Cisd2 deficiency drives premature aging and causes mitochondria-mediated defects in mice

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CISD2, the causative gene for Wolfram syndrome 2 (WFS2), is a previously uncharacterized novel gene. Significantly, the CISD2 gene is located on human chromosome 4q, where a genetic component for longevity maps. Here we show for the first time that CISD2 is involved in mammalian life-span control. Cisd2 deficiency in mice causes mitochondrial breakdown and dysfunction accompanied by autophagic cell death, and these events precede the two earliest manifestations of nerve and muscle degeneration; together, they lead to a panel of phenotypic features suggestive of premature aging. Our study also reveals that Cisd2 is primarily localized in the mitochondria and that mitochondrial degeneration appears to have a direct phenotypic consequence that triggers the accelerated aging process in Cisd2 knockout mice; furthermore, mitochondrial degeneration exacerbates with age, and the autophagy increases in parallel to the development of the premature aging phenotype. Additionally, our Cisd2 knockout mouse work provides strong evidence supporting an earlier clinical hypothesis that WFS is in part a mitochondria-mediated disorder; specifically, we propose that mutation of CISD2 causes the mitochondria-mediated disorder WFS2 in humans. Thus, this mutant mouse provides an animal model for mechanistic investigation of Cisd2 protein function and help with a pathophysiological understanding of WFS2.

[Keywords: Cisd2; Wolfram syndrome 2; autophagy; knockout mice; mitochondria; premature aging]

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CISD2 is the second member of the gene family containing the CDGSH iron sulfur domain. There are currently three members in this gene family: CISD1 (synonyms ZCD1, mitoNEET), CISD2 (synonyms ZCD2, Noxp70, and Miner1) and CISD3 (synonym Miner2). CISD1 is an outer mitochondrial membrane protein that was originally identified as a target protein of the insulin sensitizer drug pioglitazone used to treat type 2 diabetes (Colca et al. 2004). CISD1 protein contains a transmembrane domain, a CDGSH domain, and a conserved amino acid sequence for iron binding; biochemical experiments suggest that CISD1 is involved in the control of respiratory rates and regulates oxidative capacity (Wiley et al. 2007). However, CISD2 and CISD3 are novel genes with previously uncharacterized functions. The only molecular documentation for CISD2 is that CISD2 was one of the markers for early neuronal differentiation in a cell culture study (Boucquey et al. 2006).

Recently, the CISD2 gene has been identified as the second causative gene (Amr et al. 2007) associated with Wolfram syndrome (WFS; MIM 222300), which is an autosomal recessive neurodegenerative disorder. WFS is highly variable in its clinical manifestations, which include diabetes insipidus, diabetes mellitus, optic atrophy, and deafness; thus, it is also known as the “DIDMOAD syndrome” (Barrett and Bundey 1997). Positional cloning and mutation studies have revealed that WFS is a genetically heterogeneous disease with a complex molecular basis involving more than one causative gene in humans (Domenech et al. 2006). A portion of WFS patients belonging to the WFS1 group (MIM 222300) carried loss-of-function mutations in the WFS1 (wolframin) gene, which encodes a transmembrane protein primarily localized in the endoplasmic reticulum (ER) [Inoue et al. 1998; Strom et al.1998; Takeda et al. 2001]. In addition to this, a homozygous mutation of the CISD2 gene has been identified in three consanguineous families with WFS.

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Somatotype are clearly evident; it appears that there is (Supplemental Fig. 4). Growth retardation and a smaller gene was disrupted and that there was undetectable mRNA physiology, we generated role of Cisd2 involvement in development and patho-
naturally aged mice (Supplemental Fig. 3). To study the Cisd2 levels of quantitative real-time RT–PCR revealed that expression Cisd2 is a widely expressed gene in mice. Interestingly, chromosome 3G3. Northern blot analysis showed that suplemental Figs. 1, 2); the mouse syntenic region is on Cisd2 is an evolutionarily conserved gene localized on chromosome 4q where a genetic component for human longevity has been mapped. Previously, Puca et al. [2001] studied 137 sets of extremely old siblings (308 individuals in all) and conducted a genome-wide scan search for predisposing loci that might confer longevity; this linkage study revealed a single region on chromosome 4q and suggests that there may be at least one master gene contributing to life-span control; however, the responsible gene has not been identified.

