Molecular pathomechanisms and cell-type-specific disease phenotypes of MELAS caused by mutant mitochondrial tRNA<sub>Trp</sub>

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Abstract

Introduction: Numerous pathogenic mutations responsible for mitochondrial diseases have been identified in mitochondrial DNA (mtDNA)-encoded tRNA genes. In most cases, however, the detailed molecular pathomechanisms and cellular pathophysiology of these mtDNA mutations—how such genetic defects determine the variation and the severity of clinical symptoms in affected individuals—remain unclear. To investigate the molecular pathomechanisms and to realize in vitro recapitulation of mitochondrial diseases, intracellular mutant mtDNA proportions must always be considered.

Results: We found a disease-causative mutation, m.5541C>T heteroplasmy in MT-TW gene, in a patient exhibiting mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episodes (MELAS) with multiple organ involvement. We identified the intrinsic molecular pathomechanisms of m.5541C>T. This mutation firstly disturbed the translation machinery of mitochondrial tRNA<sub>Trp</sub> and induced mitochondrial respiratory dysfunction, followed by severely injured mitochondrial homeostasis. We also demonstrated cell-type-specific disease phenotypes using patient-derived induced pluripotent stem cells (iPSCs) carrying ~100 % mutant m.5541C>T. Significant loss of terminally differentiated iPSC-derived neurons, but not their stem/progenitor cells, was detected most likely due to serious mitochondrial dysfunction triggered by m.5541C>T; in contrast, m.5541C>T did not apparently affect skeletal muscle development.

Conclusions: Our iPSC-based disease models would be widely available for understanding the “definite” genotype-phenotype relationship of affected tissues and organs in various mitochondrial diseases caused by heteroplasmic mtDNA mutations, as well as for further drug discovery applications.

Keywords: Mitochondrial myopathy, encephalopathy, lactic acidosis, stroke-like episodes (MELAS), Mutant mitochondrial tRNA<sub>Trp</sub>, Mitochondrial respiratory dysfunction, Induced pluripotent stem cells (iPSCs), in vitro cellular disease models
Introduction
Mitochondrial DNA (mtDNA) has unique translation and transcription machinery and is associated with the maintenance of cellular homeostasis through intergenic crosstalk with nuclear DNA (nDNA). Numerous pathogenic mutations responsible for various mitochondrial diseases have been identified in mtDNA-encoded tRNA genes [1], and in most cases, such disease-causing heteroplasmic mutations (i.e., wild-type mtDNA and mutant mtDNA co-exist within a cell) exhibit their intrinsic pathogenic thresholds. Mutant mitochondrial tRNAs frequently induce various loss-of-function at a molecular level [2] including not only damaged mitochondrial protein synthesis, but also inhibited aminoacylation [3], tRNA molecular instability [4], altered tRNA processing [5], wobble-base modification deficiency [6, 7], or a combination of these. Therefore, the degree of accumulated mutant mitochondrial tRNAs within a cell is most likely to determine the trajectory of tissue- and organ-specific disease progression and phenotypic severity in affected individuals. Focusing on MT-TW gene, which encodes mitochondrial tRNA

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\text{\textbf{Pathomechanisms of m.5541C>T and demonstrated cell-type-specific disease phenotypes using patient-derived iPSCs carrying ~100 % mutant m.5541C>T. Our iPSC-based disease models would be widely available for understanding the definitive genotype-phenotype relationship of affected tissues and organs in various mitochondrial diseases caused by heteroplasmic mtDNA mutations, as well as for further drug discovery applications.}}
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Patients and methods
Patients
This study was approved by our Institutional Review Board and was stringently conducted in accordance with the ethical principles of the "Declaration of Helsinki". Patient biopsy was performed for diagnostic purposes only after we received written informed consent with permission to study patient-derived iPSCs. Note that 10 control subjects were also used in this study.

A partial family pedigree for this patient is shown (Fig. 1a). In this family, there was no clinical history of any neuromuscular disease. He had no growth and mental retardation until firstly presenting epileptic symptoms at age 10 years. At age 11 years, he developed weight loss, activity loss, easy fatigue, cognitive impairment, and acute heart failure. Radiographic and ultracardiographic images revealed hypertrophic cardiomyopathy at interventricular septum and left ventricular wall (Fig. 1b). Markedly increased serum lactate level (114.7 mg/dL; 3.0-17.0 mg/dL as normal), serum pyruvate level (3.86 mg/dL; 0.30–0.94 mg/dL as normal), and lactate/pyruvate ratio (29.7) were detected. At age 13 years, he developed headache, vomiting, visual disturbance, convulsion, and myoclonic status with unconsciousness. Brain MRI revealed multifocal hyper-intensity lesions at basal ganglia, cortex, and subcortical white matter of both cerebrum and cerebellum. A representative lesion showed decreased N-acetylaspartate level and increased lactate level (Fig. 1c). On this occasion, no significant abnormalities in serum lactate level (18.9 mg/dL), serum pyruvate level (0.99 mg/dL), or lactate/pyruvate ratio (19.0) were detected; however, cerebrospinal fluid lactate level (41.5 mg/dL), cerebrospinal fluid pyruvate level (1.40 mg/dL), and lactate/pyruvate ratio (29.6) were clearly high. Skeletal muscle histopathology revealed diffuse cytochrome c oxidase (COX) deficiency (less than 5 % population of COX-positive fibers) (Fig. 1d); however, no other typical pathological abnormalities such as ragged-red-fibers or strongly succinate dehydrogenase (SDH)-reactive blood vessels were observed. We diagnosed this patient as MELAS and started oral administration of l-arginine, dichloroacetate, and sodium pyruvate. He relapsed with stroke-like episodes twice in 4 years. At age 14 years, he developed multiple organ involvement...
Fig. 1 (See legend on next page.)
including acute pancreatitis, gastrointestinal malabsorption, renal tubular disturbance, and endocrine glucose intolerance. At age 15 years, he developed quadriplegia. Now, he keeps stable condition without serious trouble.

