Risk of cardiac manifestations in adult mitochondrial disease caused by nuclear genetic defects

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ABSTRACT

Objective Regular cardiac surveillance is advocated for patients with primary mitochondrial DNA disease. However, there is limited information to guide clinical practice in mitochondrial conditions caused by nuclear DNA defects. We sought to determine the frequency and spectrum of cardiac abnormalities identified in adult mitochondrial disease originated from the nuclear genome.

Methods Adult patients with a genetically confirmed mitochondrial disease were identified and followed up at the national clinical service for mitochondrial disease in Newcastle upon Tyne, UK (January 2009 to December 2018). Case notes, molecular genetics reports, laboratory data and cardiac investigations, including serial electrocardiograms and echocardiograms, were reviewed.

Results In this cohort-based observational study, we included 146 adult patients (92 women) (mean age 53.6±18.7 years, 95% CI 50.6 to 56.7) with a mean follow-up duration of 7.9±5.1 years (95% CI 7.0 to 8.8). Eleven different nuclear genotypes were identified: TWNK, POLG, RRM2B, OPA1, GFER, YARS2, TYP1, ETFDH, SDHA, TRIT1 and AGK. Cardiac abnormalities were detected in 14 patients (9.6%). Seven of these patients (4.8%) had early-onset cardiac manifestations: hypertrophic cardiomyopathy required cardiac transplantation (AGK; n=2/2), left ventricular (LV) hypertrophy and bifascicular heart block (GFER; n=2/3) and mild LV dysfunction (GFER; n=1/3, YARS2; n=1/2, TWNK; n=1/41). The remaining seven patients had acquired heart disease most likely related to conventional cardiovascular risk factors and presented later in life (14.6±12.8 vs 55.1±8.9 years, p<0.0001).

Conclusions Our findings demonstrate that the risk of cardiac involvement is genotype specific, suggesting that routine cardiac screening is not indicated for most adult patients with nuclear gene-related mitochondrial disease.

INTRODUCTION

Mitochondrial diseases are a group of neuro-metabolic disorders that can be caused by genetic defects in both the mitochondrial DNA (mtDNA) and nuclear DNA (nDNA). The estimated prevalence of all forms of adult mitochondrial disease is 1 in 4300, of which approximately a third of patients have nDNA pathogenic variants.1 Mitochondrial diseases are clinically heterogeneous and can affect organs with high-energy demand such as central nervous system, skeletal muscle and heart.2 The range of cardiac involvement such as hypertrophic cardiomyopathy, conduction abnormalities and sudden cardiac death are commonly reported in patients with certain forms of primary mtDNA disease.3–6 However, the spectrum of cardiac manifestation among patients with nDNA defects has only been described in isolated case reports or small case series.7

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What is already known about this subject?

► Approximately one-third of adult patients with mitochondrial disease have nuclear gene inheritance (ie, nuclear gene defect).

► Regular cardiac surveillance is currently advocated for patients with primary mitochondrial DNA disorders.

What does this study add?

► In this observational cohort study (n=146), we show that the majority of adult patients with mitochondrial disease of the nuclear genome (~90%) have normal cardiac investigations, and the risk of cardiac involvement is genotype specific.

► Fourteen patients (9.6%) developed cardiac abnormalities, but only half of them (4.8%) were attributed to the mitochondrial dysfunction with a mean follow-up of 8 years.

How might this impact on clinical practice?

► The risk of cardiac involvement is low in most adult patients with nuclear gene-related mitochondrial disease and we propose a cardiac algorithm to guide clinical practice.
routinely recommended for all patients with mitochondrial disease irrespective of the underlying genetic aetiology. A better understanding of cardiac abnormalities across different genotypes would guide clinicians to tailor cardiac surveillance to high-risk groups and instigate appropriate intervention at the earliest stage. On the other hand, patients with negligible risk of developing a cardiac disease related to the mitochondrial dysfunction can be reassured without requiring regular cardiac screening, potentially reducing the burden of healthcare cost. In this study, we sought to evaluate the cardiac phenotypes systematically in adult patients with mitochondrial disease resulting from pathogenic mutations in the nuclear genes. We also set out to determine the cardiac abnormalities that are attributed to their underlying primary mitochondrial disorders. To progress the current clinical practice towards individualised precision medicine, we aimed to identify patients who were most likely to benefit from regular cardiac screening in this study.

