments; Winooski, VT) (33,34). ATP production in mouse tissues was measured similarly.

**Mitochondrial respiratory chain complex I–IV activity assay**

The enzymatic activities of complexes I, II, III, and IV were assayed as described previously (35,36). Complex I (NADH dehydrogenase) activity was determined by the rotenone-sensitive NADH oxidation at 340 nm, using the coenzyme Q analog 2, 3-dimethyl-5-methyl 6-n-decyl-1, 4-benzomethyluinone as an electron acceptor. The activity of complex II (succinate dehydrogenase) was analyzed by tracking the secondary reduction of 3-dimethyl-5-methyl 6-n-decyl-1, 4-benzomethyluinone as an electron acceptor. The activity of complex III (cytochrome b c1 complex) activity was determined by measuring the reduction of cytochrome C at 550 nm with reduced decylubiquinone. Complex IV (cytochrome c oxidase) activity was measured by monitoring the oxidation of reduced cytochrome C as a decrease in absorbance at 550 nm. Citrate synthase activity was analyzed by measuring the reduction of 5, 5’-dithiobis-2-nitrobenzoic acid at 412 nm in the presence of acetyl-CoA and oxaloacetate. Complex activities were normalized relative to citrate synthase activity.

**Visual acuity**

Analysis of visual acuity was performed by the Mouse Neurobehavioral Phenotyping Core at The Jackson Laboratory using the OptoMotry system (CerebralMechanics Inc.). Untrained, unrestrained mice were placed on an elevated platform in the center of an arena surrounded by a cylinder with a 3D sine wave grid projection. The directionality of the subject’s head in response to variable frequency rotation of the cylinder was measured and an optokinetic (OKT) response score was calculated by the accompanying instrumentation software. A total of 41 mice (15 Fdxr<sup>R389Q/R389Q</sup>, 15 Fdxr<sup>R389Q/R389Q</sup>, 11 Fdxr<sup>−/−</sup>) were tested at 15–17 weeks of age. Technicians were blind to genotypes during testing. Graphing and statistical analyses were performed using Prism 7 (GraphPad Software; La Jolla, CA). Ordinary one-way ANOVA tests with a Dunnett’s post hoc test (to adjust for multiple comparisons) were used to compare each genotype (homozygous and heterozygous) to WT. No significant differences between wild type and heterozygous mice were found; therefore, unpaired, nonparametric t-tests (Mann Whitney U test) were ultimately used to demonstrate significance between WT and Fdxr<sup>R389Q/R389Q</sup>. P values of less than 0.05 were considered significant [<0.05 *], <0.01 ** and <0.001 ***].

**Cell images**

Fibroblasts were seeded in 6-well plates; 24 h later, concentrated lentivirus (titered at 2.44 × 10⁸) was added together with 4 μg/ml polybrene. Images were captured by conventional fluorescence microscopy after transduction of PLVX-FDXR-tdTomato lentivirus for 48 h. Transduced cells were sorted by flow cytometry with a FITC filter, and tdTomato-positive cells were selected for FDXR overexpression. Patient fibroblasts were sorted by FACS after 24 h in culture.

**Confocal microscopy of immunofluorescence**

Fibroblasts were cultured in DMEM containing 200mM MitoTracker Red (Molecular Probes, Thermofisher Scientific; Eugene, OR) for 30 min at 37 °C. After washing with PBS, fibroblasts were fixed in 4% paraformaldehyde in PBS for 20 min. They were incubated in 0.3% Triton X-100 for 30 min, and treated with 50 mM NH₄Cl in PBS for 15 min, followed by a PBS wash. The sections were blocked for 1 h at room temperature (RT) in blocking buffer (10% goat serum and 0.4% Triton X-100 in PBS) and then incubated with rabbit anti-Fdxr antibody (1:100) overnight at 4 °C. After three 10-min washes in PBS, goat anti-rabbit secondary antibody conjugated with Alexa Fluor® 488 (1:1500) was applied to the sections for 2 h at RT. Sections were mounted with VECTASHIELD medium containing DAPI (Vector H1200, Vector Laboratories, Burlingame, CA). Cell images were obtained via confocal microscopy using a Nikon A1R LUN-V Inverted confocal microscope (Nikon Instruments Inc.; Melville, NY) and analyzed with Imaris software (Bitplane; South Windsor, CT).
Detection of fluorescent ROS and superoxide production