**Myoblast culture**

Patient-derived skeletal muscle tissues were minced, enzymatically digested, and centrifuged to collect myoblasts according to standard protocol. Patient-derived myoblasts were maintained in DMEM/F12 (Gibco) supplemented with 20 % FBS (Gibco), 100 units/mL penicillin (Gibco), 100 μg/mL streptomycin (Gibco) at 37 °C under humidified atmosphere of 5 % CO₂. During establishment of patient-derived primary myoblasts, 0.5 μg/mL MC210 (DS Pharm) as a mycoplasmaicidal reagent and 2.5 μg/mL fungizone (Gibco) as a fungicidal reagent were also added to culture medium.

Terminal differentiation of patient-derived myoblasts into myotubes was performed as follows: Briefly, patient-derived myoblasts were seeded at high cell density onto 6-well culture plates and were maintained at 37 °C under humidified atmosphere of 5 % CO₂. After 3 days in culture, culture medium was switched to myogenic differentiation medium (Cell Applications) supplemented with 100 units/mL penicillin (Gibco), 100 μg/mL streptomycin (Gibco), and patient-derived myoblasts were maintained at 37 °C under humidified atmosphere of 5 % CO₂ for 2 weeks.

**Analysis of mtDNA mutation**

Long PCR-based whole mtDNA sequencing for the patient was performed as described elsewhere [17] with modifications to eliminate any adverse results arising from pseudo-sequences in nuclear DNA: Briefly, extracted DNA as template (10 ng for iPSCs, 20 ng for myoblasts and skeletal muscle tissues, 100 ng for blood tissues) was amplified via mtDNA-specific long-range PCR and the following mtDNA-specific nested PCR with a thermal cycler (GeneAmp PCR System 9700; Applied Biosystems). The amplified mtDNA fragments were sequenced with a DNA analyzer (ABI PRISM 3130xl; Applied Biosystems). The obtained mtDNA sequence data from each patient was compared with the databases of “Human Mitochondrial Genome Database (MITOMAP; http://www.mitomap.org/MITOMAP/)” and “Human Mitochondrial Genome Polymorphism (mtSNP; http://mtsnp.tmig.or.jp/mtsnp/)” [18] to find any genetic variants. Mutation ratio of m.5541C>T in the patient was determined as follows: Extracted DNA (1 ng) was used as template for quantitative PCR with TaqMan Universal PCR Master Mix kit (Applied Biosystems) according to the manufacturer’s instructions. A sequence detection system (ABI PRISM 7900HT; Applied Biosystems) was used, and a calibration curve was created using several copy-number standards with plasmids containing the amplified mtDNA fragments (nucleotide position in mtDNA; 5205–5767) for either wild-type or mutant sequences.

**Protein and tRNA modeling**

*in silico* modeling of bovine heart CIV homodimer in fully oxidized state [19] was performed on the database of “RCSB Protein Data Bank (http://www.rcsb.org/pdb/),” and the graphics were arranged using RasMol software. Amino acid sequences of bovine and human mtDNA-encoded CIV subunits were referenced in the database of “NCBI Protein Database (http://www.ncbi.nlm.nih.gov/protein/)”.

*in silico* modeling of wild-type and mutant mitochondrial tRNA^{TP} were performed on the database of “Vienna RNA Servers (http://rna.tbi.univie.ac.at/),” and representative physicochemical parameters (e.g., minimum free energy and base-pair probability) were simultaneously calculated.

**Reverse transcription PCR (RT-PCR) and quantitative PCR**

Reverse transcription was performed with PrimeScript RT Master Mix kit (TaKaRa Bio) according to the manufacturer’s instructions. After reverse transcription of extracted total RNA, total cDNA (25 ng) was used as template for RT-PCR with a thermal cycler (GeneAmp PCR system 9700; Applied Biosystems). The amplified PCR products were resolved via electrophoresis through 2 % agarose gel, stained with GelGreen (Biotium), and detected with a UV transilluminator (GelDoc-It Imaging System; UVP).
Quantitative PCR for measurement of transgenes expression was performed with SYBR Green I PCR Master Mix kit (Roche) according to the manufacturer’s instructions. After reverse transcription of extracted total RNA, total cDNA (10 ng) was used as template for quantitative PCR with a real-time PCR system (LightCycler 480II; Roche). The averaged threshold cycle number for housekeeping genes were adopted for ΔΔC_T-based relative quantification.

**Analysis of mitochondrial enzymatic activity**

Enzymatic activities for individual mitochondrial respiratory chain complexes were analyzed as described elsewhere [20] with modifications: Cultured and
Mutant mitochondrial tRNA<sup>Trp</sup> strongly impairs neuronal maturation, but does not affect skeletal muscle development.

**Fig. 4**

**a** Experimental design used to identify patient-specific disease phenotypes triggered by m.5541C > T. 

**b** Representative images of myotubes after 2 weeks of differentiation; ACTA1 (red). Cell nuclei were co-stained with Hoechst 33342 (blue).

**c** Differentiation efficiency of myoblasts into myotubes. The number of ACTA1-positive myotubes was counted using a cytometer to estimate differentiation efficiency. Assays were performed using 3 experimental replicates for both control and the patient. Error bars indicate as the means with SD. Statistical significance was evaluated by unpaired, two-tailed t-test. NS: Not Significant.

**d** Representative images of iPSC-derived NSCs; Nestin (green).

**e** Expression of representative NSC marker genes in iPSC-derived NSCs.

**f** Representative images of CNS neurons after 2 weeks of differentiation; TUJ1 (red). Cell nuclei were co-stained with Hoechst 33342 (blue).

