GENERAL ARTICLE

Instability of the mitochondrial alanyl-tRNA synthetase underlies fatal infantile-onset cardiomyopathy

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Abstract

Recessively inherited variants in AARS2 (NM_020745.2) encoding mitochondrial alanyl-tRNA synthetase (mt-AlaRS) were first described in patients presenting with fatal infantile cardiomyopathy and multiple oxidative phosphorylation defects. To date, all described patients with AARS2-related fatal infantile cardiomyopathy are united by either a homozygous or compound heterozygous c.1774C>T (p.Arg592Trp) missense founder mutation that is absent in patients with other AARS2-related phenotypes. We describe the clinical, biochemical and molecular investigations of two unrelated boys presenting with fatal infantile cardiomyopathy, lactic acidosis and respiratory failure. Oxidative histochemistry showed cytochrome c oxidase-deficient fibres in skeletal and cardiac muscle. Biochemical studies showed markedly decreased activities of mitochondrial respiratory chain complexes I and IV with a mild decrease of complex III activity in skeletal and...
cardiac muscle. Using next-generation sequencing, we identified a c.1738C>T (p.Arg580Trp) AARS2 variant shared by both patients that was in trans with a loss-of-function heterozygous AARS2 variant; a c.1008dupT (p.Asp337*) nonsense variant or an intragenic deletion encompassing AARS2 exons 5–7. Interestingly, our patients did not harbour the p.Arg592Trp AARS2 founder mutation. In silico modelling of the p.Arg580Trp substitution suggested a deleterious impact on protein stability and folding. We confirmed markedly decreased mt-AlaRS protein levels in patient fibroblasts, skeletal and cardiac muscle, although mitochondrial protein synthesis defects were confined to skeletal and cardiac muscle. In vitro data showed that the p.Arg580Trp variant had a minimal effect on activation, aminocacylation or misaminocacylation activities relative to wild-type mt-AlaRS, demonstrating that instability of mt-AlaRS is the biological mechanism underlying the fatal cardiomyopathy phenotype in our patients.

Introduction

Mitochondrial respiratory chain disorders are among the most common early onset metabolic disorders with an estimated minimum prevalence of 1 in 5000 live births (1). Isolated or multiple deficiencies of the five multimeric complexes (I–V) that comprise the oxidative phosphorylation (OXPHOS) system are associated with broad clinical, biochemical and genetic heterogeneity. Disorders of mitochondrial mRNA translation or protein synthesis are especially important causes of multiple mitochondrial respiratory chain deficiency, which are linked to both mitochondrial DNA (mtDNA) and nuclear gene defects (2).

Following post-transcriptional modification, a critical step of mitochondrial protein synthesis is the aminocacylation or ‘charging’ of transfer RNAs (tRNAs) (3). This step involves the recognition and conjugation of amino acids with their corresponding cognate mitochondrial transfer RNA (mt-tRNA), as dictated by the codon sequence. Attachment is catalysed by mitochondrial aminoacyl-tRNA synthetases (mt-aaRS) that are encoded by nuclear genes and imported into mitochondria. There are 17 mt-aaRS and two dual cytosolic-mitochondrial synthetases (GlyRS, LysRS), while mt-GluRS is required to efficiently misaminocylate tRNA\textsuperscript{Gln} to form Glu-tRNA\textsuperscript{Gln} in mitochondria (4,5).

All mt-aaRS and dual-localized synthetases are associated with autosomal recessive human disorders manifesting in clinically and biochemically heterogeneous phenotypes (6–8). Despite ubiquitous expression, autosomal recessive mt-aaRS disorders are associated with intriguing tissue- and cell-specific phenotypes that typically involve the central nervous system (9–18). High-throughput, next-generation sequencing technologies have greatly expanded the phenotypic continuum of mt-aaRS disorders to encompass patients presenting with additional clinical features or with the absence previously considered salient features.

