Review

Reverse genetic studies of mitochondrial DNA-based diseases
using a mouse model

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(Communicated by Takao SEKIYA, M.I.A.)

Abstract: In the situation that it would not be able to produce model animals for mitochondrial diseases caused by mitochondrial DNA (mtDNA) with pathogenic mutations, we succeeded in generating mice with pathogenic deletion mutant mtDNA (ΔmtDNA), named “mito-mice”, by direct introduction of mitochondria with ΔmtDNA into mouse zygotes. In the mito-mice, accumulation of ΔmtDNA induced mitochondrial respiration defects in various tissues, resulting in mitochondrial disease phenotypes, such as low body weight, lactic acidosis, ischemia, myopathy, heart block, deafness, male infertility, and renal failure. Thus, mito-mice are the first model animal for mtDNA-based diseases, and the mice could be valuable for understanding precise pathogeneses and testing therapies of mitochondrial diseases. In the present review, we summarized reverse genetic studies using the mito-mice.

Keywords: mouse model, mitochondrial DNA, large-scale deletion mutation, mitochondrial complementation, mitochondrial diseases

Introduction

Mitochondria are one of the organelles, which can produce 90% or more of all the energy made in the body by oxidative phosphorylation. Mammalian mitochondria have multi-copies of own genome (103 and 104 copies/cell), mtDNA, that is replicated and expressed within the organellar system.1,2) Mammalian mtDNA encodes 13 polypeptides which are essential subunits of complexes I, III, IV, and V for oxidative phosphorylation on the inner mitochondrial membrane and 22 tRNAs and 2 rRNAs which are necessary for the translation of these 13 polypeptides. The remaining mitochondrial proteins for oxidative phosphorylation, metabolic enzymes, DNA and RNA polymerases, and ribosomal proteins are all encoded by nuclear genome. In normal individual, nearly all of the mtDNA is thought to be identical. In some cases, however, especially in mitochondrial diseases, wild-type and mutant mtDNAs coexist in cells and organs.

Forward genetic studies, one of the strategies for understanding the genetic reason from phenotypes of the organism, suggested that accumulation of pathogenic mtDNAs having large-scale deletion or point mutation and the resultant mitochondrial respiratory abnormalities are associated with a wide variety of disorders, such as mitochondrial diseases, neurodegenerative diseases, and diabetes.3) Although the pathogenicities of these mtDNA mutations were proved by co-transmission of the mutant mtDNAs and respiration defects to human mtDNA-less cells (rho-0 cells),4–7) there is as yet no convincing evidence to explain whether accumulation of these pathogenic mutant mtDNAs in tissues is responsible for the expressions of various clinical phenotypes.

Reverse genetic studies, one of the strategies for understanding the mechanism of phenotypic expression of organisms from genetic manipulation, could provide model mouse systems for studying exactly how pathogenic mutant mtDNA is transmitted and distributed in tissues resulting in the pathogenesis of mitochondrial diseases that show various clinical phenotypes. However, no procedures are thus far available for introduction of
mutagenized mammalian whole mtDNA into mitochondria in living cells or even into isolated mitochondria. Therefore, generation of mouse models for mitochondrial diseases so far reported has been limited by disruption of nuclear DNA-coded factors related to mitochondrial functions.\cite{8-13} Adenine nucleotide translocator 1 (Ant1)-deficient mice were generated by targeted inactivation of the nuclear-coded Ant1 gene, and showed chronic ATP deficiency, myopathy, and hypertrophic cardiomyopathy.\cite{8} Mn superoxide dismutase (MnSOD)-deficient mice were generated by inactivating mutations in the nuclear encoded SOD2 gene, and showed disease phenotypes in skeletal muscle, heart, liver, and brain.\cite{9} Several kinds of mitochondrial transcription factor A (Tfam)-deficient mice were generated by inactivation of the nuclear Tfam gene.\cite{10} Tissue specific disruption of the Tfam gene in cardiac muscle cells, pancreatic beta-cells, and nervous cells induced dilated cardiomyopathy, diabetes, neurodegenerative diseases, respectively.\cite{11-13}