We measured ROS generation levels in fibroblasts with the Cellular ROS/Superoxide Detection Assay Kit (ab139476, Abcam; Cambridge, UK). Seeded cells were placed directly onto glass slides or polystyrene tissue culture plates to ensure 50–70% confluency. After a media change, cells were treated by 30 min with the ROS inhibitor N-acetyl-L-cysteine. Pyocyanin was used to induce ROS production; controls included vehicle alone (no inhibitor and no inducer of ROS) and vehicle with inhibitor. ROS production was observed with controlled oxidative stress detection reagent (green) and superoxide detection reagent (orange) for 30 min to 1 h. Afterwards, cells were washed twice with 1× wash buffer. Cell images were captured with a fluorescence microscope using standard excitation/emission filter sets (green 490/525 nm and orange 550/620 nm).

H&E staining and electron microscopy

Eyes were cut in 4% paraformaldehyde for 24 h at RT and then embedded in paraffin. Sagittal sections of the eye (6–8 μm) were cut through the optic nerve head and then stained with H&E. Images were obtained under light microscopy (BX63; Olympus Corporation; Center Valley, PA). Central (250 μm from the edge of the optic disc) and peripheral (100 μm from the ora serrata) segments of the retina were photographed. Nuclei in the retinal ganglion cell layer (excluding the small and flat nuclei of the endothelial cells) were counted on the photographed images. Length of the retina was measured using an ImageJ program.

For electron microscopy, mice were anesthetized and perfused with 0.9% NaCl and EM fixative (4% paraformaldehyde, 2.5% glutaraldehyde, PBS, pH 7.4–7.6). Tissues were postfixed in 0.1 M sodium cacodylate buffer (EMS; Hatfield, PA), and postfixed in 1% osmium tetroxide (EMS) for 1 h at 4°C. Samples were washed in 0.1 M sodium cacodylate buffer and dehydrated through a graded ethanol series. Samples were embedded in LX-112 (Ladd Research Industries; Williston, VT). Tissue blocks were sectioned (thickness, 0.5–1 μm) and stained with toluidine blue for light microscopy. Blocks were trimmed and cut into 90-nm-thick sections with an ultramicrotome (Leica EM UC7; Buffalo Grove, IL). The ultra-thin sections were counterstained with uranyl acetate 2% (EMS) and lead citrate. All images were taken with an 80-kV transmission electron microscope (Hitachi, H-7650, V01.07; Tokyo, Japan). Measurement of the thickness of myelin membrane in RGC axons was performed using ImageJ software (National Institutes of Health).

Western blotting

Cell lysates from mouse tissues were harvested as described above. Approximately, 50 μg of protein was mixed with 2× reducing sample buffer (Sigma-Aldrich; St. Louis, MO). The samples were heated to 95°C for 5 min before being separated on a 10% SDS-PAGE gel (21). The gel was electrophoretically transferred onto PVDF membranes (Invitrogen, Thermo Fisher Scientific; Carlsbad, CA) at 80 V for 2 h. The membrane was blocked for 1 h in 10% milk prior to overnight incubation in primary antibody FDXR (Abcam) at 1:100 dilution at 4°C. This overnight incubation was then followed by a 1-h incubation at 37°C with horseradish peroxidase-conjugated secondary antibodies (Sigma-Aldrich). ECL plus detection reagent (GE Healthcare; Chicago, IL) was used for visualization of bands, and quantification analysis of band intensity was performed in ImageJ software (version 1.45s). GAPDH at a 1:1000 dilution (Ambion, Thermo Fisher Scientific; Foster City, CA) was used as a loading control.

3D protein structure

The structure of FDXR was built from the homologous structure of FDXR for Bos taurus complexed with FAD (PDB: 1CJC) (22) and complexed with FX1 and FDXR (PDB: 1E6E) (37). The homology model of FDXR was built with the ROSETTA homology modeling module (38) with sequence alignments generated in TCOFFEE (39,40). The energy calculations for internal structure destabilization and protein-protein binding were done in the FoldX program (41). Potential phosphorylation and modification sites in FDXR were identified in Eukaryotic Linear Motif (42) to identify potential motifs, and checked in the experimental database of Phosphosite (43). The images were generated in the PyMOL Molecular Graphics System, Version 1.8 (Schrödinger, LLC; New York, NY).

Supplementary Material

Supplementary Material is available at HMG online.

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