A novel MT-CO2 variant causing cerebellar ataxia and neuropathy: The role of muscle biopsy in diagnosis and defining pathogenicity

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Abstract

Pathogenic variants in mitochondrial DNA (mtDNA) are associated with significant clinical heterogeneity with neuromuscular involvement commonly reported. Non-syndromic presentations of mtDNA disease continue to pose a diagnostic challenge and with genomic testing still necessitating a muscle biopsy in many cases. Here we describe an adult patient who presented with progressive ataxia, neuropathy and exercise intolerance in whom the application of numerous Mendelian gene panels had failed to make a genetic diagnosis. Muscle biopsy revealed characteristic mitochondrial pathology (cytochrome c oxidase deficient, ragged-red fibers) prompting a thorough investigation of the mitochondrial genome. Two heteroplasmic MT-CO2 gene variants (NC_012920.1: m.7887G>A and m.8250G>A) were identified, necessitating single fiber segregation and familial studies – including the biopsy of the patient’s clinically-affected mother - to demonstrate pathogenicity of the novel m.7887G>A p.(Gly101Asp) variant and establishing this as the cause of the mitochondrial biochemical defects and clinical presentation. In the era of high throughput whole exome and genome sequencing, muscle biopsy remains a key investigation in the diagnosis of patients with non-syndromic presentations of adult-onset mitochondrial disease and fully defining the pathogenicity of novel mtDNA variants.

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1. Introduction

The phenotypic spectrum and genetic aetiology of mitochondrial disease are heterogeneous and the diagnostic challenges are well-recognized [1]. The increased application of whole exome sequencing (WES) and whole genome sequencing (WGS) as a first step in the diagnostic pathway has successfully identified many novel Mendelian
mitochondrial genes or new pathogenic variants in known disease genes over the last decade [2,3]. In the adult population, around two thirds of patients with mitochondrial disease have pathogenic mitochondrial DNA (mtDNA) variants of which some of these variants cannot be reliably detected in blood [4]. Typical clinical phenotypes, such as chronic progressive external ophthalmoplegia (CPEO) [5] or multisystem involvement frequently prompt clinicians to consider mitochondrial disease and perform muscle biopsy to guide further genetic investigation. However, adult-onset, non-syndromic presentation of mtDNA disease remain difficult to diagnose because a high index of clinical suspicion is required to distinguish it [6,7] from other genetic mimics including autosomal recessive cerebellar ataxia and hereditary neuropathy [8]. A study from the ataxia specialist centers identified approximately a quarter of the patients with ‘idiopathic’ progressive cerebellar ataxia had canonical features of mitochondrial disease in their muscle biopsies, highlighting the utility of muscle biopsy should not be confined to the investigation of myopathy [9].

Here we report a patient who presented with adult-onset progressive ataxia, neuropathy and exercise intolerance without relevant family history. Extensive genetic testing had failed to provide a genetic diagnosis prior to muscle biopsy revealing marked mitochondrial histopathological changes. We detail the work-up of genetic investigations that identified two mtDNA variants in the same gene and how access to familial samples including the biopsy of his clinically-unaffected mother helped define the causal mtDNA genetic variant.

2. Methods

2.1. Case report

A 45-year-old man presented to the neurology clinic with slowly progressive sensory symptoms in the lower limbs and impaired balance since his early twenties. He also reported muscle pain after physical exertion but never noticed discoloured urine. He was unsteady when he walked and this became worse in the dark. He was never good at sport at school; he could run but avoided sports altogether since early teenage. Upper limb incoordination, slurred speech, worsening exercise intolerance and fatigue were reported in his thirties. As his unsteadiness progressed, he started to fall up to four times a week and began to use a walking stick when he was 40 years old. He suffered from chronic constipation since his mid-30s; he also complained of erectile dysfunction but did not have any urinary symptoms. He was aware of mild hearing deficits since his late 20s.

Other past medical history included anxiety and depression. His birth history was unremarkable. He achieved all neurodevelopmental milestones other than he walked at a slightly older age (14 months) compared to his younger sister. There was no family history of neuromuscular disease, ataxia, diabetes or deafness.

The main clinical examination findings between aged 37 and 40 years old were dysarthric speech, upper and lower limb ataxia, broad-based gait and positive Romberg’s sign. In addition, all the deep tendon reflexes were absent, there was evidence of reduced pin prick sensation following the stocking distribution and asymmetrical loss of vibration sense (left knee and right rib coastal margin, respectively). There was no eyelid ptosis, his eye movement was normal, and muscle tone and power were within the normal limit.