Methods

Study design and settings

We identified adult patients harbouring pathogenic nDNA variants from the National Health Service Highly Specialised Service for Rare Mitochondrial Disorders in Newcastle upon Tyne, UK and Mitochondrial Disease Patient Cohort (MitoCohort, UK) (REC: 13/NE/0326), between January 2009 to December 2018. Children who succumbed to fatal cardiac disease had been excluded. These patients were regularly reviewed by clinicians with expertise in mitochondrial disease every 6–24 months (AZL, AMS, GSG, RM, DMT and YSN), and their disease burden, including the degree of cardiac disease severity, was assessed objectively using the Newcastle Mitochondrial Disease Adult Scale, a validated disease rating scale. Patients had undergone regular cardiac testing (ECG and/or echocardiogram) over a similar period. The abnormal cardiac findings were reviewed by the cardiologists with expertise in mitochondrial diseases, and patients were further assessed if clinically indicated (JPB, MGDB and JO). Demographic data, clinical phenotype, molecular genetic data and blood test results were derived from the MitoCohort. Other medical co-morbidities and drug history were obtained from the clinical notes where available. The majority of genetic diagnoses were achieved via the identification of mitochondrial dysfunction in muscle biopsies (such as large number of cytochrome c oxidase deficient fibres, presence of mtDNA rearrangement/multiple deletions and/or mitochondrial respiratory chain deficiency) and Sanger sequencing of candidates genes based on the clinical phenotype. Our diagnostic algorithm for the mitochondrial disease has been described elsewhere.

Defining cardiac abnormalities in mitochondrial disease

Abnormal echocardiographic findings were reported according to the British Society of Echocardiography and European Society of Cardiology criteria. Hypertrophic cardiomyopathy was clinically recognised by a maximal LV wall thickness >15 mm in this population. The interpretation of ECG complied with consensus recommendations from professional bodies in clinical cardiology. Cardiovascular risk factors, as defined by established practice guidelines were ascertained namely, hypertension (persistent blood pressure ≥140/90 mm Hg), diabetes mellitus, hypercholesterolaemia (serum cholesterol level >5 mmol/L), obesity (body mass index >30) and smoking status. Any cardiac findings with uncertain significance were adjudicated by the cardiologists (JPB, MGDB and JO).

Patient and public involvement statement

The patients and the public were not involved in the study design, recruitment, statistical analysis and writing of this study.

Statistical analysis

Statistical analyses were performed using IBM SPSS Statistics (V.24). The incidence rate in this study was expressed as the number of cardiac abnormalities detected per number of years experienced by the cohort at risk in person-years. Binominal outcome of cardiac investigations from each genotype underwent $\chi^2$ analysis, including, Fisher’s exact test for counts <5 and parametric data were compared using analysis of variance that had been accounted for normality. Statistical significance was defined as $p \leq 0.05$.

