Title:

Transmitochondrial mice as models for primary prevention of diseases caused by mutation in the $tRNA^{Lys}$ gene

Short title:

Mice carrying a mitochondrial tRNA mutation

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ABSTRACT

We generated transmitochondrial mice (mito-mice) that carry a mutation in the $tRNA^{Lys}$ gene encoded by mitochondrial DNA (mtDNA) for use in studies of its pathogenesis and transmission profiles. Because patients with mitochondrial diseases frequently carry mutations in the mitochondrial $tRNA^{Lys}$ and $tRNA^{Leu(UUR)}$ genes, we focused our efforts on identifying somatic mutations of these genes in mouse lung carcinoma P29 cells. Of the 43 clones of PCR products including the tRNA^{Lys} or tRNA^{Leu(UUR)} genes in mtDNA of P29 cells, one had a potentially pathogenic mutation (G7731A) in the tRNA^{Lys} gene. P29 subclones with predominant amounts of G7731A mtDNA expressed respiration defects, thus suggesting the pathogenicity of this mutation. We then transferred G7731A mtDNA into mouse ES cells and obtained F₀ chimeric mice. Mating these F₀ mice with C57BL/6J (B6) male mice resulted in the generation of F₁ mice with G7731A mtDNA, named 'mito-mice-tRNALys7731'. Maternal inheritance and random segregation of G7731A mtDNA occurred in subsequent generations. Mito-mice-tRNA^{Lys7731} with high proportions of G7731A mtDNA exclusively expressed respiration defects and diseaserelated phenotypes, and therefore are potential models for mitochondrial diseases due to mutations in the mitochondrial $tRNA^{Lys}$ gene. Moreover, the proportion of mutated mtDNA varied markedly among the pups born to each dam, suggesting that selecting oocytes with high proportions of normal mtDNA from affected mothers with tRNA^{Lys}based mitochondrial diseases may be effective as primary prevention for obtaining unaffected children.

Keywords: mouse mtDNA mutation; mitochondrial $tRNA^{Lys}$ gene; transmitochondrial mice; mutated mtDNA segregation; primary prevention

Significance statement

We generated transmitochondrial mito-mice-tRNA^{Lys7731} as models for precise examination of the pathogenesis and transmission profiles of mtDNA mutations in the *tRNA^{Lys}* genes, and have obtained important information regarding primary prevention of the diseases caused by the mtDNA mutations. Although nuclear transplantation from oocytes of affected mothers into enucleated oocytes of unrelated women has been suggested, the methodology carries the technical risk of inducing nuclear abnormalities and prompts ethical concerns regarding the production of three-parent babies with normal mtDNA from unrelated oocyte donors. The current study suggests that the selection of oocytes with high proportions of normal mtDNA from affected mothers can be used to avoid these issues and therefore provides new insights into mitochondrial genetics and medicine.

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Introduction

Mitochondrial DNA (mtDNA) carrying a large-scale deletion (Δ mtDNA), single point mutations in the $tRNA^{Lys}$ gene, and in the $tRNA^{Leu(UUR)}$ gene causes CPEO (chronic progressive external ophthalmoplegia), MERRF (myoclonic epilepsy with ragged-red fibers), and MELAS (mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episodes), respectively—the three most prevalent mitochondrial diseases (1-3). However, there are slight differences among the three disease phenotypes, even though these pathogenic mtDNA mutations all induce mitochondrial respiration defects. Considering that mitochondrial respiratory function is controlled by both mitochondrial and nuclear genomes (1-3), this controversial issue can be clarified by generating transmitochondrial mice ('mito-mice') that share the same nuclear genetic background

but carry different pathogenic mtDNA mutations corresponding to the mutations found in the three prevalent mitochondrial diseases. However, no well-established effective protocols are available for introducing mutagenized mtDNA into the mitochondria of mammalian cells.

In our previous studies (4-8), we found mtDNAs carrying pathogenic mutations in mouse cell lines, transferred them into mouse female germlines, and generated several types of mito-mice, including mito-mice- Δ (4, 5) which harbor Δ mtDNA and therefore are disease models for CPEO. However, mito-mice harboring mtDNA with pathogenic mutations in the $tRNA^{Lys}$ and $tRNA^{Leu(UUR)}$ genes—and therefore prospective disease models of MERRF and MELAS, respectively—have not previously been established due to the unavailability of mouse cell lines with corresponding tRNA mutations in mtDNA.

To complement the paucity of effective technologies required for introducing mutagenized mtDNA into mitochondria of living mouse cells, we here developed an alternative strategy involving cloning and sequence analysis to detect small amounts of mtDNA with somatic mutations in the mitochondrial tRNA genes. Because pathogenic mutations responsible for mitochondrial diseases occur preferentially in the $tRNA^{Leu(UUR)}$ and $tRNA^{Lys}$ genes of humans (1-3), we sequenced 43 clones generated from PCR products carrying the mitochondrial $tRNA^{Leu(UUR)}$ and $tRNA^{Lys}$ genes of P29 mouse lung carcinoma cells (9). One of the 43 clones had a somatic G7731A mutation in $tRNA^{Lys}$, which enabled the generation of transmitochondrial mito-mice expressing respiration defects for their use as models for diseases caused by mutations in the mitochondrial $tRNA^{Lys}$ gene.

Results

Cloning and sequence analysis of PCR products including tRNA genes

To detect small proportion of mtDNA with somatic and possibly pathogenic mutations in the $tRNA^{Leu(UUR)}$ and $tRNA^{Lys}$ genes, we used two sets of primers so that the resulting PCR products would include the $tRNA^{Leu(UUR)}$ or $tRNA^{Lys}$ gene of P29 cells (Fig. S1). After cloning the PCR products, we sequenced all 43 clones obtained and compared the resulting sequences with that of mtDNA from P29 cells (Table 1). We found one and two somatic mutations in the mitochondrial $tRNA^{Leu(UUR)}$ and $tRNA^{Lys}$ genes, respectively. Given that each somatic mutation was present in only one of the 43 clones, the proportion of each in the mtDNA population of P29 cells would be about 2.2%.

The T7728C and G7731A mutations in the *tRNA*^{Lys} gene both occurred in conserved sites (Table S1) and may correspond to pathogenic mutations that induce respiration defects by their accumulation. Moreover, an orthologous mutation to mouse G7731A has been reported to occur in human mtDNA from patients with mitochondrial diseases (10, 11).