Routine laboratory investigations including normal vitamin B12 and folate, autoimmune screen, creatinine kinase, Vitamin A and E levels, were unremarkable. Serum lactate was marginally raised at 2.7 mmol/L (normal range up to 2.2 mmol/L). Urinary thymidine and uridine levels were within normal limits. Nerve conduction studies and electromyography, performed at the age of 35 years, identified axonal sensory neuropathy and mild neurogenic changes in muscles. A repeat study five years later showed little progression, with retained but reduced amplitudes in sensory nerve action potentials (SNAP) in all limbs (sural amplitude 2μV for both sides; and superficial peroneal amplitude 2μV on the right, absent on the left), and normal motor nerve function.

Given the complaint of muscle fatigue and exercise intolerance, blood spot and plasma acylcarnitine profiles were performed and showed increased medium to long chain acylcarnitine species. He reported some improvement of exercise intolerance with the supplementation of riboflavin and co-enzyme Q10. His-resting 12-lead electrocardiogram and transthoracic echocardiogram were normal. MRI head showed mild cerebellar atrophy and a T2 hyperintensity in the right cerebellar peduncle, without any contrast enhancement (Fig. 1). A detailed ophthalmological assessment identified pigmentary retinal changes and slight optic disc pallor.

The overall clinical picture was one of cerebellar ataxia with sensory neuropathy and pigmentedary retinopathy. He was tested for trinucleotide expansions in common spinocerebellar ataxia (SCA) and frataxin genes, but no abnormalities were found; a screen of the POLG gene was normal. Several gene panels including Charcot Marie Tooth neuropathy type 2 (24 genes), autosomal dominant and recessive SCA (21 genes), hereditary sensory neuropathy (11 genes) and hereditary spastic paraplegia (12 genes) were also uninformative (Gene targets identified in Supplemental appendix).

In the absence of a genetic diagnosis, he underwent skeletal muscle biopsy of vastus lateralis at the age of 44 years. His-family members were reviewed and examined in the clinic to help with the segregation studies of identified genetic variants. The patient’s mother was 72 years old at the time of review, her past medical history including breast cancer treated with mastectomy and chronic low back pain. Her neurological examination including fundoscopy was within normal limits. He has a younger sister (examined at the age of 42 years) who is fit and well and with a normal neurological assessment.
2.2. Histopathological analysis of the muscle biopsy

Standard histological (haematoxylin and eosin (H&E); Gomori trichrome; Sudan black) and histochemical (cytochrome c oxidase (COX), succinate dehydrogenase (SDH) and sequential COX/SDH activities) investigations were performed on frozen, transversely orientated skeletal muscle (10μm) sections as described [10]. Additionally, quantitative, quadruple immunohistochemical (IHC) analyses were used to assess mitochondrial OXPHOS function, interrogating complex I (NDUFB8), complex IV (COX1) and porin (VDAC1; a mitochondrial mass marker) protein expression [11].

2.3. Mitochondrial genetic studies

Total DNA was extracted from skeletal muscle by standard procedures. Screening for the common m.3243A>G variant was undertaken using a quantitative pyrosequencing assay, while large scale mtDNA rearrangements were assessed using several long-range polymerase chain reaction (PCR) assays. The entire mitochondrial genome (GenBank Accession number: NC_012920.1) was amplified, analyzed and sequenced at a minimum read depth of 200x as previously reported [10].

2.4. Single fibre segregation analysis and mtDNA heteroplasmy assessment

Two mtDNA variants, m.7887G>A (novel) and m.8250G>A (extremely rare), were identified in the MT-CO2 gene following the sequencing of the mitochondrial genome in the patient’s muscle sample. These variants were investigated further in a broad range of tissues (skeletal muscle, urinary sediments, blood and buccal epithelia) and individual (COX-deficient and COX-positive) skeletal muscle fibers isolated by laser-capture microdissection using a quantitative pyrosequencing assay designed to accurately determine the level of heteroplasmy at these sites. The two novel variants were also investigated in non-invasive tissues (urinary sediments, blood and buccal epithelia) from the patient’s mother and sister using this method. The assay was performed on a PyroMark Q24 platform (Qiagen) and used variant-specific primers (available from the authors on request). The assay can reliably detect to a level of >3% heteroplasmy. Data analysis was performed using PyroMark Q24 software (v2.0.7).

3. Results

3.1. Histochemistry and immunohistochemical analyses of patient muscle

Histological and histochemical assessment of the skeletal muscle sections showed evidence of mitochondrial dysfunction following sequential COX/SDH histochemistry, demonstrating ~80% COX-deficient fibers (Fig. 2A) as well as ragged-red fibers in the Gomori trichrome and excess lipid accumulation. IHC analyses confirmed extensive mitochondrial dysfunction, >60% of all fibres analysed (n=1030), showing complete loss of complex IV protein expression (Fig. 2B). A muscle biopsy sample from the patient’s clinically-unaffected mother revealed normal histopathological (Fig. 2C) and immunohistochemical (Fig. 2D) profiles.