Results

Clinical phenotypes and genotypes

We identified 146 adults with nDNA defects (92 women) who had complete clinical and cardiac data sets available for the analysis. The mean age of patients was 53.6±18.7 years (95% CI 50.6 to 56.7), and the mean follow-up (FU) duration was 7.9±5.1 years (95% CI 7.0 to 8.8). The common clinical features of mitochondrial disease were eyelid ptosis (51.4%), chronic progressive external ophthalmoplegia (CPEO) (51.4%), cerebellar ataxia (34.2%), myopathy (32.9%), gastrointestinal disturbance (22.6%) and peripheral neuropathy (22.6%) (Table 1). Eleven nuclear gene defects were identified: TWNK (n=3), POLG (n=37), RRM2B (n=24), OPA1 (n=24), GFER (n=3), YARS2 (n=2), TYP (n=3), ETFDH (n=2), SDHA (n=2), TRIT1 (n=2) and AGK (n=2). Clinical phenotypes associated with individual nuclear defects are summarised in online supplemental
Eight of the 14 patients developed cardiac diseases that were deemed unrelated to their genetic defect: POLG (n=1), TWNK (n=3) and RRM2B (n=3). The mean onset of cardiac abnormalities identified in these patients was 55.1±8.9 years (95% CI 46.9 to 63.4). Four patients had had a myocardial infarction secondary to coronary artery disease (RRM2B=3; POLG=1). The therapeutic interventions for three RRM2B patients included emergency coronary angioplasty (n=1), implantable cardioverter defibrillator for prolonged non-sustained VT and moderate LV dysfunction at age 60 years (n=1), and elective coronary artery bypass surgery (n=1). The patient with POLG-related mitochondrial disease had persistent AF refractory to multiple ablations and subsequently had an acute admission due to myocardial infarction. Two cases of TWNK mutation had asymptomatic, mild LV dysfunction without any structural or rhythmic changes at year of 10 years. A 27-year-old man with YARS2-related mitochondrial disease had progressive hypertrophic cardiomyopathy with LV systolic dysfunction. Two siblings with recessive GFER disease were identified to have LVH with preserved ejection fraction, and electrical abnormalities (bifascicular block and LBBB, respectively) in their twenties. The third patient with GFER mutation developed mild LV dysfunction without any structural or rhythmic changes at the age of 10 years. A 27-year-old man with YARS2-related mitochondrial disease had progressive hypertrophic cardiomyopathy with LV systolic dysfunction. Two siblings with recessive GFER disease were identified to have LVH with preserved ejection fraction, and electrical abnormalities (bifascicular block and LBBB, respectively) in their twenties. The third patient with GFER mutation developed mild LV dysfunction without any structural or rhythmic changes at the age of 10 years. A 27-year-old man with YARS2-related mitochondrial disease had progressive hypertrophic cardiomyopathy with LV systolic dysfunction.

Cardiac abnormalities were attributable to the underlying mitochondrial genotype in seven patients (4.8%): AGK (n=2), GFER (n=3), YARS2 (n=1) and TWNK (n=1). The mean age of cardiac manifestation was 14.6±12.8 years (95% CI 2.74 to 26.4). Both patients (unrelated) who harboured pathogenic variants in AGK presented with an infantile-onset (<1 year old) hypertrophic cardiomyopathy, which progressed to end-stage cardiac failure. Both patients subsequently required LV assist device insertions, followed by orthotopic cardiac transplants at age 13 years. One of the AGK cases also had symptomatic WPW syndrome and underwent two accessory pathway (AP) ablation procedures at age 4 years (left lateral AP) and 7 years (left posterolateral AP), respectively.

Two siblings with recessive GFER disease were identified to have LVH with preserved ejection fraction, and electrical abnormalities (bifascicular block and LBBB, respectively) in their twenties. The third patient with GFER mutation developed mild LV dysfunction without any structural or rhythmic changes at the age of 10 years. A 27-year-old man with YARS2-related mitochondrial disease had progressive hypertrophic cardiomyopathy with LV systolic dysfunction. Two siblings with recessive GFER disease were identified to have LVH with preserved ejection fraction, and electrical abnormalities (bifascicular block and LBBB, respectively) in their twenties. The third patient with GFER mutation developed mild LV dysfunction without any structural or rhythmic changes at the age of 10 years. A 27-year-old man with YARS2-related mitochondrial disease had progressive hypertrophic cardiomyopathy with LV systolic dysfunction.

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each of these genotypes (TWNK, POLG and RRM2B) were low even though they were accounted for the two-thirds of adult patients in our cohort. (Figure 1).

The frequency of individual cardiovascular risk factors is summarised in table 1. There was no significant association between specific nuclear gene defects and with any of the cardiovascular risk factors. Nineteen patients were treated with 3-hydroxy-3-methyl-glutaryl-coenzyme A reductase inhibitor (statin); none reported worsening of myopathy, myalgia or other significant adverse side effects. Moreover, there was no significant difference in creatine kinase level between the patients receiving statins (n=19; mean 263.1±249.8 units/L) and patients not receiving statins (n=70; mean 198.9±213 units/L) (p=0.265).