In this study, we apply a mouse genetics approach and demonstrate that Cisd2 is involved in mammalian life-span control and plays an essential role in mitochondrial integrity. Cisd2 deficiency causes mitochondria-mediated phenotypic defects in mice. Furthermore, cell culture and biochemical investigations revealed that Cisd2 is a mitochondrial protein. Additionally, Cisd2 knockout mice exhibit many clinical manifestations of WFS patients including early-onset degeneration of central [e.g., optic] and peripheral [e.g., sciatic] nerves and premature death, as well as impaired glucose tolerance. This study therefore provides an animal model for mechanistic understanding of WFS, specifically WFS2, pathogenesis.

Results

Shortened life span in Cisd2–/– mice

CISD2 is an evolutionarily conserved gene localized on human chromosome 4q24 [Supplemental Table 1; Supplemental Figs. 1, 2]; the mouse syntenic region is on chromosome 3G3. Northern blot analysis showed that Cisd2 is a widely expressed gene in mice. Interestingly, quantitative real-time RT–PCR revealed that expression levels of Cisd2 decrease in an age-dependent manner in naturally aged mice [Supplemental Fig. 3]. To study the role of Cisd2 involvement in development and pathophysiology, we generated Cisd2 knockout mice. Southern and Northern blot analyses demonstrated that the Cisd2 gene was disrupted and that there was undetectable mRNA expression in the homozygous knockout (Cisd2–/–) mice [Supplemental Fig. 4]. Growth retardation and a smaller somatotype are clearly evident; it appears that there is almost no growth after 5 wk old in the Cisd2–/– mice [Fig. 1A]. Early senescence is accompanied by a shortened life span when survival of the various genotypes is examined and there appears to be signs of haploinsufficiency for Cisd2 in view of the slightly lower survival rate for the heterozygous (Cisd2+/–) mice [Fig. 1B].

Premature aging phenotype

Starting at 8 wk old, Cisd2–/– mice begin to acquire a set of aged appearance phenomena remarkably similar to those of premature aging syndrome (Hasty et al. 2003; Kipling et al. 2004). These include prominent eyes and protruding ears (Fig. 1C). Ocular abnormalities were observed as the Cisd2–/– mice developed opaque eyes and blindness, which was accompanied by cornea damage at 20 wk old [Fig. 1D]. Histopathological examination revealed that the opacity of the cornea was due to debris deposition in the scar tissue outside the cornea [Fig. 1E]. In addition, corneal neovascularization was observed in the Cisd2–/– mice; this can impair vision and is usually associated with pathogenesis due to eye trauma or the presence of a degenerative disorder [Supplemental Fig. 5]. There was also early depigmentation in the fur at ~48 wk old [Fig. 1F, Supplemental Fig. 6]; furthermore, hair follicle atrophy and a decreased hair density could be detected in Cisd2–/– mice [Fig. 1G,H]. A decrease in the hair regrowth rate was also observed in the Cisd2–/– mice [Supplemental Fig. 7A,B]. Additionally, the skin of 48-wk-old Cisd2–/– mice exhibits a phenotype with a noticeably thickened dermis, an expanded surface, and a significant decrease in subcutaneous adipose tissue and muscle [Fig. 1I–K].

Microcomputer tomography [micro-CT] imaging showed that the trabeculae of the femur are noticeably thinner in Cisd2–/– mice [Fig. 2A]. Dual energy X-ray absorptiometer [DEXA] detected a decrease in femur density after 8 wk old; interestingly, the decrease of femur density also started to emerge in heterozygous Cisd2+/– mice, but at 24 wk old, while a progressively more severe phenotype was observed at the same age with Cisd2–/– mice [Fig. 2B]. This shows, in addition to what was observed in terms of life-span evaluation, that there is also an apparent Cisd2 haploinsufficiency with respect to femur density. The results from the gross anatomy viewpoint, from the X-ray radiography, and using micro-CT reveal a significant lorderkymphosis phenotype after 12 wk old [Fig. 2C,D, Supplemental Fig. 7C,D]; consequently, this seems to lead to a decrease in mean thoracic volume [Fig. 2E] and thence pulmonary function abnormalities. Indeed, we observed decreases in various respiratory parameters as measured by plethysmography after 20 wk old in the Cisd2–/– mice [Supplemental Fig. 8]. Muscle degeneration was detectable at 3 wk old in the Cisd2–/– mice. There was a progressive degeneration of muscle fibers and the magnitude of the degeneration exacerbated with age [Fig. 2F–J]; muscle degeneration was further confirmed by transmission electron microscopy [TEM] [Supplemental Fig. 9]. In addition, angular fibers, which are an indicator of muscle atrophy caused by neuron degeneration, could be observed in the Cisd2–/– mice [Fig. 2H].