The effective procedure to generate mouse models carrying pathogenic mutant mtDNA is to introduce exogenously mitochondria carrying the mutant mtDNA by microinjection or cell fusion techniques. Using the cell fusion technique, we have succeeded in generating mice with mitochondrial dysfunction by introduction of mitochondria with somatic mutant mtDNA into mouse zygotes.\cite{14}

**Generation of mice carrying pathogenic deletion mutant mtDNA (mito-mice)**

In generation of mito-mice, ΔmtDNA (Fig. 1A) accumulated in mouse cells was trapped into mouse rho-0 cells\cite{15,16} by isolation of cybrid cells with various proportions of ΔmtDNA, and then respiration-deficient mitochondria were introduced into mouse zygotes using electrofusion with the enucleated cybrids (Fig. 1B).

Methodologically, there are two essential points for successful generation of mito-mice. The first point is to isolate mitochondrial donor carrying mtDNA with somatic mutations from mouse cells. Since the mutation frequency in mtDNA is about 10 times higher than that in nuclear genome, it has been known that somatic mutations occur and accumulate in mtDNA molecules with aging.\cite{8} Thus, we screened the occurrence of somatic mutations in mtDNA molecules in mouse cells by trapping the mutant mtDNAs into mouse rho-0 cells and succeeded in cloning a cybrid cell line carrying both wild-type mtDNA and ΔmtDNA.\cite{15,16} The deleted region of ΔmtDNA is 4,696 bp with a breaking point from nt 7,759–12,454 and includes 6 tRNA genes and 7 structural genes (Fig. 1A). The mouse ΔmtDNA is similar to mtDNA with a common deletion, which have been shown to be responsible for the pathogenesis of a mitochondrial encephalomyopathy, Kearns-Sayre syndrome.\cite{4,17}

The second point for the successful generation of mice carrying mutant mtDNA is to introduce respiration-deficient mitochondria into mouse zygotes (B6 strain) using an electrofusion technique (Fig. 1B). The respiration-deficient cybrids with a predominant amount of ΔmtDNA were used as mtDNA donors for generation of mice with ΔmtDNA. Introduction of ΔmtDNA into mouse zygotes was attained by electrofusion of pronucleus stage embryos with several enucleated cytoplasts of the cybrids. The embryos fused with cytoplasts were cultured in vitro for 24–48 h, and transferred to the oviduct of pseudopregnant females. Since F0 progeny was healthy due to low load of ΔmtDNA (5.7%–13.0% in tails), we selected F0 females with ΔmtDNA as mothers for breeding and obtained progenies with predominant amounts of ΔmtDNA, suggesting the occurrence of ΔmtDNA transmission through female germ lines from F0 mothers to their progeny.

**Clinical and pathological features of mito-mice**

The great advantages of mito-mice are that they all share exactly the same nuclear genome background (B6 strain), and their genetic variation is restricted to the proportions of the introduced pathogenic ΔmtDNA (Fig. 2). Therefore, mito-mice can provide direct evidence that mitochondrial respiration defects induced by the accumulation of ΔmtDNA are sufficient by themselves for expression of the clinical phenotypes observed in patients with mutated mtDNA.\cite{14,18}

In various tissues of mito-mice with high load of ΔmtDNA, approximately more than 75%, mosaic composition of cells with different cytochrome c oxidase (COX, also known as complex IV) activity appeared. Since three of the 13 COX subunits are encoded in mtDNA, it has been well known that the absence of these subunits or/and tRNAs due to mtDNA mutations leads to COX deficiencies
Fig. 1. Genetic characterization of ΔmtDNA molecule and generation of the mito-mice.

