Homoplasmic mitochondrial tRNA<sub>Pro</sub> mutation causing exercise-induced muscle swelling and fatigue

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Abstract

Objective
To demonstrate the causal role in disease of the MT-TP m.15992A>T mutation observed in patients from 5 independent families.

Methods
Lactate measurement, muscle histology, and mitochondrial activities in patients; PCR-based analyses of the size, amount, and sequence of muscle mitochondrial DNA (mtDNA) and proportion of the mutation; respiration, mitochondrial activities, proteins, translation, transfer RNA (tRNA) levels, and base modification state in skin fibroblasts and cybrids; and reactive oxygen species production, proliferation in the absence of glucose, and plasma membrane potential in cybrids.

Results
All patients presented with severe exercise intolerance and hyperlactatemia. They were associated with prominent exercise-induced muscle swelling, conspicuous in masseter muscles (2 families), and/or with congenital cataract (2 families). MRI confirmed exercise-induced muscle edema. Muscle disclosed severe combined respiratory defect. Muscle mtDNA had normal size and amount. Its sequence was almost identical in all patients, defining the haplotype as J1c10, and sharing 31 variants, only 1 of which, MT-TP m.15992A>T, was likely pathogenic. The mutation was homoplasmic in all tissues and family members. Fibroblasts and cybrids with homoplasmic mutation had defective respiration, low complex III activity, and decreased tRNA<sub>Pro</sub> amount. Their respiratory complexes amount and tRNA<sub>Pro</sub> aminoacylation appeared normal. Low proliferation in the absence of glucose demonstrated the relevance of the defects on cybrid biology while abnormal loss of cell volume when faced to plasma membrane depolarization provided a link to the muscle edema observed in patients.

Conclusions
The homoplasmic MT-TP m.15992A>T mutation in the J1c10 haplotype causes exercise-induced muscle swelling and fatigue.
Mitochondrial diseases, due to oxidative phosphorylation defects, represent an extraordinary diagnostic challenge because of their diversity. The hundreds genes involved, located on either the mitochondrial or the nuclear DNA, or the frequent heteroplasmy of mitochondrial DNA (mtDNA) mutations, i.e., coexistence of wild-type and mutant mtDNA molecules, likely explain part of that diversity. In addition, deleterious homoplastic mtDNA mutations, i.e., mutations affecting all mtDNA molecules, cause diseases with incomplete penetrance and tissue-specific expression that disclose the complexity of nuclear-mitochondrial interaction. This was the case with mutations causing Leber hereditary optic neuropathy or deafness. It was particularly striking for the homoplasmic MT-TI m.4300A>G mutation whose expression pattern ranged from infantile cardiac failure with severe cardiac mitochondrial defect to apparent health. Initial identification of these deleterious homoplasmic mutations thus required either their recurrence with a characteristic phenotype or the demonstration of the mitochondrial defect in the clinically targeted organ.

Among deleterious mtDNA mutations, point mutations affecting transfer RNA (tRNA) genes are common (see Mitomap at mitomap.org). Their identification relies on a pathogenicity scoring system using criteria from clinical investigations, database analysis, and functional studies. Among the functional studies, cybrids, i.e., cytoplasmic hybrids with the patient’s mtDNA and the nuclear genome from immortal cells bring the possibility to demonstrate the mtDNA origin of the mitochondrial defect and are thus considered a gold standard for the pathogenicity assessment of mtDNA mutations. However, as expected with mutations causing tissue-specific expression impairment, these cells, being a surrogate tissue for the disease, often disclosed only a mild mitochondrial defect.

The impact of tRNA mutations depends on the minimal set of 22 mtDNA tRNAs using a nonuniversal genetic code and specific decoding rules for mitochondrial protein synthesis. In particular, all the codons with any base in the third codon position encoding the same amino acid are recognized by only 1 tRNA that carries an unmodified U in the wobble position of the anticodon.

We here report the homoplasmic mtDNA m.15992A>T mutation of the wobble position of tRNAPro in 5 families with severe exercise intolerance and combined respiratory chain defect in muscle. We demonstrate the mutation pathogenicity in cybrids by showing its association with respiratory defect, decreased tRNAPro steady state, and altered response to depolarization reminiscent of the exercise-induced ionic disturbances observed in patients.