One possible mechanism for the accelerated aging phenotypes is a defect in cellular proliferation in the Cisd2–/– mice. To test this possibility, we created several primary mouse embryonic fibroblast [MEF] cell lines from individual embryos with different genotypes. Our results revealed no significant difference in the doubling time and MEF cell growth [Supplemental Fig. 10], suggesting that accelerated aging in the Cisd2–/– mice is not due to an intrinsic defect in cellular proliferation.

A summary of the aging-related phenotypes in Cisd2–/– mice is provided in Supplemental Table 2. These mutant
mice exhibit a premature aging phenotype with 100% penetrance for both sexes using either a C57BL/6 (B6) or a 129Sv/B6 mixed background.

Mitochondrial degeneration and autophagy

The observation of premature aging phenotypes involving muscle degeneration prompted a detailed examination of the tissue ultrastructure of the homozygous knockout mice. A TEM study revealed that mitochondrial degeneration occurs in the axons of sciatic nerves, brain cells (Fig. 3A–C), cardiac muscle cells, and skeletal muscle cells (Fig. 3D–F) in the Cisd2<sup>−/−</sup> mice. Notably, the mitochondrial outer membrane (OM) appeared to have broken down prior to the destruction of the inner cristae (Fig. 3B,E). In wild-type mice, the myelinated axons are enveloped with a myelin sheath formed by the fusion of many layers of plasma membrane from Schwann cells (Fig. 3G). However, considerable disintegration of the myelin sheath and degeneration of axon was detected in the Cisd2<sup>−/−</sup> sciatic nerves (Fig. 3H,I). Importantly, these mitochondrial abnormalities, involving destruction of mitochondria, myelin sheath disintegration, and axonal lesions, are already present to a certain extent in 2-wk-old Cisd2<sup>−/−</sup> mice (Fig. 3J–L; Supplemental Fig. 11), a stage prior to the first premature aging phenotype of muscle and nerve degeneration in these mice. Interestingly, the damaged mitochondria appear to induce autophagy to eliminate the dysfunctional organelles (Kim et al. 2007) because we identified morphologically distinct autophagic vacuoles (Eskelinen 2008) in muscle, sciatic nerve, optic nerve, and brain tissue (Fig. 3J–L; Supplemental Fig. 12). The general term autophagic vacuole refers to an autophagosome, amphisome, or autolysosome. Morphologically, autophagic vacuoles can be classified into two categories: (1) early or initial autophagic vacuoles (AVis)—i.e., autophagosomes, which are double-membraned structures containing undigested cytoplasmic material or organelles; (2) late or degradative autophagic vacuoles (AVds), including amphisomes and autolysosomes, which contain partially degraded cytoplasmic material (Eskelinen 2008; Fader and Colombo 2009). Remarkably, mitochondrial degeneration exacerbates with age, and the magnitude of the autophagy increases in parallel to the development of premature aging phenotype (Fig. 3M,N). We measured the thickness of myelin...
sheaths and counted the numbers of myelinated axons in sciatic nerve; our results revealed no significant differences between different genotypes [Supplemental Fig. 13], indicating that these two factors are not involved in the nerve degeneration of Cisd2⁻/⁻ mice. We also examined the autophagosomal marker LC3-II [Kabeya et al. 2000] in skeletal and cardiac muscles, which are the most sensitive tissues to in vivo autophagic degradation [Mizushima et al. 2004]; indeed, the ratio of LC3-II/LC3-I was significantly higher in Cisd2⁻/⁻ mice than in their wild-type littermates. This biochemical evidence confirms the TEM results and provides a quantitative basis for the autophagy induction (Fig. 3O,P).

Autophagy can lead to cell death by directly activating autophagic (type II programmed) cell death that produces self-degradation of the dying cells [Shimizu et al. 2004; Mizushima et al. 2008]; alternatively, autophagy might cause cell death through activation of apoptosis [Scott et al. 2007]. To determine whether there is an increased level of apoptosis in the Cisd2⁻/⁻ mice, we performed TUNEL assays on various mouse tissues to detect apoptotic cells in situ and found no evidence of increased apoptosis in Cisd2⁻/⁻ mice [Supplemental Fig. 14]. In addition, it has been reported that starvation can induce muscle autophagy [Mizushima et al. 2004]. To test this possibility, we measured the metabolic indices including intake of food and water and generation of urine and stool. Our results revealed no significant difference in these metabolic indices between Cisd2⁻/⁻ and wild-type mice at 6 wk old [Supplemental Fig. 15A]; this is 4 wk after the detection of autophagic activation at 2 wk old. This excludes starvation/malnutrition as the cause of autophagic induction in Cisd2⁻/⁻ mice. A decrease in the metabolic index becomes evident after 12 wk old [Supplemental Fig. 15B], and this is likely to be a consequence of the aging phenotype.