Therefore, we selected these mutations in mtDNA for the generation of mito-mice. In an attempt to detect the T7728C and G7731A mutations in P29 mtDNA, we performed *Xmn* I and *Dra* I digestions of the PCR products of mtDNA, because the T7728C and G7731A mutations create an *Xmn* I site and a *Dra* I site, respectively (see SI Materials and Methods). However, the T7728C and G7731A mutations were undetectable due to their insufficient amounts in P29 cells.

Concentration of G7731A mtDNA in subclones of P29 cells

We previously showed that two mtDNA haplotypes with different mutations in single cells segregate stochastically during cell division (12). Therefore, some individual cells in the P29 population may possess detectable amounts of the mutated mtDNA. To obtain some individual cells that had accumulated either T7728C mtDNA or G7731A mtDNA from the P29 cell population, we isolated 100 subclones from P29 cells. Their mtDNA genotyping showed that two subclones, P29-42 and P29-69, possessed 34% and 48% G7731A mtDNA, respectively. However, we did not obtain any subclones carrying detectable amounts of T7728C mtDNA.

In the case of human mitochondrial *tRNA* gene mutations found in patients with mitochondrial diseases, respiration defects were apparent only when the mutated mtDNA had accumulated to more than 90% (13). To isolate P29 cells with more than 90% G7731A mtDNA, we cultured subclone P29-69, which had 48% G7731A mtDNA, for additional 3 months to allow further amplification of G7731A mtDNA through stochastic segregation; we then isolated more than 200 subclones from the P29-69 cells. We obtained one subclone, P29-69-183, which contained 92% G7731A mtDNA (Fig. 1A)—a level likely to be sufficient for the expression of respiration defects, if the G7731A mutation indeed is a pathogenic mutation.

Determination of the pathogenicity of G7731A mtDNA

Comparison of the O₂ consumption rates between parental P29 cells and the P29-69-183 cells revealed the expression of respiration defects in the P29-69-183 cells (Fig. 1B). Moreover, P29-69-183 cells demonstrated slight overproduction of reactive oxygen species (Fig. 1C), indicating the pathogenicity of the G7731A mutation. Whole-sequence analysis of mtDNA in P29-69-183 cells showed that the G7731A mutation is

the only mutation in those cells that is capable of inducing respiration defects (Table S3). However, we had to resolve two important issues before generation of the mitomice carrying G7731A mtDNA.

First, we had to confirm that the respiration defects in P29-69-183 cells were due to G7731A mtDNA and not to the selection of cells with mutations in nuclear DNA that were acquired during repeated recloning. Second, P29-69-183 cells could not be used as mtDNA donors to isolate mouse ES cells with G7731A mtDNA because of the difficulty of excluding unenucleated P29-69-183 cells from fusion mixtures of ES and enucleated P29-69-183 cells.

To simultaneously resolve these issues, we cytoplasmically transferred G7731A mtDNA from P29-69-183 cells into mtDNA-less (ρ^0) B82 cells (6). B82 cells are mouse fibrosarcoma cells that are resistant to 5-bromodeoxyuridine (BrdU) and sensitive to hypoxanthine–aminopterin–thymidine (HAT) due to their deficiency of thymidine kinase activity (6). Moreover, ρ^0 B82 cells are unable to grow in the absence of uridine and pyruvate due to their complete lack of mtDNA. Using selection medium containing BrdU and lacking uridine and pyruvate, we isolated two colonies, B82mt7731-1 and -2. Genotyping of mtDNA showed that B82mt7731-1 and -2 possessed 70% and 95% G7731A mtDNA, respectively (Fig. 2A), indicating the transfer of G7731A mtDNA from P29-69-183 cells into ρ^0 B82 cells.

We then examined the respiratory function of these 2 cybrids by estimating O₂ consumption rates and the amounts of reactive oxygen species. The B82mt7731-2 cybrids showed decreased O₂ consumption rates (Fig. 2B) and increased production of reactive oxygen species (Fig. 2C) compared with those of B82mtB6 cybrids containing

normal mtDNA from B6 mice. Therefore, respiration defects were transferred to B82mt7731 cybrids concurrently with the transfer of G7731A mtDNA from P29-69-183 cells into ρ^0 B82 cells, suggesting that the somatic G7731A mutation in mtDNA is a pathogenic mutation that can induce mitochondrial respiration defects by its predominant accumulation. Furthermore, B82mt7731 cybrids are effective as donors of G7731A mtDNA to ES cells, because unenucleated B82mt7731 cybrids can be excluded by using selection medium containing HAT.

Isolation of ES cells containing G7731A mtDNA and their chimeric mice

Our previous study showed that no chimeric mice were obtained from ES cells carrying predominant amounts of ΔmtDNA, because the significant respiration defects induced by ΔmtDNA inhibited differentiation of ES cells to various tissues and germ cells (14). Therefore, we were concerned that the transfer of mtDNA from B82mt7731-2 cybrids containing 95% G7731A mtDNA to ES cells would inhibit generation of chimeric mice, and instead used B82mt7731-1 cybrids containing lower proportions of G7731A mtDNA (Fig. 2A) as mtDNA donors.

Female (XO)-type ES cells (TT2 cells) were pretreated with rhodamine 6G (R6G) to eliminate endogenous mitochondria and mtDNA. They then were used as recipients of G7731A mtDNA and fused with enucleated B82mt7731-1 cybrids. The fusion mixture was cultured in selection medium containing HAT to exclude unenucleated B82mt7731-1 cybrids. Seven ES clones grew in the selective medium, two of which—clones ESmt7731-4 and -7—contained G7731A mtDNA (Fig. S2). The absence of G7731A mtDNA in the remaining five clones may be due to incomplete elimination of endogenous mtDNA in ES cells during R6G pretreatment.

We then aggregated the ESmt7731-4 and -7 cybrid clones with 8-cell-stage mouse embryos (ICR strain) and obtained 35 F_0 chimeric mice. Because mouse mtDNA is inherited strictly maternally (15, 16), we selected 15 F_0 chimeric female mice with 13%–76% G13997A mtDNA in their tails as founder mice (Table 2) and mated them with B6 male mice to generate F_1 mice that carried G7731A mtDNA due to its transfer through the female germline.