**Patients and methods**

**Patients**

**Patients**

Patient 1 (III-10 in family 1 in figure 1) presented with swelling of the temporal and masseter muscles during feeding since the neonatal period. Exercise intolerance occurred during early childhood, never associated with rhabdomyolysis. At age 22 years, she sought medical advice for myalgia and exhaustion after walking 500 m, followed with nausea, vomiting, and headache if effort was continued. Physical examination was normal at rest. Lactatemia was 2.1 mM in the fasting state and high after nonischemic forearm exercise test.

Patient 2 (III-13 in family 1) sought medical advice at age 14 years for clinical and biological signs similar to patient 1.

Patient 3 (II-2 in family 2) presented at age 14 years with fatigability and muscle pain after moderate exercise (1 km cycling). He also presented with bilateral cataract, which was present in his mother. His physical examination was normal apart from short stature (−2.5 SD). Lactatemia was 6 mM in the postprandial state and high after moderate effort.

Patient 4 (III-6 in family 3) complained of myalgia and fatigability since early childhood, with vomiting if effort was continued. Physical examination only disclosed short stature (−3 SD). Lactatemia was 6.3 mM in the fasting state and high after effort.

Patient 5 (III-5 in family 4) complained of myalgia and muscle fatigue since age 3 years. Vomiting and malaise after prolonged efforts started after age 6 years. At age 13 years, she could only walk 200 m. Lactatemia was high in the fasting state (3.3 mM) and after nonischemic forearm exercise test.

Patient 6 (III-5 in family 5 in figure 1) presented with congenital cataract. Severe exercise intolerance occurred in childhood and was associated with masseter swelling. At age 38 years, she could only walk 500 m because of myalgia and muscular exhaustion. Lactatemia was normal in the fasting state but high after exercise.

**Methods**

**Patients’ investigations**

All patients gave their written informed consent for their analyses for diagnostic investigations and for their use in clinical research and publication, according to our institutional ethics board. The study received approval from our Institutional Review Board.

Brain imaging used a 1.5 T General Electric MRI with a head and neck coil, including coronal and axial T1 and short-tau inversion recovery (STIR) sequences.

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**Glossary**

mtDNA = mitochondrial DNA; mt-tRNA = mitochondrial transfer RNA; STIR = short-tau inversion recovery; TMRE = tetramethylrhodamine ethyl ester.
Muscle fragments were immediately frozen and stored at −80°C until use. Standard procedures provided blood, buccal mucosa cells, urinary sediment, and cultured fibroblasts derived from a forearm skin biopsy (patients 1, 2, 3, and 4). Muscle histology followed standardized protocols.\textsuperscript{14,15}

**Molecular biology**

Extraction of DNA from muscle and fibroblasts used standard methods based on proteinase K and SDS digestion; it used QIAamp DNA Mini Kit extraction (Qiagen) for blood, cells from buccal mucosa, or urinary sediment. Extraction of RNA used the miRNA Mini Kit (Qiagen) or TRIzol (Thermo Fisher Scientific).

Long-range PCR screened for large-scale deletions, while quantitative PCR evaluated the mtDNA copy number.\textsuperscript{16} The Sanger method provided mtDNA sequence. Mispairing PCR restriction quantified heteroplasmy using the Ddel site created by the mutation. Primers are in table e-1, links.lww.com/NXG/A282.

Quantification of mitochondrial tRNAs used Northern blotting of 8% polyacrylamide 8 M urea (Tris-borate) gels on Hybond N+ membranes and T4 polynucleotide kinase—5’-end \textsuperscript{32}P-labeled probes (table e-1, links.lww.com/NXG/A282). Radioactive signals were quantified using Typhoon Trio and ImageQuant software (GE Healthcare).