**g** Differentiation efficiency of NSCs into CNS neurons. The number of TUJ1-positive CNS neurons was counted using a cytometer to estimate differentiation efficiency. Assays were performed using 3 experimental replicates for both control and the patient. Error bars indicate as the means with SD. Statistical significance was evaluated by unpaired, two-tailed t-test. *

**h** Representative images of iPSC-derived NCCs; HNK1 (green).

**i** Expression of representative NCC marker genes in iPSC-derived NCCs.

**j** Representative images of PNS neurons after 2 weeks of differentiation; Peripherin (red). Cell nuclei were co-stained with Hoechst 33342 (blue).

**k** Differentiation efficiency of NCCs into PNS neurons. The number of Peripherin-positive PNS neurons was counted using a cytometer to estimate differentiation efficiency. Assays were performed using 3 experimental replicates for both control and the patient. Error bars indicate as the means with SD. Statistical significance was evaluated by unpaired, two-tailed t-test. **P < 0.05**
harvested patient-derived myoblasts (1 × 10^5 cells/assay) were applied for measurement. A spectrophotometer equipped with thermostated unit (U-2010; Hitachi) was used, and a base line calibration was done before each measurement.

For CI activity measurement, cells were added into reaction buffer [pH 7.4; 50 mM Tris–HCl, 250 mM sucrose, 1 mM EDTA, 10 μM decylubiquinone, 50 μM NADH, 5 μg/mL antimycin A, 2 mM potassium cyanide] and were incubated at 37 °C. CI activity was monitored by time-dependent absorbance alterations.

For CII activity measurement, cells were added into reaction buffer [pH 7.4; 50 mM potassium phosphate, 20 mM succinate, 50 μM 2.6-dichlorophenolindophenol, 50 μM decylubiquinone, 5 μg/mL rotenone, 5 μg/mL antimycin A, 2 mM potassium cyanide] and were incubated at 37 °C. CII activity was monitored by time-dependent absorbance alterations.

For CIII activity measurement, cells were added into reaction buffer [pH 7.4; 50 mM Tris–HCl, 250 mM sucrose, 1 mM EDTA, 50 μM cytochrome c, 50 μM decylubiquinol (reduced form of decylubiquinone), 2 mM potassium cyanide] and were incubated at 37 °C. CIII activity was monitored by time-dependent absorbance alterations.

For CIV activity measurement, cells were added into reaction buffer [pH 7.4; 10 mM potassium phosphate, 25 μM ferrocytochrome c (reduced form of cytochrome c)] and were incubated at 37 °C. CIV activity was monitored by time-dependent absorbance alterations.

For citrate synthase (CS) activity measurement, cells were added into reaction buffer [pH 8.0; 125 mM Tris–HCl, 300 μM acetyl-CoA, 100 μM N5,5'-dithiobis (2-nitrobenzoic acid), 500 μM oxaloacetate] and were incubated at 37 °C. CS activity was monitored by time-dependent absorbance alterations.

**Electrophoretic separation of mitochondrial proteins**

Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) [21] and blue native polyacrylamide gel electrophoresis (BN-PAGE) [22] were performed as described elsewhere with modifications, respectively: Cultured and harvested patient-derived myoblasts were resuspended in isolation buffer [pH 7.4; 210 mM mannitol, 70 mM sucrose, 1 mM EGTA, 5 mM HEPES] and were homogenated on ice. Cell lysates were centrifuged to isolate mitochondrial proteins. Obtained mitochondrial proteins were quantified by Bradford assay, and a calibration curve was created using several known concentrations of BSA.

For SDS-PAGE, isolated mitochondrial proteins (100 μg) were solubilized with 0.5 % SDS containing 50 mM dithiothreitol at 70 °C for 10 min. Electrophoresis was performed on 4 –12 % NuPAGE polyacrylamide gel (Invitrogen) at room temperature under 200 V constant.

For BN-PAGE, isolated mitochondrial proteins (100 μg) were solubilized with either 0.5 % n-dodecyl-β-d-maltoside (individual complexes detection) or 1 % digitonin (supercomplexes detection) on ice for 30 min. Insoluble proteins were removed by centrifugation. Electrophoresis was performed on 3–12 % NativePAGE polyacrylamide gel (Invitrogen) at 4 °C under 150 V constant for 30 min, then resumbed at 4 °C under 250 V constant.

**Western blot for immunodetection of mitochondrial proteins**

Electrophoresed gels were blotted onto polyvinylidene fluoride (PVDF) membranes using an iBlot transfer system (Invitrogen) according to the manufacturer’s instructions. Blotted PVDF membranes were blocked at room temperature for 30 min. Primary antibody probing was performed at room temperature for 90 min. Secondary antibody probing was performed with chromogenic antibody detection kit (WesternBreeze; Invitrogen) according to the manufacturer’s instructions. Primary antibodies used for SDS-PAGE were as follows: 0.5 μg/mL anti-porin (Molecular Probes), 2.5 μg/mL anti-MT-CO1 (Molecular Probes), 2.5 μg/mL anti-MT-CO2 (Molecular Probes), 2.5 μg/mL anti-COX4 (Molecular Probes), 2.5 μg/mL anti-COX5B (Molecular Probes). Primary antibodies used for BN-PAGE were as follows: 0.5 μg/mL anti-NDUFA9 for CI (Molecular Probes), 0.5 μg/mL anti-SDHA for CII (Molecular Probes), 0.5 μg/mL anti-UQCRC2 for CIII (Molecular Probes), 2.5 μg/mL anti-MT-CO1 for CIV (Molecular Probes), 0.5 μg/mL anti-ATP5B for CV (Molecular Probes).