**Cisd2 is probably a mitochondrial OM protein**

The annotated characteristics of Cisd2 protein are very similar to Cisd1, which is an outer mitochondrial membrane protein (Supplemental Fig. 16A; Wiley et al. 2007). To address the subcellular localization, we expressed the EGFP-tagged Cisd2 protein in NIH3T3 cells. Our result indicated that Cisd2 was colocalized with the mitochondrial marker [Supplemental Fig. 16B]. However, deletion of the N-terminal 58 amino acids completely abolished the mitochondrial localization; furthermore, when the N-terminal 58 amino acids were fused to EGFP, this construct was able to redirect EGFP from a nuclear and cytoplasmic localization to the mitochondria (Fig. 4A), suggesting that Cisd2 is a nucleus-encoded mitochondrial protein and its N-terminal 58 amino acids are both necessary and sufficient to direct mitochondrial localization. To confirm the subcellular localization of the Cisd2 protein, the cytosolic and mitochondrial fractions were prepared from skeletal muscle of wild-type mice. Antibodies against Cisd1 and Cisd2 were generated. Western blot analysis revealed that Cisd2 protein, like the mitochondrial proteins Cisd1 and Hsp60 [Samali et al. 1999;
Figure 3. Mitochondrial degeneration and autophagy induction in the muscles and neurons of the Cisd2−/− mice. (A) Wild-type mitochondria in the brain (hippocampus). (B) A Cisd2−/− mitochondrion in the brain (hippocampus). Note that the outer mitochondrial membrane has broken down (arrowhead), while the inner cristae appear to be intact. (C) Cisd2−/− mitochondria in sciatic nerve. One mitochondrion (arrowhead) has a destroyed OM, but with cristae still visible; the other mitochondrion (arrow) has destroyed OMs and IMs. (D) Wild-type mitochondria in cardiac muscle. (E) Cisd2−/− mitochondria in cardiac muscle. This micrograph shows one mitochondrion (arrowhead) with a destroyed OM and two degenerated mitochondria consisting of debris (arrows). (F) A cluster of autophagic vacuoles and abnormal mitochondria was observed between the myofibrils of Cisd2−/− skeletal muscle (white arrows). (G) A wild-type myelinated axon of the sciatic nerve. (N) Nucleus of Schwann cell; (MS) myelin sheath. (H) A myelinated axon of sciatic nerve from a Cisd2−/− mouse. An ovoid with a disintegrating myelin sheath and a degenerating axonal component are shown. (I) Debris from an axon undergoing degeneration in the Cisd2−/− sciatic nerve. (J–L) Early or AVis enclosing mitochondria (arrows) and late or AVds were detected in the axonal component and cytoplasm of a Schwann cell from a 2-wk-old Cisd2−/− sciatic nerve. (M,N) Percentage of myelinated axons present in the sciatic nerves showing disintegration of their myelin sheaths and autophagic vacuoles, including AVi and AVd, in their axonal component. There were three mice for each group. (O) Western blotting to detected the presence of the proteins LC3-I and LC3-II. (P) Ratios of the LC3-II to LC3-I. There were three mice for each group. (*) P < 0.05; (**) P < 0.005. Mouse age in A–I is 4 wk old.
Colca et al. 2004; Wiley et al. 2007), is primarily localized in the mitochondrial fraction (Fig. 4B). To further define the submitochondrial localization of Cisd2, we separated mouse liver mitochondria into the following fractions: OM, mitoplasts (MP, inner membrane [IM] and matrix), and intermembrane space (IMS, soluble material between the IM and OM). Immunoblotting each fraction with antibodies against Cisd2 and known markers revealed that Cisd2 was highly enriched in the OM fraction, as was the OM marker VDAC-1; this result strongly suggests that Cisd2 is a mitochondrial OM protein (Fig. 4C).