3.2. Mitochondrial genetic studies

No evidence of the common pathogenic m.3243A>G variant or large-scale mtDNA rearrangements was detected prior to sequencing of the entire mitochondrial genome. Analysis of the mitochondrial genome in patient muscle
revealed two novel variants: m.7887G>A p.(Gly101Asp) and m.8250G>A p.(Gly222Glu), and both at high levels of heteroplasmy (78% mutant load). The m.7887G>A variant is not listed on publicly available databases including GenBank (51,836 human mtDNA sequences interrogated on 15th Jan 2021) or our own in-house database comprising >2500 mtDNA sequences, and the m.8250G>A variant is extremely rare. The m.7887G>A variant predicts the substitution of a highly-conserved, hydrophobic glycine residue with an aspartic acid residue, whilst the m.8250G>A variant is predicted to substitute a weakly-conserved glycine residue with a glutamic acid residue (Fig. 3).

Both these variants showed lower levels of heteroplasmy in a range of tissues (urinary sediment, blood and buccal...
epithelia). The m.7887G>A variant was present at levels of 21%, 6% and 3% respectively; the m.8250G>A variant was detected at levels of 17% and 5% in urinary sediment and blood, although was undetectable (within the limits of sensitivity of the assay) in buccal epithelia derived DNA.

Single muscle fibre segregation studies in individual COX-positive and COX-deficient fibres revealed that both variants segregated to COX-deficient fibres suggesting that both variants co-exist on the same mtDNA molecule. For the m.7887G>A variant, analysis of 19 COX-positive fibres showed a mean mutation load of 7±3%, whilst 17 COX-deficient fibres showed a mean mutation load of 95±1% (unpaired t-test \( p < 0.0001 \)) and for the m.8250G>A variant, analysis of 13 COX-positive fibres showed a mean mutation load of 8±5%, whilst 17 COX-deficient fibres showed a mean mutation load of 93±1% (unpaired t-test \( p < 0.0001 \)) (Fig. 3).

Quantitative pyrosequencing of non-invasive tissue samples taken from the patient’s mother revealed the m.8250G>A variant to be present at low levels in urinary sediment and buccal epithelia-derived DNA (16% and 5% respectively) and at intermediate levels in blood (36%). However, the m.7887G>A variant was undetectable in urinary sediment, blood and buccal epithelia derived samples. She underwent a muscle biopsy at aged 72 years old; the m.7887G>A was not detectable but the m.8250G>A was present at high levels (87%); in the context of her normal muscle histopathology (Fig. 2), this strongly implicates the m.7887G>A variant as causal. Interestingly, both variants were undetectable in blood, urinary sediment and buccal-epithelia-derived DNA of the patient’s sister.

4. Discussion

We describe a 45-year-old man presenting with progressive cerebellar syndrome with sensory neuropathy with onset in his 20s. Extensive genetic screening failed to identify a genetic cause, prompting diagnostic muscle biopsy, which indicated an underlying mitochondrial aetiology. Mitochondrial genome sequencing in muscle revealed two mtDNA variants - m.7887G>A (novel) and m.8250G>A (extremely rare) –
predicting missense variants in the MT-CO2 gene, compatible with the marked mitochondrial histopathological abnormalities. Although both variants attain some measures proposed for a pathogenic mtDNA variant, our wider investigations strongly suggest that only the m.7887G>A; p.(Gly101Asp) variant completely fulfills criteria necessary to prove causality of the patient’s clinical phenotype.

First, the m.7887G>A variant is absent on large, publicly accessible mitochondrial genetic databases and m.8250G>A is extremely rare. Second, both mtDNA variants are heteroplasmic and demonstrate the highest mutant loads in clinically-affected, postmitotic skeletal muscle, with lower levels in urinary epithelia, blood and buccal cells. Third, histochemical studies of the patient’s muscle biopsy showed a dramatic loss of COX activity, with quantitative, quadruple OXPHOS immunohistochemistry showing a consistent level of loss of COXI protein expression (Fig. 2); both findings would be expected for and predicted by a pathogenic MT-CO2 variant. Fourth, single fibre segregation studies established that both variants segregate with the COX histochemical defect in skeletal muscle with higher levels of mutant load in COX-deficient fibres. However, there are two lines of additional evidence to support the pathogenicity of m.7887G>A variant. The m.7887G>A variant was found to have arisen de novo and absent in all tissues of other maternal family members investigated. There was no evidence of COX deficiency in the muscle tissue of patient’s mother who harbors a higher heteroplasmy of m.8250G>A variant and she is clinically unaffected. Moreover, the m.7887G>A variant predicts the substitution of a highly-conserved, hydrophobic glycine residue with an aspartic acid residue in a region of the COX2 protein that shows strong evolutionary conservation, whilst the m.8250G>A variant predicts the change of a poorly-conserved amino acid at the very C-terminal end of the protein (Fig. 2C).