None of the patients with the following genetic defects developed cardiac abnormality during the FU: OPA1 (mean age 51.9±15.4 years, mean FU 9.2±4.4 years), TYMP (mean age 35.0±6.2 years, mean FU 3.7±1.5 years), ETFDH (mean age 43.5±17.7 years, mean FU 18.5±10.6 years), SDHA (mean age 82±8.5 years, mean FU 17.5±9.2 years) and TRIT1 (mean age 32.5±5.0 years, mean FU 9.5±2.1 years).

**DISCUSSION**

In this observational cohort study of adult patients with mitochondrial disease, we show that the majority of patients with nDNA pathogenic variants have normal cardiac studies. Abnormal cardiac findings were identified in approximately 10% of the patients and half of those with cardiac abnormalities had already presented before adulthood. In contrast, patients with pathogenic mtDNA variants have a significantly higher prevalence of cardiac involvement (25%–50%).3 4 19

Our findings of early-onset cardiac involvement caused by pathogenic AGK and YARS2 variants are consistent with the published data.18 20 The clinical presentation of two AGK cases exhibited classic features of Sengers syndrome21 characterised by congenital cataracts, hypertrophic cardiomyopathy, myopathy and exercise intolerance. We previously demonstrated that hypertrophic cardiomyopathy is common among patients who harbour the pathogenic YARS2 variants, in addition to other clinical findings such as lactic acidosis, proximal myopathy, respiratory muscle weakness and sideroblastic anaemia in some cases.18 The exact function of GFER

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**Figure 1** Summary of cardiac abnormalities identified. Left ventricle dysfunction defined as mild (LVEF 45%–55%), moderate (35%–45%) and severe (<35%). NMDAS cardiac score <3 indicates asymptomatic cardiac abnormalities. Other cardiac abnormalities includes conduction and repolarisation conditions. *The link between the underlying genetic with cardiac abnormalities is tentative. **The inheritance pattern is autosomal dominant for all cases of TWNK, RRM2B, OPA1 and SDHA except one RRM2B patient who has two pathogenic variants inherited as a recessive disease. †The inheritance pattern is autosomal recessive for the following nDNA defects: POLG (except one case with a single heterozygous variant inherited as a dominant disease), GFER, YARS2, TYMP, ETFDH, TRIT1 and AGK. AF, atrial fibrillation; ARB, angiotensin receptor blocker; BB, beta-blocker; BFB, bifascicular block; CABG, coronary artery bypass graft; CCB, calcium channel blocker; CAD, coronary artery disease; CM, cardiomyopathy; CVD, cardiovascular disease; DM, diabetes mellitus; Htn, hypertension; ICD, implantable cardioverter defibrillator; Lpd, dyslipidaemia; LVAD, left ventricular assist device; Ob, obesity; NMDAS, Newcastle Mitochondrial Disease Adult Scale; NSVT, non-sustained ventricular tachyarrhythmia; Sm, History of cigarette smoking; WPW, Wolff-Parkinson-White syndrome.
(Growth Factor, Augmenter of Liver Regeneration) protein is not well understood. The GFER genetic defect was initially reported in three children who presented with developmental delay, progressive myopathy, congenital cataracts and sensorineural hearing loss. In comparison, we describe three adult patients with recessive GFER variants from two unrelated pedigrees, who developed mild LV dysfunction; thus, expanding the phenotypic spectrum of GFER-related mitochondrial disease.

We identified mild LV dysfunction in one patient aged 27 years who harbours a heterozygous TWNK variant that was investigated for intermittent palpitations. However, cardiac investigations were normal in three of his family members, who have the same genetic mutation. Non-specific cardiac abnormalities were identified through routine screening in three other TWNK patients over the age of 50 years, and they have coexisting cardiovascular risk factors. There is no evidence of cardiac phenotype in the TWINKLE deleter mouse model and autosomal dominant TWNK mitochondrial disease typically associated with late-onset CPEO and mild myopathy in human.