A. Gene map of mouse mtDNA. The arc indicates the deleted region (4,696bp) expanded from tRNA^Lys to ND5 genes in ΔmtDNA. B. Experimental scheme for producing mice carrying wild-type mtDNA and exogenously introduced ΔmtDNA. In this scheme, mitochondria possessing wild-type mtDNA are shown in a block color, and those with ΔmtDNA in a gray color. Based on the difference of COX activity, two mitochondrial populations are classified in mouse zygotes after electrofusion. One is COX-positive mitochondria carrying only wild-type mtDNA (black color), which originally exist in mouse zygotes. The other is COX-negative (respiration-deficient) mitochondria carrying predominant amounts of ΔmtDNA (gray color), which is introduced into mouse zygotes from mouse cybrids.
in cells and tissues. In cardiac muscle tissues carrying 88% \( \Delta \text{mtDNA} \), for example, we observed three types of cells, COX-positive, COX-intermediate, and COX-negative cells. Quantitative PCR analysis demonstrated that the COX activity in individual cardiac cells was coordinated with the amount of \( \Delta \text{mtDNA} \); COX-positive, COX-intermediate, and COX-negative cells carried 76%, 83%, and 91% \( \Delta \text{mtDNA} \), respectively (see Fig. 3).

Mito-mice with high load of \( \Delta \text{mtDNA} \) showed also other mitochondrial disease phenotypes due to mitochondrial respiration defects, such as low body weight, lactic acidosis, systemic ischemia, auriculoventricular block with Wenckebach periodicity, hearing loss, renal failures, and male infertility.\(^{14,18,20–22}\) Lactic acidosis is one of the typical clinical phenotypes in patients with mitochondrial respiration defects, and the symptom suggests acceleration of glycolytic pathway for producing ATP without mitochondrial respiration. Early symptoms of the lactic acidosis, low body weight, and sperm abnormalities could be detected in mito-mice carrying approximately more than 75% \( \Delta \text{mtDNA} \).\(^{14,22}\)

\( \Delta \text{mtDNA} \) load is different among the progeny, and proportion of \( \Delta \text{mtDNA} \) in cells and tissues increased with the time\(^{14,23}\) (Fig. 2). In the case of mito-mice carrying more than 70% \( \Delta \text{mtDNA} \) at the birth, they showed mitochondrial respiration defects and disease phenotypes and consequently died on around 6 months after birth.\(^{14,18,20–22}\) On the other hand, mito-mice carrying about 30%–50% \( \Delta \text{mtDNA} \) at the birth were healthy, but they expressed mitochondrial respiration defects and disease phenotypes and died by around 1.5 years after birth when \( \Delta \text{mtDNA} \) load became 75%–85%.\(^{14,18,20–22}\) Thus, mitochondrial respiration defects and the resultant clinical phenotypes were expressed only when proportion of \( \Delta \text{mtDNA} \) exceeded approximately 80%, showing the existence of threshold effects in pathogeneses of mitochondrial diseases (Fig. 3).

Mito-mice carrying a higher amount of \( \Delta \text{mtDNA} \) did not express diabetic phenotypes, and they showed hypoglycemia caused by enhanced glycolysis.\(^{20}\) This result is unexpected and not consistent with the conventional hypothesis of mitochondrial diabetes, which claims that respiration defects caused by mutant mtDNAs in pancreatic beta-cells, skeletal muscle and liver tissues are responsible for their reduced insulin secretion and resistance of insulin and glucose, respectively, leading to expression of diabetic phenotypes.\(^{3,24}\) We cannot yet explain the exact mechanism or
biological significance of enhanced insulin secretion from pancreatic beta-cells carrying predominant /C1 mtDNA. Possibly the increased level of lactic acid in mito-mice expressing mitochondrial respiration defects could enhance insulin secretion, so that accelerated glucose uptake by respiration-deficient target tissues, such as muscle tissues, protected them from progressive glucose deficiency. Recently, it has been reported that protection of diabetic phenotypes were also observed in mice with mitochondrial respiration defects induced by muscle specific disruption of Tfam gene and apoptosis inducing factor (AIF). Thus, the results on mito-mice together with mice expressing mitochondrial respiration defects require reassessment of the conventional concept of mitochondrial diabetes, which proposes mtDNA mutations as major pathogenic factors.