Acid-denaturating gel separation and Northern blot analyzed the tRNA aminoacylation state\textsuperscript{17} while \([35S]\)-methionine in vivo labeling in the presence of emetine, a specific inhibitor of cytosolic ribosomes, assessed mitochondrial translation\textsuperscript{18} and reverse transcription followed by PCR (RT-PCR) with extended oligonucleotides and sequencing with shorter primers (table e-1, links.lww.com/NXG/A282) searched for the presence of inosine at the wobble position of tRNA anticodon.\textsuperscript{19}

**Cell biology**

Cytoplasmic hybrids (cybrids) were obtained from patients 1 and 4 fibroblasts.\textsuperscript{8} Cell respiration was analyzed in Oroboros high-resolution respirometer,\textsuperscript{20} whereas mitochondrial activities in muscle and fibroblasts used standardized spectrophotometric protocols.\textsuperscript{21} Mitochondrial production of superoxide ion was assessed by flow cytometry using 5 \textmu M MitoSOX Red (Invitrogen).\textsuperscript{22}

Mitochondrial pellet preparation and Western blot after blue native polyacrylamide gel electrophoresis were as described in

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**Figure 1** Family trees of the patients with the m.15992A>T mutation

Arrows indicate the proband in each family; stars = subjects with samples for molecular studies; white forms = subjects asymptomatic or without sufficient information; black forms = patients with severe exercise intolerance, i.e., disabling daily life; gray forms = moderate exercise intolerance, i.e., not disabling daily life but clearly experienced; striped forms = patients with isolated chewing-induced masseter swelling; question marks indicate unknown clinical status of the subject. In families 1 and 5, severe intolerance to exercise was constantly associated with chewing-induced masseter swelling, but the reverse was not true with several patients in family 5 presenting only chewing-induced masseters swelling.
Ref. 23. Primary antibodies were polyclonal antibodies against the F1 domain of bovine complex V, UQCRC2, MT-ND1, or MT-CO2 (respectively produced by Pr Joël Lunardi, Grenoble, by Dr Catherine Godinot, Lyon, or our group24) or monoclonal antibodies against the respiratory complex II SDHA subunit SDHA (Abcam), incubated overnight at 0°–4°C. Secondary antibodies were peroxidase conjugated (Sigma-Aldrich) and visualized with Pierce™ ECL Western Blotting Substrate (Life Technologies). Quantification used volumes and basal adjustment with rolling ball by Fusion FX (Vilber Company).

To analyze cell proliferation, 3,000 cells per well were cultured in four 24-well plates and Dulbecco Modified Eagle Medium with 1 mM pyruvate and either 1 g/L glucose or 200 μM glutamine without glucose.25 Cell numbering was performed, in triplicates for each condition, every 24 hours (T0, T24, T48, and T72 plates) using the neutral red method26 and expressed as fold increase relative to T0.

To evaluate plasma membrane, 100,000 cybrid cells were incubated in 96-well plate with 7.5 μM tetramethylrhodamine ethyl ester (TMRE) and 5 μM verapamil, an inhibitor of the multidrug resistance–associated proteins. Because resting membrane potential essentially depends on K diffusion gradient, progressive increase of external K concentration (from 10 to 80 mM) induced progressive depolarization. Osmolarity was kept at 300 mOsm/L. After 120-minute incubation, fluorescence was quantified by the Accuri™ C6 flow cytometer.

Statistics
Normality evaluation used the Shapiro-Wilk test. Depending on the distribution of data, comparison between 2 groups used the Mann-Whitney or t test. Differences were considered significant when p < 0.05.

Data availability
Any data not published within the article will be shared, as anonymized data, by request from any qualified investigator.

Results
The chewing-induced masticatory muscle swelling was due to edema
Nuclear MRI of facial muscles analyzed the exercise-induced muscle swelling in 2 members of family 5. Chewing a sandwich during 15 minutes induced a 20%–30% increase in thickness of the temporal and masseter muscles of patient 6 (figure 2, A and B) and her aunt (II-5 in family 5) (figure 2, C and D). Mild adipose infiltration of the masseter and lateral pterygoid muscles was present in patient 6’s aunt. Prolonged edema was indicated by hypersignal in STIR, fat-suppression images, in all masticatory muscles of patient 6 (figure 2, E and F) and in the masseter, temporal, and lateral pterygoid muscles of her aunt (figure 2, G and H).