Previously, Amr et al. (2007) reported that the Flag-tagged CISD2 protein colocalized with the ER marker calnexin in the transfected mouse P19 and human HEK293 cells. We sought to determine if there is a small portion of the Cisd2 protein sorted into the ER/sarcoplasmic reticulum (SR) using subcellular fractions prepared from skeletal muscles of 12-wk-old mice. Polyclonal antibody (Ab) against Cisd2 protein (15 kDa) was generated; this antibody cross-reacts with Cisd1 protein (12 kDa). Antibodies against mitochondrial proteins Cisd1 and Hsp60 were used as controls. Ten micrograms of each submitochondrial fraction prepared from the livers of 4-wk-old mice were analyzed by Western blot using antibodies against Cisd2 and known mitochondrial marker proteins. OM marker: [VDAC-1] voltage-dependent anion channel-1, IM marker: [ATPSB] complex V β subunit, matrix marker: [PDH] pyruvate dehydrogenase. (MP) Microplast (IM and matrix); (IMS) intermembrane space. (D) Impaired mitochondrial respiration in the skeletal muscle of 4-wk-old Cisd2−/− mice. Representative oxygraphs of the mitochondria after adding first glutamate-malate and then ADP into the closed chamber of the oxygen meter. (E) Respiratory activity was expressed as oxygen consumption rate (nanomoles of O₂ per minute per milligram of mitochondria) in the resting state, for glutamate-malate supported respiration, and for ADP activated respiration. A significant decrease in oxygen consumption was detected in the Cisd2−/− mitochondrial samples (n = 4) compared with wild-type samples (n = 3). (F) The RCR (O₂ consumption rate after ADP addition/O₂ consumption rate after glutamate-malate addition) was significantly lower in the Cisd2−/− mitochondria. (G) Comparison of electron transport activities of the respiratory enzyme complexes of mitochondria prepared from the skeletal muscles of 4-wk-old Cisd2−/− (n = 4) and wild-type mice (n = 4). [NCCR activity] Measurement of NCCR activity, which represents complexes I–III; [SCCR activity] measurement of SCCR activity, which represents complexes II and III; [CCO activity] cCCO activity, which represents complex IV. (*) P < 0.05; (**) P < 0.005.
whether the mitochondrial degeneration detected in this study has a direct functional consequence leading to a respiratory dysfunction, we assessed mitochondrial aerobic respiration using isolated mitochondria prepared from skeletal muscle. This was done by measuring the oxygen consumption after stimulating the mitochondria with glutamate-malate and ADP to activate the respiratory chain reactions. Our results revealed a significant decrease in the oxygen consumption and the respiratory control ratio (RCR) in the Cisd2−/− mitochondria [Fig. 4D–F}. To further expand this investigation, we explored the iron-sulfur proteins, which are essential electron carriers in the mitochondrial respiratory chain; there are up to 12 different iron-sulfur clusters that shuttle electrons through complex I–III [Rouault and Tong 2008]. We measured the activities of the various iron-sulfur proteins of complex I–III [NADH cytochrome c reductase, NCCR] and complex II–III [succinate cytochrome c reductase, SCCR]. In addition, we also measured the activity of complex IV [cytochrome c oxidase, CCO], which contains hemes and copper centers for electron transport [Rouault and Tong 2008]. Our results showed that there was an average 30% decrease in the electron transport activities of complex I–III, complex II–III, and complex IV in the Cisd2−/− mitochondria compared with wild-type mitochondria [Fig. 4G]. Together with the oxygen consumption experiment, these results reveal a respiratory dysfunction in the Cisd2−/− mitochondria.

To test whether an increased level of reactive oxygen species (ROS), which is a by-product of mitochondrial oxidative phosphorylation, may contribute to the phenotypes of Cisd2−/− mice, we monitored the intracellular ROS, mainly H2O2, in MEF cells and primary cells obtained from the brains and livers of different genotypes of mice. There was no significant difference in the ROS levels in these primary cells between the different genotypes [Supplemental Fig. 18A]. In addition, the mRNA levels of the enzymes that scavenge ROS were unaffected in brain, heart, liver, and skeletal muscle, suggesting that there was no ROS-induced stress response present in the Cisd2−/− mice [Supplemental Fig. 18B].

**WFS and Cisd2−/− mice**

In order to evaluate the usefulness of Cisd2−/− mice as an animal model for WFS2 and gain insight into the mechanistic basis of WFS2 pathogenesis, we compared the clinical manifestations of this disease and the phenotype of Cisd2−/− mice. WFS is a clinically heterogeneous disease; only juvenile-onset diabetes mellitus and optic atrophy are necessary criteria for WFS diagnosis. Importantly, Cisd2−/− mice exhibit a progressive neurodegenerative phenotype that includes optic nerve defects [Fig. 5A,B, Supplemental Fig. 12]. Regarding glucose homeostasis, we found that Cisd2−/− mice display a milder phenotype, namely, impaired glucose tolerance and decreased insulin secretion, which was revealed by the oral glucose tolerance test [Fig. 5C,D]. In addition, insulin tolerance tests did not show insulin resistance in the Cisd2−/− mice; in fact, these mutant mice were some-
resistance. The importance of mitochondrial dysfunction in β-cell insulin secretion defects has been previously confirmed in other mouse models, which demonstrated that mitochondrial ATP production is a critical part of the β-cell signaling system and allows insulin release [Wallace 2001; Torraco et al. 2009]. However, there was no overt diabetes observed in the Cisd2−/− mice with the C57BL/6 congenic background. This is consistent with a previous observation that C57BL/6 background confers a more diabetes-resistant phenotype [Coleman 1992], a similar finding of a genetic background effect also had been reported for WFS1 [wolframin] knockout mice [Ishihara et al. 2004]. In addition to optic atrophy and glucose intolerance, the phenotypic features of Cisd2−/− mice reflect other aspects of the clinical manifestations of WFS patients including early [juvenile] onset and premature death [Supplemental Table 3]. Thus, this mutant mouse may also provide an animal model for mechanistic investigation of Cisd2 protein function and help with the pathophysiological understanding of WFS2.

