Clinical Study

Novel m.15434C>A (p.230L>I) Mitochondrial Cytb Gene Missense Mutation Associated with Dilated Cardiomyopathy

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Background. Previously it has been shown that various types of hypertrophic and dilative cardiomyopathy (hCMP, dCMP) can be attributed to disturbed mitochondrial oxidative energy metabolism. Several studies described mutations in mitochondrial DNA-located genes encoding for subunits of respiratory chain complexes, including the cytochrome b gene (MT-CYB), causing CMPs. Methods and Results. In the present study the MT-CYB gene was analysed in 30 patients with hCMP, 40 patients with dCMP, and 50 controls for alterations. Altogether, 27 MT-CYB variants were detected. Twenty-four of them were single nucleotide polymorphisms defining common haplogroups. The variant m.15434C>A was found in a single patient with severe dCMP and assessed as novel mutation, since it was not found in healthy controls or available data sets, and was nonhaplogroup associated with Phylotree. This variant altered an amino acid (L230I) with a high interspecific amino acid conservation index (CI = 97.7%) indicative of the functional importance of the residue. Conclusions. Though the L230I mutation seems to play a causative role for dCMP, prospective studies on yeast or transgenic mice models with defined mutation are warranted to study the pathogenetic impact of this mutation.

1. Introduction

Mitochondria play a critical role in both life and death of cardiomyocytes. In healthy cells, their primary function is to meet the high energy demand of the beating heart by providing ATP through oxidative phosphorylation. Mitochondria occupy a large portion of each myocyte and are located between the myofibrils or right below the sarcolemma. The strategic positioning and abundance of mitochondria ensure a highly efficient localized ATP delivery system to support contraction, metabolism, and ion homeostasis [1]. Mitochondrial disorders (MIDs) leading to myocardial disease show a strong age-dependent clinical heterogeneity [2]. Cardiac manifestations of MIDs include arrhythmias and cardiomyopathy (CMP; hypertrophic, (hCMP), dilated (dCMP), restrictive (rCMP), histiocytoid CMP, and non-compaction) [3, 4]. Mitochondria contain their own DNA (mitochondrial DNA (mtDNA)) which is a circular, 16 569 base sequence with 37 genes that encodes for 13 subunits of the respiratory chain complexes, 22 transfer RNAs, and 2 ribosomal RNAs [5]. Mutations in mtDNA-encoded genes influence the production of reactive oxygen species in mice [6] and have been implicated in a large number of human and murine diseases [7]. Some amino acid changes may also improve aerobic capacity and adaptation to new thermal environments [8]. One of the mtDNA located genes associated with cardiac disease is MT-CYB encoding for cytochrome-b. Base substitutions in the human MT-CYB gene are associated with a wide spectrum of CMPs [9, 10] or exercise intolerance [11–14]. MT-CYB is an extremely conserved protein, reflecting its fundamental role in energy production in mitochondria. It catalyses the reversible electron transfer from ubiquinol to cytochrome c coupled to
proton translocation (Q-cycle) [15]. The aim of the present study was to analyse the MT-CYB gene from CMP patients and controls and to identify potential pathogenic mutations associated with hCMP or dCMP phenotypes.

2. Materials and Methods

2.1. Selection of Patients. Blood samples were collected from 70 patients with CMPs who originated from the area of Tunis and were recruited from the department of cardiology in the public teaching Hospital la Rabta Tunis, Tunisia. Age at onset of the disease varied from adolescence to adulthood. They comprised 40 patients with dCMP with a mean age of 39.3 ± 15.2 years and 30 patients with hCMP with a mean age of 41.87 ± 19.13 years (Table 1). All patients had angiographically normal coronary arteries and were not affected by other cardiac or systemic disease. CMP was diagnosed according to the WHO/ISFC criteria [16]. dCMP was diagnosed in case of left ventricular dilatation (LVEDD > 55 mm) and reduced ejection fraction. hCMP was diagnosed if there was myocardial thickening >14 mm on echocardiography (Table 1). Blood samples were collected with informed consent also from 50 unrelated healthy controls originating from the same area as the patients without clinical evidence or a family history of CMP.