**Cytochemical staining**

Patient-derived myoblasts were seeded onto 4-well culture slides and were maintained at 37 °C under humidified atmosphere of 5 % CO2. After 3 days in culture, cytochemical staining was performed as follows:

For cytochemical COX staining, cells were stained with reaction buffer [pH 5.5; 100 mM sodium acetate, 0.1 % MnCl2, 0.001 % H2O2, 10 mM diaminobenzidine] at 37 °C for 1 h, followed by subsequent incubation with 1 % CuSO4 at 37 °C for 5 min. Cell nuclei were co-stained with hematoxylin. Stained cells were rinsed, fixed, and dehydrated according to standard histological protocol. Samples were sealed with cover glass and were observed under an optical microscope (BX50 System; Olympus).

For cytochemical SDH staining, cells were stained with reaction buffer [pH 7.4; 50 mM succinate, 1 mM nitrotetrazolium blue] at 37 °C for 1 h. Cell nuclei were co-stained with hematoxylin. Stained cells were rinsed, fixed, and dehydrated according to standard histological protocol.
protocol. Samples were sealed with cover glass and were observed under an optical microscope (BX50 System; Olympus).

**Immunocytochemical detection of CIV structural subunits**

Patient-derived myoblasts were seeded onto 4-well culture slides and were maintained at 37 °C under humidified atmosphere of 5 % CO₂. After 3 days in culture, cells were fixed, permeabilized, and blocked according to standard immunocytochemical protocol. Primary antibody probing was performed at room temperature for 2 h. Secondary antibody probing was performed with 2.5 μg/mL Alexa Fluor 568 (Molecular Probes) at room temperature for 1 h. Mitochondria were co-stained with 0.25 μg/mL MitoTracker Green (Molecular Probes). Stained cells were observed under a fluorescent microscope (IX71 System; Olympus). Primary antibodies used were as follows: 2.5 μg/mL anti-MT-CO1 (Molecular Probes), 2.5 μg/mL anti-COX4 (Molecular Probes).

**Analysis of ATP level**

Cultured and harvested patient-derived myoblasts (100 cells/assay) were applied for measurement. ATP amount was monitored with rLuciferase/Luciferin chemiluminescence-based ATP detection kit (Promega) according to the manufacturer’s instructions. A chemiluminescent multi-well plate reader (Centro LB 960; Berthold Technologies) was used, and a calibration curve was created using several known concentrations of ATP.

**Analysis of oxidative stress level and membrane potential level**

Patient-derived myoblasts were seeded onto 96-well culture plate and were maintained at 37 °C under humidified atmosphere of 5 % CO₂. After 3 days in culture, cells were stained at 37 °C for 1 h. Stained cells were rinsed and were measured on a fluorescent multi-well plate reader (ARVO SX; Perkin Elmer); first at excitation/emission of 545/595 nm (red fluorescence) and then sequentially at excitation/emission of 485/535 nm (green fluorescence). Fluorescent dyes used were as follows: 0.25 μg/mL MitoTracker Green (Molecular Probes), 0.25 μg/mL MitoSOX Red (Molecular Probes), 0.25 μg/mL JC-1 (Molecular Probes).

**Generation of patient-derived iPSCs with episomal vector**

Patient-derived iPSCs were generated using episomal vectors as described previously [23] with modifications: Briefly, each 1 μg of episomal plasmid vectors (Plasmid #27077, #27078, #27080; Addgene) were electroporated into patient-derived myoblasts (5 × 10⁵ cells) with an electroporator (Neon; Invitrogen). Transformed patient-derived myoblasts (1 × 10⁵ cells) were reseeded onto mouse embryonic fibroblasts (MEF; ReproCELL) 4 days after electroporation. The next day, culture medium was replaced with primate ESC culture medium (ReproCELL) supplemented with 10 ng/mL bFGF (ReproCELL), 100 units/mL penicillin (Gibco), 100 μg/mL streptomycin (Gibco), and transformed patient-derived myoblasts were maintained at 37 °C under humidified atmosphere of 5 % CO₂. Emergent colonies with ESC-like morphology were manually picked up to establish patient-derived iPSCs, and these iPSCs were expanded either on MEF-seeded dishes in primate ESC culture medium or on Geltrex (Gibco)-coated dishes in mTeSR1 medium (STEMCELL Technologies) supplemented with 100 units/mL penicillin (Gibco), 100 μg/mL streptomycin (Gibco) for long-term maintenance.

**Characterization of patient-derived iPSCs**

Characterization of patient-derived iPSCs via detection of pluripotency markers was performed as follows: Briefly, cultured and harvested patient-derived iPSCs were transferred onto MEF-seeded 6-well culture plates and were maintained in primate ESC culture medium at 37 °C under humidified atmosphere of 5 % CO₂. After 3 days in culture, patient-derived iPSCs were characterized by immunocytochemical staining. Fluorophore-conjugated primary antibodies used were as follows: 5 μg/mL Cy3-conjugated anti-OCT4 (Millipore), 5 μg/mL Cy3-conjugated anti-NANOG (Millipore), 5 μg/mL AlexaFluor 488-conjugated anti-TRA-1-60 (Millipore), 5 μg/mL AlexaFluor 488-conjugated anti-TRA-1-81 (Millipore).

_in vitro_ spontaneous differentiation of patient-derived iPSCs into EB-mediated three germ layers was performed as follows: Briefly, cultured and harvested patient-derived iPSCs were transferred onto ultra-low-adherent culture dishes (HydroCell; CellSeed) and were maintained in primate ESC culture medium without bFGF at 37 °C under humidified atmosphere of 5 % CO₂. After 7 days in floating culture, emergent EBs were transferred onto gelatin-coated 6-well culture plates and were maintained in primate ESC culture medium without bFGF at 37 °C under humidified atmosphere of 5 % CO₂. After 14 additional days in adherent culture, spontaneously differentiated cells were characterized by immunocytochemical staining. Primary antibodies used were as follows: 5 μg/mL anti-TUJ1 for ectoderm (Abcam), 5 μg/mL anti-αSMA for mesoderm (Abcam), 5 μg/mL anti-AFP for endoderm (Abcam). Secondary antibody used was 2.5 μg/mL Alexa Fluor 568 (Molecular Probes).