Discussion

For more than a decade, physicians and researchers have fiercely debated as to whether WFS is associated with mitochondria and a defect in ATP supply. Most WFS patients die prematurely with severe neurological disabilities involving the central nervous system and peripheral nerves [Barrett and Bundey 1997; Domenech et al. 2006]. In 1993, Bu and Rotter proposed a dual genome defect model and hypothesized that mitochondrial DNA mutation and nuclear genetic defects that interfere with the normal function of mitochondria can independently lead to WFS [Bu and Rotter 1993]. This hypothesis was based on the clinical observations that the affected tissues and organs in WFS patients have a high metabolic demand and most of the clinical manifestations of WFS are consistent with an ATP supply defect, which is often seen in mitochondria-mediated disorders. There were several studies supporting this hypothesis [Rötig et al. 1993; Vora and Lilleyman 1993; Barrientos et al. 1996]. Notably, Bundey et al. [1992] described a WFS patient having morphologically and biochemical abnormal mitochondria in the muscle biopsy; this finding indicated that a mitochondrial defect may be involved in the pathogenesis of WFS. However, other clinical studies revealed no evidence supporting the hypothesis of mitochondrial deficiency [Hofmann et al. 1997; Barrett et al. 2000]. This controversy seems to have been resolved by the identification of different causative genes for WFS, and this hypothesis is supported by the mouse works carried out in this study.

WFS1 is associated with an ER defect

Previous studies in patients had identified WFS1 [wolframin] as the causative gene for WFS1 [Inoue et al. 1998; Strom et al. 1998]. Biochemical and cell culture investigations revealed that wolframin is a transmembrane protein primarily localized in the ER and may be involved in the regulation of ER stress and calcium homeostasis [Takeda et al. 2001; Fonseca et al. 2005; Zatyka et al. 2008]. In animal studies, a pancreatic phenotype related to glucose intolerance and impaired insulin secretion, but not the neurodegenerative phenotype, has been reported in wolframin knockout mice. Wolframin deficiency in mice leads to progressive loss of β cells and impaired glucose homeostasis [Ishihara et al. 2004], which appears to be caused by increased ER stress and apoptosis in the pancreatic β cells [Riggs et al. 2005; Yamada et al. 2006]. Obviously, the pathogenesis of WFS1 patients with wolframin mutations is mechanistically related to an ER rather than a mitochondrial defect. This provides an explanation for the discrepancy as to why there were contradictory observations in some WFS [specifically WFS1] patients who do not have any detectable abnormality in their mitochondria.

WFS2 is a mitochondria-mediated disorder

Recently, Amr et al. [2007] identified CISD2 homozygous mutations in WFS2 patients and suggested that CISD2 is the causative gene responsible for WFS2. Our Cisd2 gene knockout mouse work provides strong evidence supporting the hypothesis that WFS is a mitochondria-mediated disorder, thus, specifically, WFS2, which is caused by a CISD2 mutation, is a mitochondria-mediated disorder. Previous clinical studies in WFS patients suggested that optic atrophy probably represents a degeneration of the optic nerve [Mtanda et al. 1986; Barrett et al. 1997]. Indeed, our mouse work has revealed that progressive degeneration of the optic nerve is one of the earliest phenotypic features detected at 2–3 wk of age, which is before weaning; this phenotype exacerbates with age in the Cisd2−/− offspring [Fig. 5; Supplemental Fig. 12]. Regarding glucose homeostasis, although the phenotype is relatively milder and only glucose intolerance was observed in the diabetes-resistant C57BL/6 background, in the future, it will be of great interest to introduce the Cisd2 mutant allele into C57BLKS/J [Mao et al. 2006], 129/Sv [Terauchi et al. 2003], or other diabetes-prone strains of mice, which may contain genetic modifier(s) that increase susceptibility to diabetes. This will allow the effect of the genetic background on the severity of diabetes to be examined.