Mitochondrial myopathy underlay the muscle symptoms
Muscle histology disclosed mild alterations in patients 2, 3, 4, and 5, including moderate lipidosis and increased subsarcolemmal...
mitochondria (figure e1A, links.lww.com/NXG/A279). It was considered normal for patient 6.

Spectrophotometric assays of the respiratory chain revealed severe combined defect of respiratory complexes I, III, and IV with increased citrate synthase in the muscle biopsies from all probands (table 1). Respiratory complex II, the only complex without mtDNA-encoded subunit, was either normal or elevated.

**All patients shared the same homoplasmic MT-TP m.15992A>T mutation**

Long-range PCR excluded the presence of large size rearrangement, whereas quantitative PCR ruled out depletion. Sequencing of the whole mtDNA sequence revealed that all patients had the same mtDNA, sharing 31 of 32 variants (table e2, links.lww.com/NXG/A283). Thirty-one of these 32 variants were highly likely polymorphisms: 28 reported at least 1,462 times in GenBank database, a MT-RNR1 variant reported 37 times in different ethnic backgrounds, and 2 synonymous mutations (MTND2 c.5024C>T and MT-CO1 m.7028C>T) (mitomap.org). These polymorphisms defined the mtDNA haplotype as J1c10 for all the families. The last variant, common to all patients, was the m.15992A>T mutation in the MT-TP gene. Several criteria indicated its potential pathogenicity: (1) it was reported only once in more than 49,000 GenBank full-length mtDNA sequences; (2) it modified the strictly phylogenetically conserved wobble base in the tRNA anticodon (see MitotRNAdb/Mamit-tRNA website at mttrna.bioinf.uni-leipzig.de); and (3) alteration of the wobble position in mt-tRNA anticodon is a major deleterious factor in several mt-tRNA confirmed mutations.27

The mutation appeared homoplasmic (figure 3) with PCR restriction in the DNA samples from muscle of all the probands, from blood of all patients marked with a star in figure 1, and from urinary sediment and buccal cells of patients 1 and 2, their mother, and grandmother.

**Transfer of the m.15992A>T mutation into cybrids demonstrated its deleterious potential on OXPHOS activities**

Fibroblasts derived from a skin biopsy of patients 1, 2, 3, and 4 had 100% mutation (figure 3). They disclosed a mild but significant decrease of their basal respiration and respiration linked to adenosine triphosphate (ATP) production (figure 4A). Citrate synthase activity was significantly increased (figure 4B). Complexes I, III, IV, and V appeared decreased, reaching significance for complex III after normalization to citrate synthase (figure 4, B and C).

Cybrid clones, derived from patient 2 and patient 4 fibroblasts, had homoplasmic m.15992A>T mutation (figure 3). They showed decreased respiration, either basal, linked to ATP production, or maximal (figure 4A). As in fibroblasts, cybrids had decreased activity for all the mtDNA-depending complexes, reaching significance for respiratory complex III (figure 4, B and C). The amount of the respiratory complexes appeared normal when analyzed by Western blot after blue native polyacrylamide gel electrophoresis (figure e1B, links.lww.com/NXG/A279).

Mutant cells disclosed significantly slower proliferation than control cells in the absence of glucose, whereas their proliferation rate was identical in its presence (figure 5A). The mild oxidative phosphorylation pathway (OXPHOS) defect was therefore relevant, affecting the cell proliferation capacity in a medium without glucose.25,28,29 It had no apparent impact on the mitochondrial production of superoxide ion, which was similar in mutant and control cybrids in the basal state (figure 5B).30

### Table 1 Mitochondrial activities in the muscle of patients with homoplasmic m.15992A>T mutation

| Activity                  | P2       | P3       | P4       | C1 (n = 200) | P5       | P6       | C2 (n = 140) |
|---------------------------|----------|----------|----------|-------------|----------|----------|-------------|
| CI total                  | 11±      | 8±       | 7±       | 25 ± 7      | 17±      | 21±      | 48 ± 15      |
| CI rot sens               | 0±       | 6±       | 0±       | 22 ± 7      | 12±      | 18±      | 42 ± 14      |
| CII                       | 39       | 90       | 30       | 31 ± 8      | 77       | 136      | 62 ± 19      |
| CIII total                | 80±      | 26±      | 23±      | 124 ± 274   | 70±      | 63±      | 235 ± 64     |
| CIII am sens              | 22±      | 11±      | 5±       | 107 ± 26    | 6±       | 23±      | 160 ± 59     |
| II + CIII                 | 3±       | 5±       | 2±       | 18 ± 7      | 14±      | 13±      | 50 ± 15      |
| CIV                       | 43±      | 63±      | 9±       | 56 ± 17     | 109±     | 89±      | 199 ± 57     |
| CS                        | 314      | 1,005    | 248      | 153 ± 35    | 276      | 470      | 209 ± 64     |