In vivo inter-mitochondrial complementation

Why could mito-mice with low load of ΔmtDNA escape mitochondrial respiration defects and the resultant disease phenotypes? We were able to answer this question by using an electron micrographic technique that identifies the COX activity at individual mitochondrial levels. As shown in Fig. 1B, it can be considered that mouse zygotes that were introduced donor mitochondria carrying ΔmtDNA possess two mitochondrial populations. One is COX-positive host mitochondrion carrying wild-type mtDNA that originally exists in zygotes. The other is COX-negative donor mitochondrion carrying ΔmtDNA that is introduced exogenously.

Considering that individual mitochondria in living cells undergo constant migration, fission, and fusion, there is a possibility that mitochondria could exchange gene products through their fusion and fission. In this case, all mitochondria in single cells of mito-mice would be COX-positive or COX-negative uniformly (Fig. 4), and exogenous ΔmtDNA would be detected even in cells with only COX-positive mitochondria. If exchange of gene products did not occur between exogenous donor COX-negative mitochondria carrying ΔmtDNA and COX-positive mitochondria from recipient zygotes, or even if it occurred, but infrequently and discontinuously, mitochondria in single cells of the mito-mice would continue to show mosaic
distribution of host COX-positive and donor COX-negative mitochondria.

COX electron micrographs of cardiac muscle tissues from mito-mice clearly showed that all mitochondria in single cells with low load of ΔmtDNA, less than approximately 75% ΔmtDNA, were COX-positive (Fig. 3). The appearance of COX-negative mitochondria was limited to cells with more than approximately 85% ΔmtDNA (Fig. 3). In cells with 75%–85% ΔmtDNA load, moreover, COX-intermediate mitochondria were observed (Fig. 3). The appearance of COX-intermediate mitochondria that possess intermediate characters of COX-positive and COX-negative mitochondria indicate the exchange of gene products between COX-positive host mitochondria and COX-negative donor mitochondria. Importantly, it was not observed that the coexistence of COX-positive, COX-intermediate, and COX-negative mitochondria within single cells, irrespective of whether the tissues contained low or high concentrations of ΔmtDNA (Fig. 3). These observations could be explained by in vivo mitochondrial complementation between COX-negative donor mitochondria carrying ΔmtDNA and COX-positive host mitochondria carrying wild-type mtDNA (18,19) (Fig. 4).

Since a threshold level of exogenous ΔmtDNA required to cause mitochondrial dysfunction in all mitochondria in single cells that would be more than approximately 80%, mitochondrial respiratory function could be maintained quite normally, if wild-type mtDNA exists more than 20% in single cells. Although ΔmtDNA has lost 6 tRNA genes, proteins encoded by ΔmtDNA could be translated with the help of the corresponding tRNAs transcribed from wild-type mtDNA. In this case, all mitochondria in single cells could show normal mitochondrial respiratory function (mitochondrial positive complementation) (see Fig. 4).

On the contrary, in somatic cells where the proportion of wild-type mtDNA is less than 20%, mitochondrial translation may become limiting due to the lack of sufficient tRNAs transcribed solely from the wild-type mtDNA. Thus, the translation phase may be shifted from complementation to competition of the tRNAs in cells with more than 80% ΔmtDNA. In this case, progressive inhibition of overall mitochondrial translation and subsequent reduction of the mitochondrial respiratory function would be induced in all mitochondria of single cells (mitochondrial negative complementation) (see Fig. 4), leading to the onset of the disease phenotypes in various tissues.