Generation of mito-mice-tRNA^{Lys7731} via female germ line transfer of G7731A mtDNA

Of the 15 F₀ female chimeras, 11 produced a total of 121 F₁ pups, 63 of which had G7731A mtDNA in their tails (Table 2). This finding suggests that G7731A mtDNA was transmitted maternally from F₀ female mice to F₁ progeny. Mice that carried G7731A mtDNA derived from lung carcinoma P29 cells were named 'mito-mice-tRNA^{Lys7731}'. Because the proportion of G7731A mtDNA did not differ significantly between tissues of the same F₁ male mito-mouse-tRNA^{Lys7731} (Fig. S3), we can deduce the approximate overall proportion of G7731A mtDNA in mito-mice-tRNA^{Lys7731} by estimating the proportion of G7731A mtDNA in tail, which we can do without euthanizing them.

Of the 63 F₁ mice with G7731A mtDNA in their tails, seven F₁ female mice carrying high proportions of G7731A mtDNA were mated with B6 male mice to obtain subsequent generations (F₂-F₅) of mito-mice-tRNA^{Lys7731} with sufficient G7731A mtDNA for the expression of respiration defects and resultant disorders. The proportions of G7731A mtDNA varied significantly among the pups born to each dam, but none of the pups carried more than 85% G7731A mtDNA (Fig. 3A).

To examine the reasons for the lack of mito-mice-tRNA^{Lys7731} with more than 85% G7731A mtDNA, we estimated the proportion of G7731A mtDNA in the oocytes obtained by ovarian hyperstimulation of F_5 female mito-mice-tRNA^{Lys7731} with high

proportions of G7731A mtDNA. The results again showed significant variation in G7731A mtDNA proportions among the oocytes and the absence of the oocytes with more than 85% G7731A mtDNA (Fig. 3B), indicating that lethality of oocytes with high levels of G7731A mtDNA is responsible for the absence of mito-mice-tRNA^{Lys7731} carrying more than 85% G7731A mtDNA.

Expression of disorders in mito-mice-tRNA^{Lys7731} with predominant G7731A mtDNA

We used B6 mice as controls and three groups of F₅ mito-mice-tRNA^{Lys7731} with different heteroplasmic conditions (low, intermediate, and high levels of G7731A mtDNA) in their tails to examine various phenotypes relevant to mitochondrial diseases (Fig. 4).

First, we analyzed body length (Fig. 4A) and muscle strength (Fig. 4B), because abnormalities in these characteristics frequently occur in patients with mitochondrial diseases (10, 11), and these parameters can be examined without euthanizing the mice. Unlike mito-mice tRNA^{Lys7731} with low and intermediate levels, those with high levels of G7731A mtDNA showed short body length (Fig. 4A) and muscle weakness (Fig. 4B), which are closely associated with the clinical phenotypes caused by mutations in the mitochondrial *tRNA*^{Lys} gene (2, 10, 11).

We then quantitatively estimated mitochondrial respiratory function and revealed respiration defects in skeletal muscle and kidney from mito-mice-tRNA^{Lys7731} with high levels of G7731A mtDNA (Fig. 4C). Therefore, accumulation of G7731A mtDNA likely is responsible for the respiration defects in mito-mice-tRNA^{Lys7731} with high levels of G7731A mtDNA (Fig. 4C). These respiration defects in the skeletal muscle subsequently result in the expression of muscle weakness (Fig. 4B), which corresponds to a phenotype relevant to mitochondrial diseases (2).

In contrast, other metabolic parameters relevant to mitochondrial diseases were normal (Fig. S4). Histochemical analysis showed that ragged-red fibers (RRF) frequently observed in MERRF patients (2) and renal failures frequently observed in mito-mice-Δ (4, 5) were not found in mito-mice-tRNA^{Lys7731} (Figs. S5 and S6). Absence of these disorders in mito-mice-tRNA^{Lys7731} may be due in part to that G7731A mtDNA proportions in them were not sufficient to induce significant respiration defects. For example, G8344A mtDNA proportions exceed 85% in patients with MERRF (2). Our failure to obtain mito-mice-tRNA^{Lys7731} with more than 85% G7731A mtDNA can be explained by the lethality of mouse oocytes carrying these levels of G7731A mtDNA (Fig. 3B).

These observations suggest that mito-mice-tRNA^{Lys7731} at least in part can serve as models to investigate pathogenesis of mitochondrial diseases that arise due to mutations in the mitochondrial *tRNA* genes. In addition, the transmission profiles of G7731A mtDNA showed that selecting oocytes with lower levels of the mutated mtDNA likely would be effective to prevent maternal transmission of the disease phenotypes to the progeny.

Discussion

The current study generated mito-mice-tRNA^{Lys7731} carrying G7731A mtDNA with a pathogenic G7731A mutation in the mitochondrial *tRNA^{Lys}* gene. Specifically, we concentrated a small proportion of mtDNA with a somatic G7731A mutation that was present in P29 cells and then introduced it into ES cells to generate mito-mice-tRNA^{Lys7731}. The resulting mice were used to investigate the pathogenesis and transmission profiles of G7731A mtDNA.

Regarding the pathogenesis of G7731A mtDNA, mito-mice-tRNA^{Lys7731} with 74%-84% G7731A mtDNA demonstrated respiration defects and the resultant muscle weakness and short body length (Fig. 4). These abnormalities in our mice are very similar to those found in patients with mitochondrial diseases due to human orthologous G8328A mutation (10, 11) and in MERRF patients carrying the G8344A mutation in the tRNA^{Lys} gene (2, 17-19). Although these abnormalities were expressed in mito-mice-tRNA^{Lys7731} with 74%-84% G7731A mtDNA (Fig. 4) and in the patients with 57% (10) and 82% G8328A mtDNA (11), more than 85% G8344A mtDNA was required for the onset of severe abnormalities in patients with MERRF (2, 17-19). However, we could not obtain mito-mice-tRNA^{Lys7731} with more than 85% G7731A mtDNA (Fig. 3A), which would be expected to induce significant respiration defects and severe abnormalities corresponding to MERRF, owing to the lethality of mouse oocytes with more than 85% G7731A mtDNA (Fig. 3B). Therefore, the expression of the significant respiration defects and severe clinical disorders in MERRF patients can be explained by supposing the absence of lethality in human oocytes even at G8344A mtDNA levels exceeding 85%. In contrast, our previous studies (4, 5) showed that mito-mice- Δ possessed more

than 85% Δ mtDNA with a large-scale deletion, and expressed significant respiration defects and severe abnormalities, even though their oocytes did not have more than 80% Δ mtDNA (20). The increase in the proportion of Δ mtDNA after birth may be due to its replication advantage (4, 5).