Our present study reveals that Cisd2 is primarily localized in the mitochondria. Cisd2 deficiency causes mitochondrial dysfunction accompanied by autophagic cell death, and these events precede neuron and muscle degeneration; together, they lead to a panel of phenotypic features suggestive of premature aging. Since muscles and nerves have the highest energy needs and are therefore the most dependent on mitochondrial function, this explains why neuronal lesions and muscle abnormalities are the two earliest manifestations and why they precede the gross premature aging phenotype. Accordingly, mitochondrial degeneration appears to have a direct phenotypic consequence that triggers the accelerated aging process in Cisd2−/− mice [Fig. 6]. Our results thus provide strong evidence for the causal involvement of mitochondrial
A specific pathogen-free facility and treated according to the was obtained from their agouti progeny. The mice were bred in male mice were bred with B6 females. Germline transmission 3. Targeted ES cells were injected into B6 blastcysts. Chimeric probe, specifically a 1.7-kb BamHI–EcoRI fragment from exon

dysfunction in driving mammalian aging as suggested previously by other studies in mitochondrial DNA mutator mice [Trifunovic et al. 2004; Kuroh et al. 2005; Vermulst et al. 2008]. There are many genetic factors that have the potential to shorten life span [Kuro-o et al. 1997; Hasty et al. 2003; Mounkes et al. 2003; Niedernhofer et al. 2006]. However, the human genetic factor that has been specifically identified as present on human chromosome 4q is present in the same region as Cisd2, and this is highly suggestive. Nonetheless, experiments that shorten life span might be less informative than those that prolong a healthy life span [Kurosu et al. 2005; Schriner et al. 2005; Pinton et al. 2007]. Accordingly, it will be of great interest to evaluate the life history of transgenic mice expressing elevated levels of Cisd2 protein to see whether Cisd2 is the genetic determinant on human chromosome 4 that may regulate mammalian longevity and allow an unusually long life span to be achieved.

Materials and methods

Generation of the Cisd2 knockout mouse

Mouse Cisd2 genomic DNA was obtained by screening a BAC library [Research Genetics, Inc.] derived from the C57BL/6 (B6) mouse strain. A SpeI–BamHI 6.4-kb DNA fragment, which contains part of intron 1, exon 2, and part of exon 3 of the Cisd2 gene, was used as the homologous recombination arms for construction of an insertion-type targeting vector [Supplemental Fig. 4]. The Cisd2 targeting vector, containing the puromycin selection cassette, was linearized with Apal and transfected into AB2.2 ES cells using electroporation. Targeted ES cell clones were screened by Southern blot analysis using a 3′-flanking probe, specifically a 1.7-kb BamHI–EcoRI fragment from exon 3. Targeted ES cells were injected into B6 blastcysts. Chimeric male mice were bred with B6 females. Germline transmission was obtained from their agouti progeny. The mice were bred in a specific pathogen-free facility and treated according to the
was performed using paraffin-embedded pancreas sections (3 μm). Pancreas sections were soaked in antigen retrieval buffer containing 10 mM sodium citrate (pH 6.0) and heated in a microwave oven twice for 10 min [Surnpentown, SM-1220, 650W]. The sections were then incubated with primary antibody against insulin [1:100; Abcam, ab7842 guinea pig polyclonal antibodies] for 18–24 h at 4°C, detected by biotinylated secondary antibodies [1:500, Abcam, ab6907], and visualized by the LSAB Kit (DakoCytomation, K0690).

**Rabbit anti-mouse Cisd1 and Cisd2 polyclonal antibodies**

Mouse cDNA fragments of Cisd1 (corresponding to amino acids 27–108) and Cisd2 (corresponding to amino acids 52–135) were amplified by PCR and cloned into the pQE-31 (Qiagen) vector, which contains a His tag sequence. The expression plasmids for His-Cisd1 and His-Cisd2 were transformed into M15 bacteria, induced by 2 mM IPTG (isopropylthio-β-D-galactoside), and purified using Nickel-resin (Novagen). These proteins were injected into rabbits to generate antisera containing polyclonal antibodies against the mouse Cisd1 and Cisd2, respectively.

**Subcellular localization**

The EGF-tagged Cisd2 expression plasmids were transfected into NIH/3T3 cells using lipofectamin 2000 [Invitrogen, 11668-019]. Cells transiently expressing EGF-Fis-Cisd2 fusion proteins were plated on gelatin-coated glass coverslips, stained with various organelle probes including mitochondria (MitoTracker Red CMXRos; Invitrogen Life Technologies), ER [anti-calnexin; Sigma, C7617], and Golgi apparatus (anti-Golgi 97; Molecular Probes). The coverslips were then fixed and visualized by confocal microscopy (Olympus Fluoview FV300). Nuclei were counterstained with DAPI (4′,6-diamidino-2-phenylindole; Sigma).