**Directed differentiation of iPSCs into neural stem cells (NSCs)**

Directed differentiation of patient-derived iPSCs into NSCs was performed according to a previous report.
[24] with modifications: Briefly, patient-derived iPSCs were seeded onto Geltrex-coated dishes and were maintained in mTeSR1 medium at 37 °C under humidified atmosphere of 5 % CO₂. After 3 days in adherent culture, culture medium was switched to NSC induction medium [1:1 mixture of DMEM/F12 (Gibco) and Neurobasal medium (Gibco) supplemented with 1 × N2 (Gibco), 1 × B27 minus vitamin A (Gibco), 1 × GlutaMAX (Gibco), 100 units/mL penicillin (Gibco), 100 μg/mL streptomycin (Gibco), 10 μM SB431542 (Wako), 100 nM LDN193189 (Wako), 20 ng/mL EGF (Peprotech), 20 ng/mL bFGF (Peprotech), and patient-derived iPSCs were maintained at 37 °C under humidified atmosphere of 5 % CO₂. Emergent NSCs were expanded in NSC induction medium and were characterized by immunocytochemical staining. Fluorophore-conjugated primary antibody used was 5 μg/mL AlexaFluor 488-conjugated anti-Nestin (Millipore).

Directed differentiation of iPSCs into neural crest cells (NCCs)

Directed differentiation of patient-derived iPSCs into NCCs was performed according to a previous report [25] with modifications: Briefly, patient-derived iPSCs were seeded onto Geltrex-coated dishes and were maintained in mTeSR1 medium at 37 °C under humidified atmosphere of 5 % CO₂. After 3 days in adherent culture, culture medium was switched to NCC induction medium [mTeSR1 medium (STEMCELL Technologies) supplemented with 100 units/mL penicillin (Gibco), 100 μg/mL streptomycin (Gibco), 2 μM (2Z,3E)-6-bromoindirubin-3’-oxime (Wako), 20 μM SB431542 (Wako)], and patient-derived iPSCs were maintained at 37 °C under humidified atmosphere of 5 % CO₂. Emergent NCCs were expanded in NCC induction medium and were characterized by immunocytochemical staining. Fluorophore-conjugated primary antibody used was 5 μg/mL FITC-conjugated anti-HNK1 (Miltenyi Biotec).

Terminal differentiation of NSCs and NCCs into neurons

Terminal differentiation of patient-derived NSCs and NCCs into neurons was performed as follows: Briefly, patient-derived NSCs and NCCs were seeded at high cell density onto Geltrex-coated 6-well culture plates and were maintained at 37 °C under humidified atmosphere of 5 % CO₂. After 3 days in culture, culture medium was switched to neuron induction medium [Neurobasal medium (Gibco) supplemented with 1 × N2 (Gibco), 1 × B27 minus vitamin A (Gibco), 1 × GlutaMAX (Gibco), 100 units/mL penicillin (Gibco), 100 μg/mL streptomycin (Gibco), 10 ng/mL BDNF (Peprotech), 10 ng/mL GDNF (Peprotech), 10 ng/mL NGF (Peprotech), 500 μM dbcAMP (Sigma), 200 μM ascorbic acid (Wako)], and patient-derived NSCs and NCCs were maintained at 37 °C under humidified atmosphere of 5 % CO₂ for more than 2 weeks. For immunocytochemical detection of emergent NSC-derived neurons, 5 μg/mL anti-TUJ1 (Abcam) and 2.5 μg/mL Alexa Fluor 568 (Molecular Probes) were used. For immunocytochemical detection of emergent NCC-derived neurons, 5 μg/mL PE-conjugated anti-Peripherin (Santa Cruz) was used.

Results

Molecular pathomechanisms of mutant mitochondrial tRNA_{TnP}

We found a patient who was clinically diagnosed as MELAS with multiple organ involvement including hypertrophic cardiomyopathy, acute pancreatitis, gastrointestinal malabsorption, renal tubular disturbance, and endocrine glucose intolerance (Fig. 1a-d). We identified a disease-causative mutation, m.5541C > T heteroplasmy in MT-TW gene, in this patient (Fig. 1e and Additional file 1: Table S1). Skeletal muscle tissues and the established myoblasts showed quite high mutant proportions (~100 %), whereas blood tissues showed relatively low mutant proportions (~50 %). The same m.5541C > T heteroplasmy was also observed in blood tissues from this patient’s asymptomatic mother at quite low mutation levels (~10 %). The mutated position in the anticodon-stem of MT-TW gene was evolutionarily conserved through most parts of primates and typical eukaryotes (Fig. 1f). We also performed in silico calculation of mitochondrial tRNA_{TnP} stability for both wild-type and m.5541C > T mutant (Additional file 1: Figure S1). Mutant mitochondrial tRNA_{TnP} was destabilized by m.5541C > T and was probably existed more physicochemically stable but biochemically inappropriate conformation. Our findings suggest that m.5541C > T presumably induces defects in mitochondrial tRNA_{TnP}-associated translation machinery.