The occurrence of frequent and continuous
interaction throughout mitochondria in living mice, human cultured cells, and mammalian cells, have provided new concepts on the mitochondrial genetic system in living animals. One was the loss of the individuality of each mitochondrion within single cells. Mammalian cells have been thought to contain hundreds of independent mitochondria, but our observations gave a totally different view that mitochondria function as a single dynamic cellular unit. The other important concept was a novel defense system in mitochondria for avoiding phenotypic expression of respiration defects caused by pathogenic mutant mtDNAs accumulated in somatic tissues. The mitochondrial theory of aging proposed that age-associated accumulation of somatic mutations in mtDNAs is responsible for age-associated mitochondrial dysfunction, since mtDNA is a target of most carcinogens and mutagens and is continuously exposed to reactive oxygen species produced in mitochondria. In fact, accumulation of various kinds of acquired mutations in mtDNAs with age was reported in human subjects. However, this mitochondrial theory of aging is based on above circumstantial evidence, and there have been no reports directly proving that mtDNAs in aged subjects could induce age-associated mitochondrial defects. We found that extensive and continuous interaction occurred between mitochondria in various somatic tissues of mice with ΔmtDNA, resulting in protection of mice from expressing severe respiration defects.

Based on these views, we would like to propose a novel hypothesis on mitochondrial biogenesis, “interaction theory of mammalian mitochondria”: they exchange gene products, and thus lose the individuality and function as a single dynamic cellular unit. This hypothesis has been confirmed by creation of recombinant mtDNA molecules in vivo. For examination of in vivo mtDNA recombination, mito-mouse possessing ΔmtDNA of Mus musculus domesticus and wild-type mtDNA of Mus spretus were used. To avoid PCR jumping artifacts, PCR products were not used, but whole length mtDNAs purified from mito-mouse tissues by EtBr-CsCl centrifugation were used for cloning and sequence analyses. Of the 318 mtDNA molecules, only three molecules possessed specific mutation sites of both M. m. domesticus and M. spretus mtDNAs. Presence of recombinant mtDNA indicates that the exogenous and endogenous mtDNAs come into close enough proximity to convert their sequences.

**New insights for understanding the energy dependency in biological processes using mito-mice**

Mito-mice can be used as model systems for understanding in vivo energy-dependent checkpoints. Since most mice with the genetic disruption of nuclear DNA-encoded proteins functional for mitochondrial biogenesis showed embryonic lethal phenotypes, it is rather difficult to examine the significance of energy supplied from mitochondrial respiration in in vivo events, such as cell differentiation, maintenance, and regeneration. On the other hand, mito-mouse embryos with ΔmtDNA can escape from the lethal phenotypes, because these embryos do not express mitochondrial respiration defects, due to mitochondrial positive complementation. Using mito-mice, we have recently succeeded in showing experimental evidence for mtDNA-based male infertility, and showing that the mitochondrial respiration activity is essential for the mammalian spermatogenesis, especially for progression to the pachytene stages during meiosis. It has been reported that synopsis of homologous chromosomes in the mammalian spermatogenic meiotic process begins at the zygote stage through chromosome movement and attachment, and that several factors for synaptonemal complexes of homologous chromosomes, such as RAD51, DMC1, and cohesin, contain functional ATP binding domains. Actually, abnormal and incomplete attachment of homologous chromosomes occurred only in mito-mice with high load of ΔmtDNA. Our in vivo study using mito-mice directly demonstrates that there is an energy-dependent checkpoint in meiotic zygote stage during mammalian spermatogenesis, and that meiotic arrest due to pathogenic mtDNA-derived mitochondrial respiration defects results in male infertility. On the basis of these findings, we have proposed that some cases of human male infertility with unknown etiology might result from mitochondrial respiratory dysfunction.