Abbreviations: CI total = NADH ubiquinone oxidoreductase activity; CI rot sens = CI fraction sensitive to rotenone inhibition, i.e., specific respiratory complex I activity; CI = succinate ubiquinone oxidoreductase activity, i.e., respiratory complex II activity; CIII total = ubiquinol cytochrome c oxidoreductase activity; CIII am sens = CIII fraction sensitive to antimycin inhibition, i.e., specific respiratory complex III activity; CIV = cytochrome c oxidase activity, i.e., respiratory complex IV activity; CS = activity of citrate synthase, a Krebs cycle enzyme considered as representing the mitochondrial mass; n = number of different controls analyzed in each series.

Activities in bold font indicate values below the 10th centile of control values; activities in italics font indicate values above the 90th centile of control values.

*After the respiratory activities indicate values below the 10th centile of control values after their normalization to citrate synthase activity. P2, P3, P4, P5, and P6 = results obtained in the muscle biopsy from patients 2, 3, 4, 5, and 6; C1, C2 = successive control series due to the change of spectrophotometric assays in 2009.21
The mutant tRNA\textsuperscript{Pro} had a lower steady-state but normal aminoacylation

Northern blot analysis in several cell lines revealed significant decrease of the amount of the mitochondrial tRNA\textsuperscript{Val} in all the mutant cell lines, fibroblasts, or cybrids (figure e-2A, links.lww.com/NXG/A280). Cybrids also disclosed an increase of tRNA\textsuperscript{Val}, which could suggest an increase in the mitochondrial ribosomes because tRNA\textsuperscript{Val} is one of their integral components.\textsuperscript{31}

Abnormal decrease of the cell volume in mutant cybrids on depolarization by external potassium was reminiscent of the exercise-induced muscle edema observed in patients

Ionic disturbances were a striking observation in 2 families with the m.15992A>T mutation. They were reminiscent of ionic disturbances observed in patients with homoplasmic MT-ATP6 mutations and recurrent paralysis episodes associated with significant plasma membrane depolarization in fibroblasts.\textsuperscript{22} Core facilities providing patch-clamp electrophysiologic analyses are scarce. Therefore, to analyze the plasma membrane potential in cybrids, we used an indirect determination based on TMRE, a fluorescent probe that follows Nernst equation and thus may be used to analyze membrane potentials.\textsuperscript{4} Because 7.5 \(\mu\)M TMRE significantly decreases respiration,\textsuperscript{4} the mitochondrial membrane potential does not influence the TMRE fluorescence signal, which thus essentially represents both the cell size and plasma membrane potential. As expected, increasing in isotonic conditions the external K concentration from 5 to 10, 20, 30, 40, and 80 mM led to progressive decrease of TMRE fluorescent signal to 91 ± 9, 78 ± 12, 73 ± 7, 61 ± 9, and 38 ± 10% of its initial value, respectively (analysis of 33 independent cell populations, mutant and wild type grouped). At the basal state, the size (evaluated by the forward scatter) and the TMRE fluorescence (evaluated by the fluorescence area FLA) did not differ between wild-type and mutant cybrids with either the m.15992A>T mutation or the m.9185T>C mutation previously associated with permanent plasma membrane depolarization\textsuperscript{22} (figure 5, C and D). In contrast, at 80 mM external K concentration, both types of mutant cybrids had significantly decreased their size, whereas wild-type cybrids had maintained a size similar to the basal state (figure 5C). All 3 cybrid types had
decreased their TMRE signal, in accordance with the induced decrease of plasma membrane potential, but mutant cybrids to a level significantly lower than wild-type cybrids.