Although m.5541C > T was previously reported and predicted as "definitely pathogenic" [26], and our patient was the second case to show this mutation, the detailed molecular pathomechanisms of m.5541C > T —how this mutation influences mitochondrial pathophysiology, which is closely related to the variation and the severity of clinical symptoms in affected individuals— remain unclear. We comprehensively evaluated mitochondrial function using patient-derived myoblasts carrying quasi-homoplasmic m.5541C > T [i.e., ~100 % mutant mtDNA exists within a cell]. On mitochondrial respiratory chain complexes, severely decreased CIV activity and moderately decreased CI activity were both detected in the patient, whereas the other respiratory chain complexes showed within normal ranges (Fig. 2a,b). CIV holoenzyme and CIV-containing respiratory supramolecular architectures were also diminished in the patient, whereas...
the other respiratory chain complexes showed almost normal levels with the exception of moderately decreased CI holoenzyme amount (Fig. 2c,d). Such CIV holoenzyme deficits in the patient were most likely because of decreased mRNA and protein expression levels of mtDNA-encoded CIV structural subunits (Fig. 2e-g). On mitochondrial physiology, widespread dysfunction such as decreased ATP level, increased oxidative stress level, and damaged membrane potential level were all observed in the patient (Fig. 2h-j). We also performed in silico prediction on mitochondrial tryptophan contents and their locations in each mtDNA-encoded CIV structural subunit (Additional file 1: Figure S2). Amino acid sequences in all mtDNA-encoded CIV structural subunits showed high homology between bovine and human, and bovine mitochondrial tryptophan residues were predominantly located in α-helix and β-sheet domains essential for the maintenance of CIV structural subunit conformations. Some human mitochondrial tryptophan residues in mtDNA-encoded CIV structural subunits were also located at the boundary between CIV structural subunits necessary for CIV holoenzyme assembly. Our results clearly indicate that m.5541C > T primarily induces the aberrant steady-state of mitochondrial respiratory chain complexes, followed by severely injured mitochondrial homeostasis.

Cell-type-specific disease phenotypes of MELAS using patient-derived iPSCs carrying all mutant mitochondrial tRNA<sup>Trp</sup>

We next generated each 3 lines of integration-free disease-relevant iPSCs derived from myoblasts of both control and the patient carrying quasi-homoplasmic m.5541C > T as in vitro disease models. No apparent differences in embryonic stem cell (ESC)-like pluripotent characteristics were confirmed between iPSCs derived from control and the patient (Fig. 3 and Additional file 1: Figure S3). To elucidate patient-specific cellular disease phenotypes triggered by quasi-homoplasmic m.5541C > T, we used myotubes and iPSC-derived neurons of both control and the patient (Fig. 4a). In myogenic lineage, no significant differences in in vitro differentiation propensity into ACTA1-positive myotubes were observed between control and the patient (Fig. 4b,c). This phenomenon indicates that m.5541C > T seems not to affect skeletal muscle development in the patient regardless of serious mitochondrial dysfunction (see also Figs. 1d and 2). We differentiated patient-derived iPSCs into central nervous system (CNS) lineage. Efficient differentiation into nestin-positive NSCs (>95 % conversion) was observed in both control and the patient (Fig. 4d,e and Additional file 1: Figure S4a); however, the number of TUJ1-positive mature CNS neurons was markedly decreased only in the patient (Fig. 4f,g and Additional file 1: Figure S4b), and most parts of patient-derived differentiating NSCs finally died during long-term terminal differentiation (data not shown). We also differentiated patient-derived iPSCs into peripheral nervous system (PNS) lineage. A trend quite similar to CNS lineage, stable differentiation into HNK1-positive NCCs (>95 % conversion) was confirmed in both control and the patient (Fig. 4h,i and Additional file 1: Figure S4c); however, significant decrease of peripherin-positive mature PNS neurons was detected only in the patient (Fig. 4j,k and Additional file 1: Figure S4d), and patient-derived differentiating NCCs no longer survived during extended periods of neuronal maturation (data not shown). Several recent studies have demonstrated that mitochondria are gradually rejuvenated to an ESC-like "quiescent state" during cellular reprogramming [27–30]. Our results also suggest that stem/progenitor cells of both CNS and PNS lineages are minimally influenced by m.5541C > T, most likely because these cell types may possess a less active mitochondrial respiration state similar to ESCs and iPSCs. Therefore, we conclude that the molecular pathogenicity of m.5541C > T is strongly visible in terminally differentiated post-mitotic neurons, but not their stem/progenitor cells, which is probably associated with the degree of mitochondrial maturation during cellular lineage-commitment process.

Discussion

The molecular pathomechanisms of m.5541C > T can be summarized as follows: This mutation firstly loses the appropriate base pair interaction, from Watson-Crick to T-G mismatching, in the anticodon-stem of MT-<i>TW</i> gene and induces defects in mitochondrial tRNA<sup>Trp</sup>-associated translation machinery most likely due to inadequate anticodon recognition of mitochondrial tryptophan by its altered conformation. Mutant mitochondrial tRNA<sup>Trp</sup> disturbs the synthesis of mtDNA-encoded respiratory chain complexes subunits; in this case, markedly decreased amounts of mtDNA-encoded CIV subunits predominantly inhibit CIV holoenzyme formation at each assembly process [31]. In fact, some patients carrying this mutation or other reported pathogenic mutations in the anticodon-stem of MT-<i>TW</i> gene also present severe COX deficiency [8, 26, 32]. Therefore, our findings clearly demonstrate why mutant mitochondrial tRNA<sup>Trp</sup> is able to cause severe COX deficiency as one of common clinical phenotypes. Induced mitochondrial respiratory dysfunction triggered by loss of CIV holoenzyme severely impairs mitochondrial biogenesis and bioenergetics such as decreased ATP level, increased oxidative stress level, and damaged membrane potential level. Increased oxidative stress level may promote the accumulation of oxidative damages to other mitochondrial enzymes, substrates, lipids, and mtDNA, all of which lead to premature cell senescence. Damaged membrane potential level may also
accelerate the leakage of cytochrome c molecules in mitochondrial electron transport system, which induces apoptotic cell death. Thus, m.5541C>T causes widespread mitochondrial dysfunction, which is closely related to cell-type-specific physiological impairment in various post-mitotic tissues and organs in this patient.