2.2. MT-Cytochrome b Gene Analysis. Total DNA extracted from lymphocytes (according to conventional methods) was submitted to polymerase chain reaction (PCR) amplification, by using two pairs of specific primers designed from the genomic sequence of the mitochondrial MT-CYB gene (14747–15887). Forward primers and reverse primers (5′-3′) were as follows: pair 1 (14667–14687 and 15289–15269) and pair 2 (15189–15209 and 15941–15921). Amplification was carried out in a total volume of 50 μL containing 100 ng of total DNA, 10 pmol of each primer, 4 μL MgCl2, 5 μL dNTP, and 0.2 μL Taq polymerase. The amplification conditions included 3 min at 94°C, 30 cycles of 30 s at 94°C, 60 s at 52°C, and a final extension of 10 min at 72°C terminated the PCR. The PCR products were checked by a 1.5% agarose gel electrophoresis and an ethidium bromide staining. After amplification, the PCR product was subjected to denaturing gradient gel electrophoresis (DGGE). It was run on a Tris-EDTA-acetate buffer (TEA: Tris 40 mM; Na-acetate 20 mM; EDTA 1 mM) at 60°C, using a 6.5% polyacrylamide gel containing a gradient of denaturing agents (60% denaturant = 4.2 M and 24% formamide). Gels were stained with ethidium bromide and examined under ultraviolet light, and images were saved electronically (Figure 1).

2.3. Direct Sequencing of PCR-Amplified Fragments. Direct sequencing of the PCR fragments was performed using an ABI 3730 automated DNA sequencer (Applied Biosystem) by using a BigDye Terminator V.3.1 cycle sequencing kit (applied 4337455 (100 reactions)) according to the manufacturer’s recommendations. MT-CYB gene sequences from patients and controls were compared with the mtDNA reference, the revised Cambridge Reference Sequence (rCRS; NC_012920) [17].

2.4. Software and Databases. We used the tool PolyPhen-2 (http://genetics.bwh.harvard.edu/pph2/) [18] for predicting the damaging effect of a missense mutation. The tool PolyPhen-2 compiles two pairs of datasets, the HumDiv and the UniProt. The conservation index proposed by Ruiz-Pesini et al. [19] was computed by using MitoTool (http://www.mitotool.org/) [20]. A total of 43 primate species were considered and a conservation index (CI) of 0.744 meant that 74.4% of 43 primate species had the wild-type allele in common with the human sequence (GenBank Accession number NC_012920). The presence of the mutation as a haplogroup-specific variant was scored relative to the available global mtDNA phylogenetic tree by using Phylotree (http://phylotree.org/tree/main.htm; [21]).
In case MIDs exclusively affected a limited number of cells and resulting in tissue-specific phenotypes \[22, 23\].

The diagnosis of mitochondrial CMPs may be established clinically indistinguishable from CMPs of other cause \[3, 24\].

### 4. Discussion

It is well known that mutations in the MT-CYB gene are often sporadic and arise during embryogenesis, affecting a limited number of cells and resulting in tissue-specific phenotypes \[22, 23\]. In case MIDs exclusively affect the heart, mitochondrial hCMP or mitochondrial dCMP may be clinically indistinguishable from CMPs of other cause \[3, 24\].