Since the ΔmtDNA load differs in each cell, spermatogenic cells carrying a relatively lower proportion of ΔmtDNA can complete meiosis and...
transformation into sperms, but the number and motility are reduced.\textsuperscript{22) In comparison with disease model mice generated by the manipulation of the nuclear genome, mito-mice are not always suitable for the drug screening, because it is difficult to obtain a large population of mito-mice with the same \(\Delta\text{mtDNA}\) load. Sperm samples from mito-mice could be utilized for large-scale screening drugs designed to restore mitochondrial respiratory dysfunction, based on the recovery of decreased sperm motility. Moreover, drugs capable of improving mitochondrial respiratory dysfunction would be useful for treating not only mitochondrial diseases but also male infertility from mitochondrial causes.\textsuperscript{22) }

\textbf{Gene therapy of mito-mice by nuclear transplantation}

Considering the pathogeneses of mitochondrial diseases regulated by mitochondrial complementation, dilution of pathogenic mutated mtDNA population is one of the prenatal therapy strategies for the diseases. Effective procedures to obtain normal progeny from affected mothers would be either

\begin{figure}
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\caption{Pathological observations of mito-mouse testes.}
\begin{enumerate}
\item[A] Schematic presentation of mammalian spermatogenesis. Mammalian spermatogenesis occurs continuously with individual maturation of sperm and comprises the entire sequence of events by which spermatogonia are transformed into sperm, through meiotic division of spermatocytes. MCA indicates expression of male meiotic metaphase chromosome-associated acidic protein (meichroacidin).

\item[B] Distribution of spermatogenic cells in the state of meiotic processes. To visualize spermatocytes and spermatids, testis sections from B6 mouse, Mouse-11, Mouse-75, and Mouse-85 were stained with anti-MCA antiserum, because anti-MCA is specific for the cytoplasm of spermatocytes undergoing meiosis and early round spermatids, but not for cytoplasm of other cells in the testis (see Fig. 5A). Fewer cells stained positive for the anti-MCA decreased in Mouse-75 testis than in Mouse-11 or B6 mouse testes and the number of such cells further decreased in Mouse-85, indicating meiotic abnormalities in spermatogenesis. The Mouse-11, Mouse-75, and Mouse-85 possessed 11\%, 75\%, and 85\% \(\Delta\text{mtDNA}\) in their tails. Scale bar, 50 \(\mu\text{m}\).

\item[C] Homozygous chromosomes at the zygote stage in Mouse-82. Abnormal homozygous chromosomes were seen in spermatocyte nuclei at the zygote stage (see Fig. 5A). Single and double arrowheads indicate the region with self-attachment and the degenerated chromosome, respectively. The homozygous chromosomes in a right picture were avoided to form their synapsis due to the self-attachment. The Mouse-82 possessed 82\% \(\Delta\text{mtDNA}\) in the tail. Scale bar, 2 \(\mu\text{m}\).
\end{enumerate}
\end{figure}
nuclear transplantation from mito-mouse zygotes to enucleated normal zygotes, or cytoplasmic transplantation of mitochondria from normal to mito-mouse zygotes. However, the amount of mtDNA copies that could be introduced into mouse zygotes by cytoplasmic transplantation is relatively small. Nuclear transplantation inevitably co-introduces a small volume of cytoplasm and mitochondria, but the amount of mtDNA co-introduced with nucleus is 6% of all mtDNA in zygotes. Theoretically, nuclear transplantation would result in production of nuclear transplanted-zygotes with 6% mtDNA from mito-mouse zygotes, even when the mite-mouse zygotes possess 100% mtDNA. Since the maximum proportion of mtDNA in mito-mouse zygotes was 78%, nuclear transplantation therapy could hold the proportion of co-transplanted mtDNA at 5% or less. Therefore, nuclear transplantation was considered to be more effective than cytoplasmic transplantation for sufficient dilution of mtDNA in zygotes of mito-mice.

In nuclear transplantation therapy, we first estimated proportions of mtDNA in mito-mouse zygotes using polar bodies as biopsy samples. Then, from 39 zygotes possessing 35% mtDNA, karyoplasts were prepared and fused with enucleated normal zygotes. They were transferred to pseudo-pregnant females, and the resultant 11 mice at the weaned stage possessed 11% mtDNA in their tails. On the other hand, when 34 mite-mouse zygotes with 32% mtDNA were transferred without nuclear transplantation, obtained 9 mice possessed 66% mtDNA. They showed mitochondrial respiration deficiencies in various tissues and died due to renal failure at 218-277 days after birth, while nuclear-transplanted mite-mice showed no abnormalities throughout their lives.