This case is consistent with MELAS with multiple organ involvement characterized by its various clinical symptoms; however, we did not find any typical abnormalities in patient-derived skeletal muscle tissues other than diffuse COX deficiency. Generally, 80-90 % of MELAS patients those carrying mutant mtDNAs (e.g., m.3243A>G in MT-TL1 gene) exhibit ragged-red-fibers and/or strongly SDH-reactive blood vessels in skeletal muscle tissues, which indicate morphological, numerical, and/or physiological abnormalities in mitochondria. These findings suggest that quasi-homoplasmic m.5541C>T may cause rare pathological signatures in skeletal muscle development, presumably because this patient exhibits no significant differences in in vitro differentiation propensity into terminally differentiated myotubes as compared with control regardless of serious mitochondrial dysfunction.

Recently, several groups have also reported iPSC-based disease models for other heteroplasmic mutant mtDNAs [33-37]. As the common perceptions, intracellular mutant mtDNA proportions must always be considered for actual in vitro recapitulation of mitochondrial diseases at each cellular fate-determining process such as reprogramming, self-renewal, or differentiation. From this viewpoint, we generated integration-free patient-derived iPSCs carrying ~100 % mutant mtDNA and used them as in vitro cellular disease models to investigate the definitive genotype-phenotype relationship. We also demonstrated that terminally differentiated iPSC-derived CNS and PNS neurons, but not their stem/progenitor cells, are strongly influenced by m.5541C>T, most likely because the molecular pathogenic severity of mutant mitochondrial tRNA\textsuperscript{TTP} may be determined by the degree of physiological and morphological maturation in mitochondria. Although our presenting results do not completely elucidate the relationship between in vitro cellular disease phenotypes and in vivo clinical symptoms of this patient, our approach would be widely available for understanding bona fide molecular pathomechanisms and cellular pathophysiology of affected tissues and organs in patients carrying heteroplasmic mtDNA mutations, as well as for further drug discovery applications.

Conclusions

Throughout this study, we identified the "definite" molecular pathomechanisms of m.5541C>T and demonstrated cell-type-specific in vitro disease phenotypes triggered by mutant mitochondrial tRNA\textsuperscript{TTP} using integration-free disease-relevant iPSCs derived from myoblasts of the patient. Our iPSC-based strategy therefore holds enormous promise for the development of evidence-based, personalized diagnostics and therapeutics to patients exhibiting various mitochondrial diseases.

Additional file

**Additional file 1: Supplementary Figures and Tables.** (DOC 4252 kb)

**Figure S1.** Comparison of mitochondrial tRNATrp stability between wild-type and m.5541C>T mutant (related to Fig. 1). **Figure S2.** Protein modeling and amino acid sequences of each mtDNA-encoded CIV subunit (related to Fig. 2). **Figure S3.** Generation of disease-relevant iPSCs carrying all mutant mitochondrial tRNATrp (related to Fig. 3).

**Figure S4.** Mutant mitochondrial tRNATrp strongly impairs neuronal maturation (related to Fig. 4). **Table S1.** mtDNA sequence variants in this patient. **Table S2.** Primer list. **Table S3.** TaqMan probe list.

Competing interests

The authors declare that they have no competing interests.

Authors' contribution

HH conceived the study, designed and performed experiments. AK, HK, and IN collected the patient's clinical data. HH and YG analyzed and interpreted data, wrote the manuscript. All authors read and approved the final manuscript.

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References

1. Taylor RW, Turnbull DM (2005) Mitochondrial DNA mutations in human disease. Nat Rev Genet 6:389–402
2. Suzuki T, Nagao A, Suzuki T (2011) Human mitochondrial tRNAs: biogenesis, function, structural aspects, and diseases. Annu Rev Genet 45:299–329
3. Enriquez JA, Chomyn A, Attardi G (1995) MtDNA mutation in MERRF syndrome causes defective aminocylaization of tRNAGlu and premature translation termination. Nat Genet 10:47–55
4. Hao H, Moraes CT (1997) A disease-associated G5703A mutation in human mitochondrial DNA causes a conformational change and a marked decrease in steady-state levels of mitochondrial tRNA\textsuperscript{Glu}. Mol Cell Biol 17:6831–6837
5. Kaufmann F, Koga Y, Shanske S, Hirano M, DiMauro S, King MP (1996) Mitochondrial DNA and RNA processing in MELAS. Ann Neurol 40:172–180
6. Kirino Y, Yasukawa T, Ohta S, Akira S, Ishihara K, Watanabe K (2004) Codon-specific translational defect caused by a wobble modification deficiency in mutant tRNA from a human mitochondrial disease. Proc Natl Acad Sci U S A 101:15070–15075

7. Kirino Y, Goto Y, Campos Y, Arezes J, Suzuki T (2005) Specific correlation between the wobble modification deficiency in mutant tRNAs and the clinical features of a human mitochondrial disease. Proc Natl Acad Sci U S A 102:7127–7132

8. Nelson J, Hanna MG, Alsanjari N, Scaravilli F, Morgan-Hughes JA, Harding AE (1995) A new mitochondrial DNA mutation associated with progressive dementia and chorea: a clinical, pathological, and molecular genetic study. Ann Neurol 37:400–403

9. Santorelli FM, Tanji K, Sano M, Shanske S, El-Shawaw M, Kranz-Eble P (1997) Maternally inherited encephalopathy associated with a single-base insertion in the mitochondrial tRNA<sub>Gly</sub> gene. Ann Neurol 42:256–260

10. Silvestri G, Rana M, DMuzio A, Uncini A, Tonal P, Servidei S (1998) A late-onset mitochondrial myopathy is associated with a novel mitochondrial DNA (mtDNA) point mutation in the tRNA<sub>Glu</sub> gene. Neuromuscul Disord 8:291–295

11. Maniura-Weber K, Taylor RW, Johnson MA, Chrzansowa-Lightowlers Z, Morris AA, Charlton CP (2004) A novel point mutation in the mitochondrial tRNA<sub>Glu</sub> gene produces a neurogastrointestinal syndrome. Eur J Hum Genet 12:212–218