The absence of abnormalities in mito-mice-tRNA^{Lys7731} with intermediate levels (37%-56%) of G7731A mtDNA (Fig. 4A and B) appears to be different from what is observed for mtDNA with human orthologous G8328A mutation, because the patient with only 57% G8328A mtDNA in the skeletal muscles expressed disorders (10). This apparent discrepancy may be due to the aging effects in the patient with 57% G8328A mtDNA, considering that the phenotypes of the mito-mice-tRNA^{Lys7731} were examined 4 months after the birth (Fig. 4), and that the patient expressed disorders 45 years after the birth (10). Thus, we have to examine whether mito-mice-tRNA^{Lys7731} with intermediate levels of G7731A mtDNA express abnormalities along with their aging.

Regarding transmission profiles, those of mito-mice-tRNA^{Lys7731} are highly similar to those of MERRF patients, in that the mutated mtDNA is inherited to subsequent generations through the female germline, and the proportion of mutated mtDNA varied markedly among the pups born to each dam (Fig. 3A). The pups always carry wild-type mtDNA (Fig. 3A), due to the significant pathogenicity of G7731A mtDNA and resultant induction of oocyte lethality in the absence of wild-type mtDNA (Fig. 3B). In contrast, we previously generated mito-mice-COI⁶⁵⁸⁹ (6) and mito-mice-ND6¹³⁹⁹⁷ (7, 8), which carry a 100% (homoplasmic) T6589C mtDNA mutation and a homoplasmic G13997A mtDNA mutation in the structural genes *COXI* and *ND6*, respectively, due to the mild pathogenicity of the mutations. Because patients with MERRF and mito-mice-

tRNA^{Lys7731} always carry both wild-type mtDNA and mutated mtDNA, mito-mice-tRNA^{Lys7731} are appropriate models for further investigating the transmission profiles of mutated mtDNA with marked pathogenicity.

The heteroplasmic mtDNA (wild-type mtDNA and mutated G7731A mtDNA) of our mito-mice-tRNA^{Lys7731} segregated stochastically, like those derived from different mouse strains (BALB and NZB) with polymorphic mutations (21) and those with and without point mutations (22). The significant variation in the G7731A mtDNA proportions among pups (Fig. 3A) may reflect 'bottleneck effects' with (23, 24) or without decrease in the mtDNA copy number (25, 26) during female germline transmission of the heteroplasmic mtDNAs.

Regarding primary prevention of mitochondrial diseases, the results in Figs. 3 and 4 indicate that the selection of oocytes with low proportions of G7731A mtDNA likely would yield phenotypically normal mice. In contrast, our previous study (20) showed that nuclear transplantation from zygotes of transmitochondrial mito-mice-Δ into enucleated zygotes with normal mtDNA is effective as germline gene therapy. Many recent reports similarly have noted that the use of the nuclear transplantation from human oocytes of affected mothers into enucleated oocytes donated by unrelated women would prevent the transmission of mitochondrial diseases caused by mtDNA mutations to their children (27-29). However, this technology includes the risk of inducing nuclear abnormalities, even though it excludes the risk of mitochondrial abnormalities. This problem would be resolved by the selection of oocytes with low proportions of mutated mtDNA. Previous study showed that polar bodies are effective for preimplantation genetic diagnosis (PGD) to deduce the proportion of mutated

mtDNA in mouse oocytes (20). However, subsequent studies using polar bodies from human oocytes (30) and blastomeres from human embryos (31, 32) indicated that blastomeres are more appropriate than are polar bodies for PGD to deduce the proportion of mutated mtDNA. Although this procedure would not completely exclude the mutated mtDNA from the affected mothers, our previous studies (33, 34) showed the presence of inter-mitochondrial complementation to maintain normal respiratory function in the presence of the mutated mtDNA. Therefore, in light of the results in Figs. 3 and 4, we propose that the selection of embryos with low proportions of mutated mtDNA from affected mothers would be effective for obtaining unaffected children.

The risk of late-onset disorders in subjects with low proportions of mutated mtDNA appears to be negligible, because no subjects with less than 40% MERRF G8344A mutation in the blood expressed any abnormal symptoms throughout their lifetime (17). Moreover, mito-mice-COI⁶⁵⁸⁹ carrying homoplasmic T6589C mtDNA in the *COX1* gene and expressing mild respiration defects showed normal lifespans and no late-onset disorders (6, 8). Although mito-mice-ND6¹³⁹⁹⁷ carrying homoplasmic G13997A mtDNA in the *ND6* gene developed late-onset and age-related disorders (8), the onset was not due to respiration defects but to overproduction of reactive oxygen species, which can be suppressed by the administration of antioxidants (8). Because high proportions of G7731A mtDNA also induced slight overproduction of reactive oxygen species (Fig. 2C), we will examine whether mito-mice-tRNA^{Lys7731} with high proportions of G7731A mtDNA express late-onset disorders due to overproduction of reactive oxygen species.

Materials and Methods

Cell lines and cell culture; Cloning and sequencing of PCR products; Genotyping of mtDNA; Isolation of B82mt7731 cybrids; Isolation of ESmt7731 cybrids; Generation of chimeric mice and mito-mice-tRNA^{Lys7731}; Mice; Oxygen consumption; Measurement of reactive oxygen species in mitochondria; Histopathologic analyses; Grip strength test; Biochemical measurements of respiratory enzyme activity; Measurement of blood glucose, lactate, and blood urea nitrogen (BUN); Measurement of hematocrit values; Sequence analysis; Statistical analysis. These are detailed in *SI Materials and Methods*.