As such, we believe that the risk of early cardiac disease is low in this patient group. Several nuclear genes involved in the mtDNA maintenance such as POLG, TWNK, RRM2B and OPA1 account for a substantial proportion of the genetic diagnoses in adult patients. Our results suggest that these common nuclear gene defects do not have an inherent cardiac phenotype apart from a single case of TWNK mutation. Based on our longitudinal data, we have proposed a clinical algorithm of cardiac screening for nDNA related mitochondrial disease in adults (figure 2). A baseline cardiac screening is recommended for any novel or ultra rare nuclear gene defects with limited natural history data on their respective cardiac involvement.

A small number of older patients with cardiovascular risk factors developed coronary artery disease similarly seen in the general population, suggesting that the conventional management of cardiovascular risk factors should be instigated accordingly. There is a theoretical concern that using statins could exacerbate myopathy in patients with neuromuscular and mitochondrial disease.

Figure 2 Clinical algorithm to guide cardiac surveillance for the adult mitochondrial disease of nuclear gene defects. This algorithm is proposed guidance and is not intended to replace the clinical judgement of healthcare professionals in the context of individual circumstances. The clinical guidelines for mitochondrial DNA mutations have been published. Clinical team refers to clinicians who regularly follow-up the patients with mitochondrial disease, and they include general neurologists, clinical geneticists, metabolic medicine physicians and general practitioners/primary care physicians. CK, creatine kinase; CVS, cardiovascular system.
However, we have not observed any significant adverse effect of statin use in our patients. We have not been able to determine the risk of cardiac disease in rarer nuclear gene defects associated with the mtDNA maintenance such as MGME1, DNA2, SLC25A4 (also known as ANT1), DGUK and MPV17. The published literature has not revealed that cardiac involvement is a consistent finding in these patients who presented with CPEO and multiple mtDNA deletions on muscle biopsy. However, robust systematic evaluation of the risk of cardiac disease in these genes is warranted to inform clinical practice. A few recent studies proposed that cardiac surveillance for mitochondrial patients can be stratified based on the clinical syndrome or phenotype alone. However, such recommendations have limitations, given that the phenotype–genotype correlation is not always evident. For instance, CPEO and myopathy are common clinical features seen in adult patients with mitochondrial disease and the genetic causes are highly heterogeneous, which include primary mtDNA point mutations such as m.3243A>G, single, large-scale mtDNA deletions or multiple mtDNA deletions secondary to nuclear gene defects as outlined previously. If cardiac surveillance were to perform in all patients with CPEO or myopathy, a significant portion of patients with nDNA pathogenic would subject for unnecessary investigations and hospital visits.

There are several limitations in this study. First, we only capture three nuclear genes associated with an early-onset cardiac phenotype in this cohort, and the patients have survived into adulthood. However, there are ultra-rare nuclear genetic defects, such as aminoacyl-tRNA synthetase that can cause fatal cardiac disease in childhood. Second, most of our patients did not have cardiac MR imaging performed. Cardiac MR has a higher sensitivity for the detection of myocardial changes ahead of ventricular dysfunction over routine transthoracic echocardiography. However, the majority of our patients had serial echocardiograms over long FU to reduce the likelihood of any clinically significant pathology. Third, we only studied the dominant form of mitochondrial disease for these genes: TWNK, RRM2B, OPA1 and SDHA. We had not recruited patients with the autosomal recessive disease who typically have a more severe phenotype, frequently manifesting in childhood and much less prevalent in the population. Finally, the numbers of patients with specific genotypes (eg, TRIT1 and SDHA) are small in our cohort, and an international, multicentre study would be necessary to establish their risk of cardiac involvement definitively.

CONCLUSIONS

The overall risk of early cardiac involvement in adult mitochondrial disease caused by nuclear gene defects is low (4.8%), and is genotype specific. Our study provides evidence that routine cardiac screening is not indicated for most adult patients with nuclear gene-related mitochondrial disease.

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Contributors DMT, RM, GSG and YSN conceived the idea and designed the study. AZL, DMU and YSN carried out the data collection and performed the analysis. MGDB, AMS, JG, CF, MEF, JPB, DMT, RM, GSG and YSN had provided substantial acquisition of data and interpretation. JG, JPB and MGDB gave critical feedback on the cardiac data. AZL and YSN wrote the first version of the manuscript. All authors critically reviewed the manuscript and approved the final version of this manuscript.

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Competing interests None declared.

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