Our case study also illustrates several diagnostic caveats to underpin the diagnosis of mtDNA disease in clinical practice. The genetic aetiology of non-syndromic ataxia with axonal neuropathy is highly heterogeneous [12,13] with a Mendelian pattern of recessive inheritance often assumed in the absence of any relevant family history. Recent studies have suggested that de novo mtDNA variants are not uncommon [14] with over half of all reported pathogenic MT-CO2 variants having arisen de novo [7]. Pathogenic MT-CO2 variants have been linked to several clinical phenotypes with different disease onset, such as myopathy with [15] or without [16] recurrent myoglobinuria, multisystem disease characterized by neurodevelopmental delay, gait disorder, cardiac involvement, retinitis pigmentosa and lactic acidosis [17,18], MELAS syndrome [19] and progressive cerebellar ataxia [7]. WES and WGS are increasingly integrated with mainstream genetic investigations in a “genetics first” approach and their abilities of detecting pathogenic mtDNA variants have been widely demonstrated [20,21]. However, the complexity of assigning pathogenicity of any mtDNA alterations is further complicated by the presence of mtDNA heteroplasmy as well as skewed tissue segregation of mtDNA variants. Low mutant heteroplasmies detected in EDTA-blood DNA samples can be inadvertently overlooked or regarded as incidental in some diagnostic pipelines. In agreement with earlier reports [6,7,22,23], muscle biopsy remains crucial in the diagnostic algorithm of adult-onset mitochondrial disease, particularly when evaluating the pathogenicity of novel mtDNA variants [24].

We identified the accumulation of medium-chain acylcarnitine species in the patient’s plasma, similar to the findings reported in other pathogenic MT-CO2 variants [7,15,16], which mimic the findings in patients with primary fatty acid oxidation disorder due to multiple acyl-CoA dehydrogenase deficiency (MADD). Abnormal acylcarnitine profiles in the fibroblasts of patients with mitochondrial respiratory chain deficiencies, resembling patients with various primary fatty acid oxidation defects, were previously reported [25]. More recently, a metabolomics study of nine patients with LRPPRC-related mitochondrial disease (deficiency in this nuclear-encoded mitochondrial protein also causes isolated complex IV deficiency) identified elevated C2, C6, C12, C14, C14:1, C16, C18:1, and C18:2 acylcarnitines, suggesting the perturbation of fatty acid oxidation pathway is likely to be secondary to mitochondrial respiratory chain dysfunction [26].

Co-segregation of two (possible) pathogenic mtDNA variants has occasionally been reported in the literature. In some cases, the co-presence of two pathogenic mtDNA variants was thought to contribute the clinical manifestations, such as identification of m.11778G>A and a single, large-scale mtDNA deletion – the so-called “common mtDNA deletion” - in a young man who with childhood-onset CPEO who later developed symptomatic, bilateral optic neuropathy [27]; co-existence of single, large-scale deletion and m.3243A>G has also been reported in a woman with Kearns-Sayre syndrome and multiple endocrine disorder [28]. The exact roles of co-segregated mtDNA variants cannot be robustly determined in some patients, leading to speculation of a synergistic, pathogenic effect or that one of the mtDNA variants may act to modulate the penetrance of the second mtDNA variant [29–32]. On the other hand, some of these variants can be classified as neutral polymorphisms retrospectively when more stringent diagnostic criteria have been applied [33–35]. Our case study is unique as we have established that only the de novo m.7887G>A variant is pathogenic, whilst the maternally-inherited m.8250G>A variant is a polymorphism through the analysis of the patient’s and his mother’s muscle biopsies. Based on these findings, we have been able to provide genetic counseling to other maternal family members.

In conclusion, we describe a novel pathogenic m.7887G>A p.(Gly101Asp) MT-CO2 gene variant in a patient who manifests with an ataxia neuropathy spectrum mimicking autosomal recessive cerebellar ataxia and highlight the continuing requirement for diagnostic muscle biopsy in cases of adult-onset mtDNA disease, particularly those necessitating the characterization of novel sequence variants [22].
Declarations of Competing Interest

Nothing to declare.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.jnd.2021.05.014.

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