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Figure legends

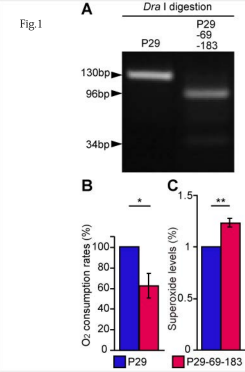
Fig. 1 Characterization of P29-69-183 cells to determine the pathogenicity of a G7731A mutation in the mitochondrial $tRNA^{Lys}$ gene. (A) Estimation of the proportion of

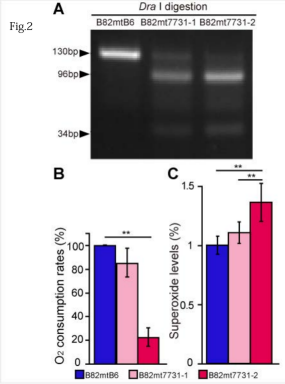
G7731A mtDNA in P29 and P29-69-183 cells by Dra I digestion of the PCR products. The G7731A mtDNA produced 96-bp and 34-bp fragments due to the gain of a DraI site through G7731A substitution in the $tRNA^{Lys}$ gene, whereas mtDNA without the mutation produced a 130-bp fragment. Quantitative estimation of G7731A mtDNA showed that P29-69-183 cells contained 92% G7731A mtDNA. (B) Estimation of O2 consumption rates of P29 and P29-69-183 cells. *, P < 0.05. (C) Estimation of mitochondrial superoxide levels in P29 and P29-69-183 cells after their treatment with MitoSOX Red. **, P < 0.01.

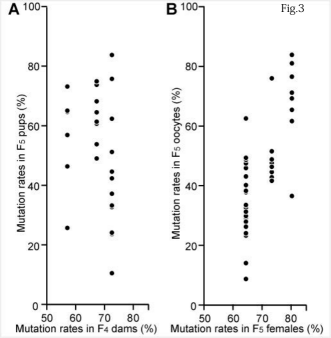
Fig. 2 Characterization of transmitochondrial B82mt7731 cybrids for their use as G7731A mtDNA donors to ES cells. (A) Estimation of the proportion of G7731A mtDNA in cybrids clones B82mt7731-1 and B82mt7731-2 by Dra I digestion of the PCR products. Quantitative estimation of G7731A mtDNA showed that B82mt7731-1 and B82mt7731-2 had 70% and 95% G7731A mtDNA, respectively. (B) Estimation of O2 consumption rates. **, P < 0.01. (C) Estimation of mitochondrial superoxide levels after treatment with MitoSOX Red. **, P < 0.01.

Fig. 3 Variation of G7731A mtDNA proportions among pups or oocytes from individual female mice. (A) Variation of G7731A mtDNA proportions among F₅ pups born to three F₄ dams. G7731A mtDNA proportions were estimated by using tails from F₅ pups that were obtained from three F₄ dams with high proportions of G7731A mtDNA. (B) Variation of G7731A mtDNA proportions among oocytes obtained from three F₅ female mice. G7731A mtDNA proportions were estimated by using oocytes obtained by ovarian hyperstimulation of F₅ female mito-mice-tRNA^{Lys7731} with high proportions of G7731A mtDNA.

Fig. 4 Characterization of F_5 male mito-mice-tRNA^{Lys7731} according to phenotypes associated with mitochondrial diseases. Comparison of (A) body length and (B) muscle (grip) strength between B6 mice (n=3) and F_5 mito-mice-tRNA^{Lys7731} carrying low (13%–15%; n = 3), intermediate (37%–56%; n = 3), and high (76%–84%; n = 3) proportions of G7731A mtDNA. Body length and grip strength were measured at 4 months after birth. Data are presented as means \pm SD. *, P < 0.05; **, P < 0.01. (C) Comparison of activities of mitochondrial respiratory complexes (I+III, II+III, and IV) between B6 mice and F_5 mito-mice-tRNA^{Lys7731} carrying high proportions (76–84%) of G7731A mtDNA in the skeletal muscle and the kidney at 4 months after birth. Respiratory complex I (NADH dehydrogenase), complex II (succinate dehydrogenase), complex III (cytochrome c reductase), and complex IV (cytochrome c oxidase) are components of the electron-transport chain. Enhanced activity of complex II+III in mito-mice-tRNA^{Lys7731} would be due to compensatory activation of complex II, which is controlled exclusively by nuclear DNA. Data are presented as means \pm SD (n = 3). *, P < 0.05; **, P < 0.01.







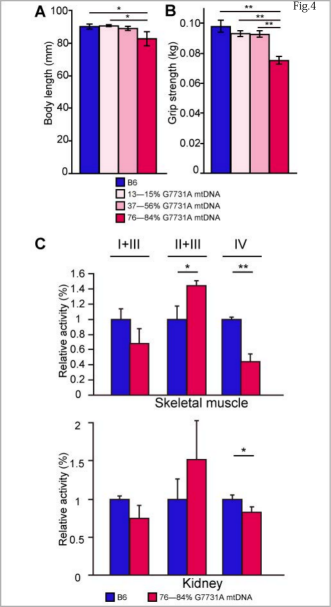


Table 1 Somatic mutations in P29 mtDNA according to cloning and sequence analysis of the PCR products including the *tRNA*^{Leu(UUR)} and *tRNA*^{Lys} genes

	tRNA genes	Leu ^{UUR}	lle	Met	Ser ^{CUN}	Lys	Lys
	nucleotide position	2721	3760	3883	6931	7728 [†]	7731 [‡]
No. of clones	P29 sequence*	Т	G	-	Т	Т	G
1		del	-	-	-	-	=
1		-	Α	-	-	-	=
1		-	-	insA	-	-	-
1		-	-	-	С	-	-
1		-	-	-	-	С	-
1		-	-	-	-	-	Α
37		-	-	-	-	-	-

^{*} Registered under the GenBank accession number EU312160.

[†] The T7728C mutation in the *tRNA^{Lys}* gene occurred in a site highly conserved throughout animals and fungi (see Table 2). The mutation affects the first base of an anticodon triplet (TTT), resulting in an anticodon swap from TTT to CTT (Lys to Glu).

[‡] The mutation equivalent to the G7731A mutation in the *tRNA^{Lys}* gene of mice has been reported to occur in the mtDNA from human patients with mitochondrial diseases (10, 11).