In human cases, however, there are several biological problems to be resolved before applying gene therapy to zygotes. One is that in human cases maternal transmission of mtDNA with a common deletion is rare. When deletion mutant mtDNA is transmitted to oocytes, the 13 times longer gestation period in humans than in mice would result in accumulation of a sufficient proportion of deletion mutant mtDNA in nuclear-transplanted embryos to be lethal before birth. Therefore, application of nuclear transplantation would be restricted to patients with mitochondrial diseases, only when pathogenic mtDNAs in patients were inherited maternally, and did not possess significant replication advantages over wild-type mtDNA. For example, zygotes from patients with MELAS (mitochondrial myopathy, encephalopathy and lactic acidosis, and stroke-like episodes) and MERRF (myoclonus epilepsy associated with ragged-red fibers) caused by point mutations in mtDNAs, respectively, would be appropriate for applying nuclear transplantation to rescue their progeny. In these cases, a small proportion of wild-type mtDNAs was sufficient to suppress disease phenotypes. The other problem is that nuclear transplantation results in zygotes possessing the heteroplasmic state of mtDNA molecules from two mothers. At the present stage, we could not rule out the possibility that the heteroplasmic state itself or the resultant creation of mtDNA recombinants might induce mitochondrial abnormalities.

Recently, it was reported that the UK Human Fertilization and Embryology Authority (HFEA) would allow scientists at the University of Newcastle upon Tyne to transfer the nucleus of a human fertilized egg into an egg donated by a second woman. Although the study using mito-mice unambiguously showed that the nuclear transplantation technique could also be applied to human patients with mitochondrial diseases as a prenatal therapy, we would like to stress that we have to discuss not only biological safety, but also ethical points in the application of this therapy.

Future works
All genes encoded by human mtDNA are required for mitochondrial respiratory function, so that any pathogenic mutations in the mtDNA could induces mitochondrial respiration defects, leading to various clinical phenotypes associated with mitochondrial diseases. Disease phenotypes, however, are different among the types of the mutation. For example, mutations in tRNA genes are responsible for MELAS, MERRF, and cardiomyopathy, whereas ones in structural genes induce Leigh syndrome and Laber’s disease. Furthermore, nuclear genome backgrounds could involve in the difference of disease phenotypes, since most subunits for respiration complexes are encoded in...
nuclear genome. For addressing precise pathogeneses in mitochondrial diseases, we have to generate new mito-mice carrying various mutated mtDNAs responsible for human cases and various nuclear genome backgrounds. Recently, we have succeeded in generating new mito-mice carrying various mutated mtDNAs with a point mutation in COXI gene, and this mouse have showed lactic acidosis and growth retardation. In addition, we continue to generate mouse lines with mutated mtDNAs, so we will be able to report several mouse models for mtDNA-based diseases in near future.

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Profile

Kazuto Nakada was born in Tochigi, 1969. He received his Ph.D. from University of Tsukuba in 1999 under the supervision of Prof. Tamio Hirabayashi. After working at Prof. Hirabayashi’s laboratory as a Research Fellow at the Japan Society for the Promotion of Science (JSPS), he was promoted to Assistant Professor of University of Tsukuba in 2000, and he started mitochondrial studies at Prof. Jun-ichi Hayashi’s laboratory. For 3 years from 2001, he also worked as a Researcher at Precursory Research for Embryonic Science and Technology (PRESTO), Japan Science and Technology Agency. Since 2004, he is Associate Professor of Cell Biology at University of Tsukuba. He received Young Scientist Award of the Ministry of Education, Culture, Sports, Science and Technology-Japan in 2006 and JSPS Prize in 2008. His research interest focuses on biogenesis of mammalian mitochondria carrying mutated mitochondrial genome.