**Discussion**

In this article, we demonstrate the pathogenicity of an original homoplasmic MT-TP mutation targeting skeletal muscle. In muscle, the severe combined respiratory chain defect demonstrated the mtDNA origin of the disease. However, we could not exclude a nuclear DNA alteration solely affecting muscle mtDNA expression because the absence of paternal transmission by the few affected fathers was insufficient to demonstrate maternal inheritance. Therefore, we had to provide demonstration of the mutation deleterious potential.

As observed with other deleterious homoplasmic mtDNA mutations with restricted clinical expression, fibroblasts and cybrids presented with very mild enzymatic defect. However, that defect was statistically significant with unbiased nonparametric statistical tests. In addition, it induced significant reduction of the mutant cells proliferation in the absence of glucose showing its relevance to cell physiology. Because most organs essentially comprise postmitotic cells, cell proliferation is not relevant for clinical symptoms. Therefore, we addressed the ionic disturbances that were a striking aspect of the disease, prominent in 2 families. When faced to an isosmotic high potassium concentration, inducing plasma...
membrane depolarization, cybrids with the MT-TP mutation differed from wild-type cybrids by losing part of their cell volume, most probably through loss of internal fluid. That behavior fitted with the edema observed in patients using muscle imaging after 15-minute chewing effort.

At the molecular level, the mutant mitochondrial tRNA<sub>Pro</sub> had a reduced steady-state level in cells. It bore an unmodified adenosine at the wobble position of its anticodon, which thus should optimally recognize only codon CCU that represents less than 25% of the total mtDNA proline codons. Special mitochondrial decoding rules probably allowed the mutant tRNA to recognize all the 4 proline codons, but less efficiently. These 2 anomalies (reduced tRNA steady-state and decoding problems) might have a cumulative effect on mitochondrial translation. Why the impact on OXPHOS was mild in fibroblasts and cybrids but drastic in muscle remained hypothetical in the absence of appropriate muscle fragments.

The role of the J1c10 haplotype in the mutation pathophysiology was disputable. That haplotype was constant in the 5 families here reported and in the sole GenBank report of the mutation. It is not pathogenic in itself, being observed in 2.5%–5% of the people living in the western half of France, but it is a significant modifier in Leber hereditary optic neuropathy<sup>37,38</sup> and in the cybrid resistance to rotenone toxicity.<sup>39</sup>
According to the pathogenicity scoring system for tRNA mutations updated in 2011,6 mutation m.15992A>T has a score of 13, which indicated a mutation definitely pathogenic (2 for independent pedigrees, 2 for phylogenetic conservation, 2 for muscle histology, 2 for biochemical defects, and 5 for hybrid analysis).

The homoplasmic m.15992A>T MT-TP mutation in the J1c10 haplotype causes severe intolerance to exercise, often associated with exercise-induced muscle edema.

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| Name                  | Location                              | Contribution                                      |
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| Karine Auré, MD, PhD  | Department of Neurophysiology, Foch Hospital, Suresnes, France | Acquisition of data; analysis of data; and drafted and revised the manuscript |
| Guillemette Fayet, MD, PhD | Centre de Référence Maladies Neuromusculaires Hôtel-Dieu, CHU Nantes, France | Acquisition of data; analysis of data; and drafted and revised the manuscript |
| Ivan Chicherin, PhD   | M.V. Lomonosov State University, Moscow, Russia | Acquisition of data and analysis of data |
| Benoit Rucheton, PharmD | Service de Biochimie Métabolique CHU Pitié-Salpêtrière, AP-HP, Paris, France | Acquisition of data; analysis of data; and drafted and revised the manuscript |
| Sandrine Filaut       | Service de Biochimie Métabolique CHU Pitié-Salpêtrière, AP-HP, Paris, France | Acquisition of data and analysis of data |

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**Appendix 1 (continued)**

| Name                  | Location                              | Contribution                                      |
|-----------------------|---------------------------------------|--------------------------------------------------|
| Julie Eichler, MSc    | CNRS UMR 7156 GMGM, University of Strasbourg, France | Acquisition of data and analysis of data |
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