12. Saccozzi S, Salvatori L, Nishigaki Y, Walker WF, Hernandez-Rosa E, Trevison E (2008) A functionally dominant mitochondrial mutation in the mtDNA. Mol Genet Metab 94:103–108

13. Mikoaar-Beal E, Chamlia K, Kammoun F, Kammoun T, Aloulou H, Hachicha M (2009) Two new mutations in the MT-W gene leading to the disruption of the secondary structure of the tRNA<sub>Glu</sub> in patients with Leigh syndrome. Mol Genet Metab 97:179–184

14. Sproule DM, Kaufmann P (2008) Mitochondrial encephalopathy, lactic acidosis, and stroke-like episodes: basic concepts, clinical phenotype, and therapeutic management of MELAS syndrome. Ann NY Acad Sci 1142:133–158

15. Takahashi K, Tanabe K, Ohnuki M, Narita M, Ichisaka T, Tomoda K (2007) Induction of pluripotent stem cells from adult human fibroblasts by defined factors. Cell 131:861–872

16. Yu J, Vodyanik MA, Smuga-Otto K, Antosievicz-Bourget J, Frané JL, Tian S (2007) Induced pluripotent stem cell lines derived from human somatic cells. Science 318:1917–1920

17. Akanuma J, Muraki K, Komaki H et al (2000) Two pathogenic point mutations exist in the authentic mitochondrial genome, not in the nuclear pseudogene. J Hum Genet 45:337–341

18. Tanaka M, Takeyasu T, Fuku N, Li-Jun G, Kurata M (2004) Mitochondrial genome single nucleotide polymorphisms and their phenotypes in the Japanese. Ann NY Acad Sci 1011:7–20

19. Tsukihara T, Aoyama H, Yamashita E, Tomizaki T, Yamaguchi H, Shinzawa-Itoh K (1996) The whole structure of the 13-subunit oxidized cytochrome c oxidase. Science 272:1136–1144

20. Trounce IA, Kim YL, Jun AS, Wallace DC (1996) Assessment of mitochondrial oxidative phosphorylation in patient muscle biopsies, lymphoblasts, and transmigrational cell lines. Methods Enzymol 264:484–500

21. Schägger H (2006) Tricine-SDS-PAGE. Nat Protocol 1:16

22. Wittig I, Braun HP, Schägger H (2006) Blue native PAGE. Nat Protocol 1:16–22

23. Witting I, Braun HP, Schägger H (2006) Blue native PAGE. Nat Protocol 1:418–428

24. Chambers SM, Fasano CA, Papapetrou EP, Tomishima M, Sadelain M, Studer L (2009) Highly efficient neural conversion of human ES and iPS cells by dual inhibition of SMAD signaling. Nat Biotechnol 27:275–280

25. Menendez L, Yatskievych TA, Antin PB, Dalton S (2011) Wnt signaling and a dual inhibition of SMAD signaling. Nat Biotechnol 27:275–280

26. Blakely EL, Yahr JM, Alston CL, Craig K, Poulton J, Brierley C (2013) Pathogenic mitochondrial tRNA point mutations: nine novel mutations affirm their importance as a cause of mitochondrial disease. Hum Mutat 34:1260–1268

27. Armstrong L, Tiltner K, Saretzki G et al (2010) Human induced pluripotent stem cell lines show stress defense mechanisms and mitochondrial regulation similar to those of human embryonic stem cells. Stem Cells 28:661–673

28. Prigione A, Fauler B, Luz R, Lehrach H, Adjaye J (2010) The senescence-related mitochondrial/oxidative stress pathway is repressed in human induced pluripotent stem cells. Stem Cells 28:721–733

29. Holmes CD, Nelson TJ, Martinez-Fernandez A, Arell O, Lindor JZ, Dzeja PP (2011) Somatic oxidative bioenergetics transitions into pluripotency-dependent glycolysis to facilitate nuclear reprogramming. Cell Metab 14:264–271

30. Holmes CD, Dzeja PP, Nelson TJ, Terzic A (2012) Metabolic plasticity in stem cell homeostasis and differentiation. Cell Stem Cell 11:596–606

31. Shoubridge EA (2001) Cytochrome c oxidase deficiency. Ann J Med Genet 106:46–52

32. Silvestri G, Mongini T, Odoardi F, Modoni A, De Rosa G, Doriguzzi C (2000) A new mtDNA mutation associated with a progressive encephalopathy and cytochrome c oxidase deficiency. Neurology 54:1693–1696

33. Fujikura J, Nakao K, Sone M, Naguchi M, Mori E, Naito M (2012) Induced pluripotent stem cells generated from diabetic patients with mitochondrial DNA A3243G mutation. Diabetologia 55:1689–1698

34. Cherry AB, Gagne KE, McLoughlin EM, Bacci A, Gorman B, Hartung O (2013) Induced pluripotent stem cells with a mitochondrial DNA deletion. Stem Cells 31:1287–1297

35. Holmes CD, Martinez-Fernandez A, Perales-Clemente E, Li X, McDonald A, Ogliesbee D (2013) Disease-causing mitochondrial heteroplasmy segregated within induced pluripotent stem cell clones derived from a patient with MELAS. Stem Cells 31:1298–1308

36. Hämäläinen RH, Manninen T, Koivukul H, Kíslín M, Otonkoski T, Suomalainen A (2013) Tissue- and cell-type-specific manifestations of heteroplasmic mtDNA 3243A→G mutation in human induced pluripotent stem cell-derived disease model. Proc Natl Acad Sci U S A 110:3622–3630

37. Kodaira M, Hatakeyama H, Yuasa S, Seki T, Egashira T, Tohyama S (2015) Impaired respiratory function in MELAS-induced pluripotent stem cells with high heteroplasmacy levels. FEBS Open Bio 5:219–225

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