**Isolation of mitochondria from skeletal muscle**

Fresh skeletal muscles were washed twice with PBS and homogenized immediately in ice-cold SEH buffer [0.25 M sucrose, 1 mM EGTA, 3 mM HEPES, protease inhibitor cocktail at pH 7.2]. The mitochondrial pellet was obtained by low-speed centrifugation (10,000 g) to separate the OM and IMS of the mitochondria. The mitoplast (containing the IM and matrix) was pelleted, and the supernatant was subjected to high-speed centrifugation (12,000 g) to further separate the OM and IMS of the mitochondria.

**Mitochondrial subfractionation**

Mitochondrial subfractionation was performed according to the method described by Pagliarini et al. (2005) with some modifications. Briefly, 0.5 mL of mitochondrial suspension (10 mg/mL) was incubated with 10 mg of purified digitonin on ice for 30 min, and the mixture was gently inverted every 10 min. After high-speed centrifugation (12,000g), the mitoplast (containing the IM and matrix) was pelleted, and the supernatant was subjected to ultracentrifugation (150,000g) to separate the OM and IMS of the mitochondria.

**Measurement of oxygen consumption**

The oxygen consumption rate was measured using a 782 Oxygen Meter (Strathkelvin Instruments). An aliquot of 300 μL of assay buffer [125 mM sucrose, 65 mM KCl, 2 mM MgCl2, 20 mM Na+,K+-phosphate buffer at pH 7.2] containing ~0.2–0.5 mg of mitochondria was delivered into the closed chamber of the oxygen meter at 37°C to measure the steady-state oxygen consumption rate of the mitochondria (Chen et al. 2008). In order to further estimate the respiratory function of mitochondria, we measured the glutamate-malate-supported respiration and RCR of mitochondria. First, we used a Hamilton syringe (Strathkelvin) to add 10 mM glutamate and 10 mM malate (Sigma-Aldrich) into the chamber as the electron donor and recorded the glutamate–malate-supported oxygen consumption rate. After 5 min, we injected 3 μL of 100 mM ADP to attain a final ADP concentration of 1 mM in the assay medium. The rate of activated respiration was recorded to measure the RCR of the mitochondria.

**Respiratory enzyme complex activity**

The following activity assays were performed according to the method described by Wei et al. (1998). The activities of NCCR (which represents complex I–III activity) and SCCR (which represents complex II–III activity) were measured by following the reduction of exogenous oxidized cytochrome c. An aliquot of 20–50 μg of sub mitochondrial particles [SMP] was preincubated with the assay buffer [1.5 mM KCN, 50 mM K2HPO4 at pH 7.4] containing β-NADH or succinate for 15 min at 37°C. After addition of cytochrome c to the mixture, the change in absorbance at 550 nm was recorded on a UV/visible spectrophotometer. CCO (which represents complex IV activity) was determined by following the oxidation of exogenous reduced cytochrome c. An aliquot of 20–50 μg of SMP was preincubated in the assay buffer [5 mM K2HPO4 at pH 7.4] for 10 min at 30°C. After addition of ferrocytochrome c to the assay mixture, the change in absorbance at 550 nm was recorded on a UV/visible spectrophotometer.

**Oral glucose tolerance test and insulin tolerance test**

Mice after a 10-h fast (10 p.m. to 8 a.m.) were orally administrated with glucose solution [1.5 g/kg body weight] using a feeding needle [Juan et al. 2004]. Blood samples were collected from tail tips before [0 min] and after glucose load at the indicated time points. The blood glucose levels were measured using glucose test strips (LifeScan, Johnson & Johnson) and SureStep Brand Meter. Serum insulin levels were determined by an ELISA kit (Mercodia). The insulin tolerance test was performed after a 2-h fast [9 a.m. to 11 a.m.] and involved an intraperitoneal injection of insulin [0.75 U/kg body weight, Novolin human regular insulin; Novo Nordisk] [Tran et al. 2008]. There were three mice for each group and three independent measurements for each mouse.

**Statistics**

Results are presented as means ± SD. Differences among multiple groups were analyzed by a one-way ANOVA (SPSS 14.0 statistical software). Comparisons between two groups were done using a Student’s t-test. Mouse survival rates were calculated by the Kaplan-Meier method, and differences in the survival of different groups of mice were determined by the log-rank (Mental-Cox) test. When analyzing statistical differences between the knockout and wild-type mice, P < 0.05 was considered significant.

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Cisd2 deficiency drives premature aging and causes mitochondria-mediated defects in mice

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