Table 2 Generation of F_0 chimeric mice and their F_1 progeny that carry G7731A mtDNA in their tails.

		15 F ₀ chimeric	F ₁ mice		
			% G7731A	No. of F ₁ pups with	
	% G7731A		mtDNA	G7731A mtDNA / No.	
ES clones	mtDNA	% Chimerism*	in tails	of F ₁ pups	
ESmt7731-4	49.8	95	68	4/4	
		95	53	2/12	
		90	45	24/24	
		50	48	0/17	
		40	16	0/0	
		35	21	0/8	
ESmt7731-7	48.8	95	72	19/19	
		95	76	4/4	
		95	72	2/12	
		95	68	8/8	
		80	52	0/8	
		60	61	0/5	
		40	26	0/0	
		15	13	0/0	
		10	19	0/0	
				Total 63/121	

^{*} Chimerism was judged by coat color.

SI Figure Legends

Fig. S1 Nucleotide positions of PCR primers on the mouse mtDNA map. To detect small proportions of mtDNA with somatic but possibly pathogenic mutations in the tRNA^{Leu(UUR)} and tRNA^{Lys} genes of P29 cells, we used two sets of PCR primers so that the resultant PCR products included the $tRNA^{Leu(UUR)}$ and $tRNA^{Lys}$ genes, respectively. These products also included five additional tRNA genes (tRNA^{Ile}, tRNA^{Gln}, tRNA^{Met}, $tRNA^{Ser(CUN)}$, and $tRNA^{Asp}$).

Fig. S2 Identification of G7731A mtDNA in ESmt7731 cybrid clones grown in selection medium. We were able to obtain two ESmt7731 cybrid clones carrying G7731A mtDNA (ESmt7731-4 and -7) but were unable to exclude ES cells that lacked G7731A mtDNA, owing to incomplete elimination of endogenous mtDNA by R6G treatment. Accordingly, ESmt7731-1, -2, -3, -5, and -6 correspond to ES cells that lacked G7731A mtDNA but contained endogenous mtDNA.

Fig. S3 Proportions of G7731A mtDNA in the tissues from a single F₁ male mouse. Note that the proportion of G7731A mtDNA did not differ significantly among different tissues from the same F₁ male mito-mice-tRNA^{Lys7731}, suggesting that we can deduce the overall proportion of G7731A mtDNA in mito-mice-tRNA^{Lys7731} from that in the tail without euthanizing mice.

Fig. S4 Analyses of metabolic parameters related to mitochondrial diseases by using peripheral blood from mito-mice-tRNA^{Lys7731} with high proportions of G7731A mtDNA. (A) Blood urea nitrogen. (B) Hematocrit. (C) Blood glucose levels before and after glucose administration. (D) Blood lactate levels before and after glucose administration. No statistically significant differences between B6 and mito-mice-tRNA^{Lys7731} with high

proportions of G7731A mtDNA were obtained. Data are presented as mean \pm SD (n = 3).

Fig. S5 Histopathologic analysis to identify ragged red fibers (RRF) in skeletal muscle. Cryosections (thickness, $10~\mu m$) of skeletal muscle were stained by using modified Gomori trichrome for histopathologic analysis to identify RRF. No RRF were present in mito-mice-tRNA^{Lys7731} with high proportions of G7731A mtDNA. Bar, $50~\mu m$.

Fig. S6 Morphology and histopathologic analysis of kidney. (A) Morphology of kidneys from B6 mice (left) and mito-mice-tRNA^{Lys7731} with 76% G7731A mtDNA (right). Bar, 1 cm. (B) Histopathology of the renal cortex of kidneys from B6 mice (left) and mito-mice-tRNA^{Lys7731} with 76% G7731A mtDNA (right). Formalin-fixed, paraffin-embedded sections (thickness, 5 μ m) were stained with hematoxylin and eosin. No renal failure was noted in mito-mice-tRNA^{Lys7731}. Bar, 50 μ m.

Sl Materials and Methods

Cell lines and cell culture. Mouse P29 cells, their subclones, mtDNA-less (ρ^0) B82 cells, and their transmitochondrial cybrid clones were grown in DMEM (Sigma-Aldrich) containing 10% fetal calf serum, uridine (50 ng/ml), and pyruvate (100 ng/ml). Mouse ES cells (TT2-F, an XO subline established from XY TT2 cells; Riken BioResource Center) and transmitochondrial ES cybrids were cultivated on mitomycin-C-inactivated feeder cells derived from fetal fibroblasts in DMEM (Invitrogen) supplemented with 15% Knockout Serum Replacement (Invitrogen), non-essential amino acids (MP Biomedicals LLC), leukemia inhibitory factor (105 U/ml, Invitrogen), and 100 μ M 2-mercaptoethanol (Sigma–Aldrich).

Cloning and sequencing of PCR products. Total DNA extracted from P29 cells was used as the template for PCR amplifications, which generated two mtDNA fragments, F1 and F2, by using unique primer pairs that were designed to generate 2.6- and 2.7-kb fragments, respectively. The sequences of the primers were based on the standard mtDNA sequence of B6 mice (GenBank accession no., AY172335): F1 forward, nucleotides 2045–2064; F1 reverse, 4623–4603; F2 forward, 5931–5950; and F2 reverse, 8626–8606 (Fig. S1). All PCR amplifications were performed in 50 μl of

solution consisting of PCR buffer, 0.2 mM dNTPs, 0.6 mM primers, 1 U Ampli*Taq* Gold DNA polymerase (Perkin-Elmer Applied Biosystems), and 1 μg of cellular DNA as template. Reaction conditions were 95 °C for 10 min, followed by 35 cycles of 60 s at 95 °C, 60 s at 50–56 °C, and 150 s at 72 °C. PCR products were ligated into pUC118 (Takara) and then introduced into DH5α cells (Takara). Sequence templates were prepared by using a TempliPhi DNA Sequencing Template Amplification Kit (Amersham Pharmacia Biosciences) according to the manufacturer's protocol. Sequence reactions were performed by using dye termination methods (Takara PCR Thermal Cycler GP), and samples were sequenced on a MegaBACE1000 automated sequencer (Amersham Pharmacia Biosciences).