The diagnosis of mitochondrial CMPs may be established by mtDNA sequence analysis for known pathogenic mtDNA mutations. Reviewing published literature, we detected eight nonsynonymous mutations distributed across the MT-CYB and the majority were rare (allele frequency < 5%). The amino acid change induced by these eight mutations was considered to be mild because either the modified amino acid or an amino acid with similar physicochemical characteristics was commonly encountered at that position in other animal species variants and defined common haplogroups. One of the eleven non-synonymous mutations was novel and was detected in a patient with severe dCMP. The novel m.15434C>A variant could not be found in 50 healthy controls, among confirmed variants of MID in MITOMAP: a Human Mitochondrial Genome Database (http://www.mitomap.org/, 2011) in previous studies on CMPs, or in human mitochondrial genome databases (http://www.genpat.uu.se/mtDB/). The variant was deemed pathogenic on the basis of the following findings: (1) the m.15434C>A mutation resulted in the substitution of a glycine by an aspartate residue at position 251 of cytochrome b (G251D) associated with histiocytoid CMP \[10\]. This mutation was located in the intermembrane space loop connecting the fifth and sixth transmembrane segments. Analyses on mitochondria purified from patients’ hearts have detected a defect in the succinate cytochrome c oxidoreductase activity and in the cytochrome b assembly, showing the importance of residue 251 for bc1 (complex III) function. A finding which challenges the high conservation of the adenine residue at position 230 of MT-CYB, suggesting that it may have resulted from oxidative damage since it could not be inherited \[25, 26\]; (2) the mutation was located in the transmembrane region of MT-CYB and altered an amino acid (L230I) with a high interspecific amino acid CI of 97.7%, indicative of the functional importance of the residue; (3) nonhaplogroups were associated on Phylotree; (4) from all 3,155 damaging alleles annotated in the UniProt database causing human Mendelian diseases and affecting protein stability or function, and from 6,321 differences between human proteins and their closely related mammalian homologues, it was assumed that the mutation was non-damaging. The second pair, HumVar, consists of all the 13,032 human disease-causing mutations from UniProt and 8,946 human nonsynonymous single-nucleotide polymorphisms (nsSNPs), and predicting the mutation as possibly damaging with a score of 0.563. All these findings indicated that the novel mutation had an effect on the mitochondrial oxidative metabolism and cellular physiology.

Generally, missense mutations in MT-CYB may impair assembly or stability of the complex, leading to a dramatic decrease in the levels of assembled enzyme, or they may alter the catalytic activity of the complex with little effect on its assembly \[27\]. Several pathogenic variants in the MT-CYB gene have been previously reported linked to CMPs \[9, 10\]. One mutation that substituted a glycine by an aspartate residue at position 251 of cytochrome b (G251D) was associated with histiocytoid CMP \[10\]. This mutation was located in the intermembrane space loop connecting the fifth and sixth transmembrane segments. Analyses on mitochondria purified from patients’ hearts have detected a defect in the succinate cytochrome c oxidoreductase activity and in the cytochrome b assembly, showing the importance of residue 251 for bc1 (complex III) function. A finding which challenges the high conservation of the m.15434C>A variant is the presence of the polymorphism m.15434C>T (p.230L>F) in two individuals and the fact that phenylalanine is not much different from isoleucine \[28\].