Genotyping of mtDNA. To detect the T7728C mutation, a 71-bp fragment containing the 7728 site was PCR-amplified by using the nucleotide sequences 7703 to 7723 (5'-TATGAAGCTAAGAGCGgaAAC-3', small letters indicate mismatch) and 7773 to 7754 (5'-TGTGGCATATCACTATGGAG-3') as oligonucleotide primers. Combination of the PCR-generated mutation with the T7728C mutation creates a restriction site for *Xmn* I and generates 50- and 21-bp fragments on *Xmn* I digestion of the PCR products. To detect the G7731A mutation, a 130-bp fragment containing the 7731 site was PCR-amplified by using the nucleotide sequences from 7633 to 7653

(5'-GCCCATTGTCCTAGAAATGGT-3') and 7762 to 7732 (5'-ACTATGGAGATTTTAAGGTCTCTAACTTTAA-3') as oligonucleotide primers. The G7731A mutation creates a restriction site for *Dra* I and generates 96- and 34-bp fragments on *Dra* I digestion of PCR products. The restriction fragments were separated by electrophoresis in a 3% agarose gel. For quantification of G7731A mtDNA, we used the NIH Image program.

Isolation of B82mt7731 cybrids. We used mtDNA-less (ρ^0) B82 cells as recipients for G7731A mtDNA. ρ^0 B82 cells are resistant to BrdU and sensitive to HAT due to their deficiency of thymidine kinase activity. Furthermore, ρ^0 B82 cells are unable to grow in the absence of uridine and pyruvate due to their absence of mtDNA. mtDNA donor P29-69-183 cells were pretreated with cytochalasin B (10 μg/ml) for 10 min and centrifuged at 7,500 × g for 10 min at 37 °C for enucleation; the resultant cytoplasts were fused with ρ^0 B82 cells by using polyethylene glycol. The fusion mixture was cultured in selection medium containing BrdU (30 μg/ml) and lacking uridine and pyruvate. The selection medium excluded unenucleated P29-69-183 cells and unfused ρ^0 B82 and thus allowed exclusive growth of transmitochondrial B82mt7731 cybrids.

Isolation of ESmt7731 cybrids. Host ES cells were pretreated with rhodamine 6G (R6G; 0.38–1.5 μg/ml in 3% ethanol) for 48 h to eliminate endogenous mitochondria and mtDNA (1, 2). The ES cells then were washed with PBS and suspended in R6G-free medium for 2 h to allow recovery. mtDNA donor B82mt7731 cybrids were pretreated with cytochalasin B (10 μg/ml) for 10 min and centrifuged at 22,500 × g for 30 min at 37 °C for enucleation. The resultant cytoplasts were fused with R6G-pretreated ES cells by using polyethylene glycol, and the fusion mixture was cultured in selection medium containing HAT to exclude unenucleated B82mt7731 cybrids. At 7 days after fusion, growing colonies were harvested for mtDNA genotyping. Under these selection conditions, ES cells containing recovered endogenous mtDNA were not eliminated completely, owing to insufficient R6G treatment.

Generation of chimeric mice and mito-mice-tRNA^{Lys7731}. Frozen 8-cell-stage embryos of ICR mice (ARK Resource Company) were thawed, and their zona pellucidae were removed by treatment with acidified Tyrode's buffer (Sigma–Aldrich). Each treated embryo was placed with about 50 ESmt7731 cybrids in a well of a 35-mm culture dish and incubated overnight to enable aggregation. The next day, the embryos were transferred into pseudopregnant ICR female mice (Japan SLC). The resulting

progeny were identified by their coat-color chimerism. Founder (F_0) chimeric female mice were mated with C57BL/6 J (B6; CLEA Japan) male mice to produce the F_1 generation, and F_1 female mice with G7731A mtDNA in their tails $(F_1$ female mito-mice-tRNA^{Lys7731}) were backcrossed with B6 male mice to obtain F_2 mito-mice-tRNA^{Lys7731}.

Mice. Inbred B6 mice generated through more than 40 rounds of brother–sister mating were obtained from CLEA Japan. We maintained mito-mice-tRNA^{Lys7731} by repeated backcrossing of these female mice with B6 male mice. Animal experiments were performed in accordance with protocols approved by the Experimental Animal Committee of the University of Tsukuba, Japan (approval number, 12070).

Oxygen consumption. Cells were washed once in PBS and then resuspended in PBS at a density of 4×10^6 cells/ml. Oxygen consumption was measured by using an oxygraph equipped with a Clark-type electrode (model 5300, Yellow Springs Instruments Company) at 37 °C. The cell suspension (2 ml) was transferred to a polarographic cell, and basal respiration was measured immediately under constant stirring.

Measurement of reactive oxygen species in mitochondria. Reactive oxygen species were detected by using the mitochondrial superoxide indicator MitoSOX-Red

(Invitrogen). Cells were incubated with 1 mM MitoSOX-Red for 15 min at 37 °C in PBS, washed twice with PBS, and then immediately analyzed by using flow cytometry (FACScan, Becton Dickinson).

Histopathologic analyses. Formalin-fixed, paraffin-embedded sections (thickness, 5 μ m) were stained with hematoxylin and eosin to identify cases of renal failure. Cryosections (thickness, 10 μ m) of skeletal muscle were stained by using a modified Gomori trichrome to identify RRF.

Grip strength test. Muscle strength was estimated by using a Grip Strength Meter (Columbus Instruments); three sequential trials were performed on each mouse.

Biochemical measurements of respiratory enzyme activity. Mitochondrial respiratory complex I (NADH dehydrogenase), complex II (succinate dehydrogenase), and complex III (cytochrome c reductase) are components of the electron-transport chain and are located in the mitochondrial inner membrane. The activity of these enzymes was assayed as described previously (4). Briefly, to estimate the activity due to complex I + III, NADH and cytochrome c (oxidized form) were used as substrates, and the reduction in cytochrome c was monitored by measuring absorbance at a wavelength of 550 nm. To estimate the activity due to complex II + III, sodium succinate and

cytochrome c (oxidized form) were used as substrates, and the reduction of cytochrome c was monitored as described previously.

Measurement of blood glucose, lactate, and blood urea nitrogen (BUN). To determine fasting blood lactate and glucose concentrations, peripheral blood was collected from the tail veins of mice after overnight fasting from food. After oral administration of glucose (1.5 g/kg body weight), blood was collected again, and lactate and glucose concentrations were measured by using an automatic blood lactate test meter (Lactate Pro; Arkray) and glucose test meter (Dexter ZII; Bayer), respectively. BUN was measured by using a Urea N B test (Wako Pure Chemical, Osaka, Japan) in accordance with the manufacturer's protocol.

Measurement of hematocrit values. To determine hematocrit levels, capillary blood was collected from each mouse into heparinized capillary tubes and centrifuged at $10,500 \times g$ for 5 min. Packed cell volumes were measured by using a hematocrit reader.

Sequence analysis. Each DNA (3 μ g) extracted from P29-69-183 cells and from kidney of mito-mouse-tRNA^{Lys7731} were used to prepare the sequencing libraries. These sequencing libraries were constructed by using the TruSeq DNA LT Sample Prep Kit (Illumina) according to the instructions in TruSeq DNA Sample Preparation Guide Rev.

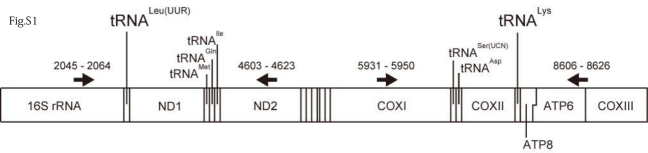
C (Illumina). These sample DNA were sheared by using an acoustic solubilizer (Covaris), and the overhangs resulting from the fragmentation were converted to blunt ends by phosphorylation of the DNA fragments. The blunt fragments were adenylated at their 3' ends were ligated to the ends of the DNA fragments, which were subsequently enriched by PCR. Clonal template clusters were generated from the sequencing library, and sequencing was performed by the reversible terminator-based method (5). Base calling and data conversion into Fastq files were performed by using software providing the Burrows-Wheeler Alignment Tool (6).

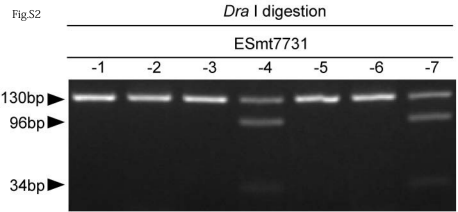
Statistical analysis. Data are presented as means \pm SD and were analyzed by using Student's t test; P values less than 0.05 were considered significant.

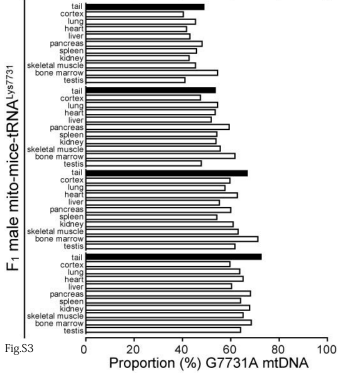
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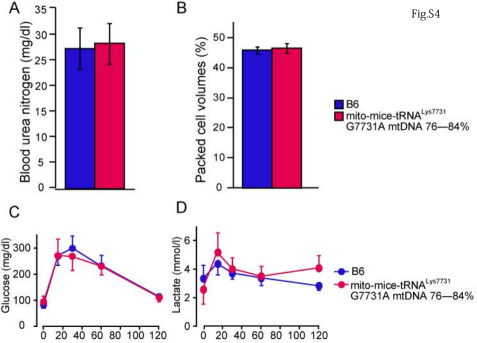
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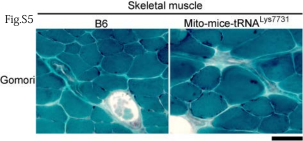
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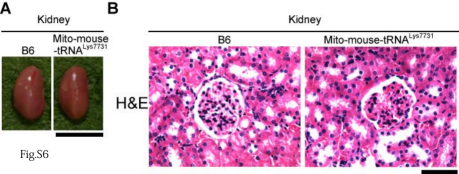


Table S1 Comparison of mutations in the mouse mitochondrial *tRNA* gene with the wild-type *tRNA* genes of other species

Comparison of mutations in the mouse mitochondrial *tRNA* gene with the wild-type *tRNA* genes of other species

	Leu(UUR)	Lys		
	del2721T	T7728C	G7731A	
Mutation	_	С	Α	
Mouse	Т	Т	G	
Human	С	Т	G	
Xenopus	Т	Т	G	
Zebrafish	С	Т	G	
Drosophila	Т	Т	Α	
Nematoda	_	Т	Т	
Yeast	С	Т	G	
		<u></u>		

The GenBank accession numbers of the sequences used in the alignment are: mouse, AY172335; human, NC_012920; *Xenopus*, NC_001573; zebrafish, NC_002333; *Drosophila*, NC001709; Nematoda, NC_001328; yeast, NC_001224.

Table S2 Whole-sequence analysis of the mtDNA from P29-69-183 cells

Whole-sequence analysis of the mtDNA from P29-69-183 cells

genes	tRNA ^{Lys}		tRNA ^{Arg}		D-loop	
nucleotide position	7731	9818~	9821~		16099	16105
P29 sequence*	G	(T) ₃	(A) ₉	(A) ₉	Α	Т
variant sequence	А	(T) ₂	(A) ₈	(A) ₁₀	С	С
P29-69-183**	89.6%†	16.4%	8.2%	6.5%	31.6%	47.6%
mito-mouse-tRNA ^{Lys7731} ‡	80.0%	3.6%	0.5%	18.1%	26.9%	37.0%

^{*} P29 mtDNA sequence, GenBank accession number EU312160.

[‡]Kidney of F₅ mito-mouse-tRNA^{Lys7731} possessing 81.3% G7731A mtDNA quantified by using restriction fragment length polymorphism was used for whole-sequence analysis of the mtDNA. All polymorphic mutations found in mtDNA of P29-69-183 cells have persisted in mtDNA of a mito-mouse-tRNA ^{Lys7731}. Registered under GenBank accession no. AP014541.

[†] Whole-sequence analysis of mtDNA from P29-69-183 cells showed that they had 89.6% G7731A mtDNA. This value was almost equivalent to that obtained in Fig. 1A (92%), suggesting the reliability of our quantification of G7731A mtDNA proportion by using restriction fragment length polymorphism.

^{**}Registered under GenBank accession no. AP014540.