In conclusion, the present study implies that the MT-CYB gene mutation L230I plays a pathogenetic role in the
| Variant   | dCMP (n = 40) | hCMP (n = 30) | Co (n = 50) | Amino acid position, physicochemical property | CI     | Haplogroup specific variant | Status          |
|-----------|---------------|---------------|-------------|----------------------------------------------|--------|---------------------------|----------------|
| C14766T  | 30/40         | 7/30          | 26/50       | T7I, polar (T) to non polar (I)               | 0.279  | B2a1a, M2b, E1a1, HV     | Polymorphism    |
| A14769G  | 2/40          | 0/30          | 0/50        | N8S, medium, polar (N) to small, polar (S)   | 0.581  | L1b, L3f1b, L3f1b       | Polymorphism    |
| T14798C  | 1/40          | 0/30          | 5/50        | F181L, large, aromatic (F) to medium, hydrophobic (L) | 0.581  | J1c, T2g, K, L1c6      | Polymorphism    |
| A14927G  | 1/40          | 0/30          | 0/50        | T61A, medium, polar (T) to small, hydrophobic (A) | 0.581  | D4b1a, D5c, U6a1       | Polymorphism    |
| G15257A  | 1/40          | 0/30          | 0/50        | D171N, medium, acidic (D) to medium, polar (N) | 0.860  | K1b1a, M71a1, J2      | Possibly deleterious |
| G15314A  | 1/40          | 0/30          | 1/50        | A190T, small, hydrophobic (A) to medium, polar (T) | 0.326  | R9b2, L3j, L3k, M38, D4a1a1, P4a, L3j, | Polymorphism    |
| A15326G  | 8/40          | 5/30          | 49/50       | T194A, medium, polar (T), to small, hydrophobic (A) | 0.442  | Z3a1, H2a2a, R7b1a, R8a | Polymorphism    |
| C15434A  | 1/40          | 0/30          | 0/50        | L230I, both residues are medium and hydrophobic | 0.977  | Non-haplogroup variants | Novel missense mutation |
| C15452A  | 2/40          | 0/30          | 6/50        | L236I, similar physicochemical property, both residues are medium size and hydrophobic | 0.744  | JT            | Polymorphism    |
| G15803A  | 1/40          | 0/30          | 0/50        | V353M, similar physicochemical property, both residues are medium size and hydrophobic | 0.395  | L2a2a        | Polymorphism    |
| G15884C  | 1/40          | 0/30          | 0/50        | A380P, small, hydrophobic (A) to medium, hydrophobic (P) | 0.047  | L0F2b        | Polymorphism    |

The tool PolyPhen-2 (http://genetics.bwh.harvard.edu/pph2/) was used for predicting the damaging effect of a missense mutation. The conservation index (CI) was computed by using MitoTool (http://www.mitotool.org/). The presence of the mutation as a haplogroup-specific variant was scored relative to the available global mtDNA phylogenetic tree by using Phylotree (http://phylotree.org/tree/main.htm). Co: controls.
Table 3: Silent mutations (synonymous single nucleotide polymorphisms) in the mtDNA cytochrome b gene (MT-CYB) identified in patients with CMP.

| Variants   | dCMP n = 40 | hCMP n = 30 | Controls n = 50 | Amino acid position | Status          |
|------------|-------------|-------------|-----------------|---------------------|-----------------|
| T14935C    | 1           | 0           | 0               | F63F                | Polymorphism    |
| T15344C    | 1           | 0           | 0               | L200L               | Polymorphism    |
| C15632T    | 1           | 0           | 0               | L296L               | Polymorphism    |
| A15679G    | 1           | 1           | 0               | K311K               | Polymorphism    |
| A15799G    | 1           | 0           | 0               | G351G               | Polymorphism    |
| T14783C    | 0           | 0           | 2               | L13L                | Polymorphism    |
| C14905A    | 1           | 1           | 1               | M53M                | Polymorphism    |
| G15043A    | 1           | 1           | 4               | G99G                | Polymorphism    |
| T15115C    | 1           | 0           | 0               | T123T               | Polymorphism    |
| G15148A    | 1           | 0           | 0               | P134P               | Polymorphism    |
| A15244G    | 1           | 0           | 1               | G166G               | Polymorphism    |
| G15301A    | 1           | 1           | 4               | L185L               | Polymorphism    |
| T15454C    | 0           | 1           | 0               | L236L               | Polymorphism    |
| T15514C    | 1           | 0           | 0               | Y256Y               | Polymorphism    |
| T15530C    | 1           | 0           | 0               | L262L               | Polymorphism    |
| T15787C    | 1           | 0           | 0               | F347F               | Polymorphism    |

The 16 variants were interpreted as synonymous mutations, which do not cause any amino acid change, and were reported as single nucleotide polymorphisms in MITOMAP: A Human Mitochondrial Genome Database. (http://www.mitomap.org/, 2011).

development of primary CMPs. The novel potentially deleterious MT-CYB gene mutation L230I has been described for the first time in a patient with primary dCMP. However, prospective studies on yeast or transgenic mice with defined mutation are warranted to study the impact and contribution of this mutation on the pathophysiology of dCMP.

Conflict of Interests

None of the authors had any personal or financial conflict of interests.

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