Recessively inherited variants in AARS2 (NM_020745.2), encoding mitochondrial alanyl-tRNA synthetase (mt-AlaRS), were first described in patients presenting with fatal infantile cardiomyopathy and multiple OXPHOS defects (19), with additional patients subsequently identified (20–24). However, the spectrum of AARS2-related disease has expanded to include childhood to adulthood-onset leukoencephalopathy with premature ovarian failure (POF) in females (9,25–28), retinopathy and optic atrophy (29) and fatal non-immune hydrops fetalis (30); all with conspicuous absence of cardiac involvement. Currently, AARS2-related fatal infantile cardiomyopathy is associated with a recurrent pathogenic c.1774C>T (p.Arg592Trp) founder mutation that is either homozygous or compound heterozygous in all described patients. This founder mutation has not been reported in patients presenting with other AARS2-related phenotypes. Consequently, the spectrum of AARS2-related disease phenotypes has been attributed to the location of pathogenic variants in the protein and the effect on protein function (22). It has been previously hypothesized that the p.Arg592Trp AARS2 founder mutation, which occurs in a conserved editing domain, causes a severe decrease in aminocacylation due to impaired tRNA binding and positioning of the 3′-end within the active site (22). On the other hand, other AARS2-related disease phenotypes were predicted to result from only a partial reduction in aminocacylation activities (22). This mt-AlaRS editing domain is required for the deacylation of mischarged tRNAs, since the aminocacylation domain is unable to discriminate alanine with serine and glycine (31,32). This proofreading activity is essential to clear mischarged Ser-tRNA\textsuperscript{Ala} and avoid misincorporation of serine at alanine codons, since a slight decrease results in embryonic lethality in mice (33). Of all mt-aaRS, only mt-AlaRS and mt-ThrRS have demonstrable editing activities to prevent the formation of mischarged mt-tRNAs (31,32,34).

In this study, we describe two unrelated patients presenting with fatal infantile cardiomyopathy, lactic acidosis and respiratory failure, with severe multiple OXPHOS deficiency and who both harboured an unreported AARS2 variant (c.1738C>T, p.Arg580Trp) in trans with a loss-of-function AARS2 variant, but not the recurrent p.Arg592Trp founder mutation. We validate pathogenicity of this shared novel mt-AlaRS editing domain variant through post-mortem molecular studies, in silico modelling and in vitro assays. This data supports the genotype-phenotype correlation between AARS2 variants in the β-barrel domain with fatal cardiomyopathy and that instability of mt-AlaRS due to the novel p.Arg592Trp and loss-of-function alleles is the underlying biological mechanism in our patients.

Results

Case reports

Patient 1. Patient 1 was a male infant born at term by normal vaginal delivery to non-consanguineous parents with a birth weight of 3.11 kg. He developed respiratory distress and poor respiratory drive soon after birth, requiring ventilation. He had generalized hypotonia and evidence of diaphragmatic paralysis with paradoxical abdominal wall movements. There was persistent lactic acidemia (9–30 mmol/L; normal <2.5 mmol/L), Urine organic and amino acids were unremarkable apart from increased lactate excretion. There were no seizures and he tolerated nasogastric feeding. Initial echocardiography showed no evidence of cardiomyopathy, but he developed mild biventricular hypertrophy by 5 weeks of age. This was associated with periods of cardiac electrical inactivity, lasting 6–7 s and later up to 30 s. He was weaned off ventilatory support at 6 weeks and died 1 week later. Whole mitochondrial genome sequencing failed to detect a pathogenic variant, while quantitative real-time polymerase chain reaction (PCR) assay of skeletal muscle mtDNA copy number was normal (data not shown). A previous daughter...
was born at term and died within 24 h with lactic acidemia and coagulopathy. Post-mortem analysis of this female sibling is said to have shown pulmonary hypoplasia. There is one healthy son and the child’s mother had two previous miscarriages.

**Patient 2.** Patient 2, a male infant, was a dizygotic twin born at 33 weeks gestation by caesarean section to non-consanguineous parents with a birth weight of 1.595 kg. He presented at 2 months of age with respiratory failure secondary to respiratory syncytial virus bronchiolitis. An echocardiogram revealed severe concentric left ventricular hypertrophy and dilation and severe systolic dysfunction. (Fig. 1D) Brain magnetic resonance imaging noted a thin corpus callosum, but neurological examination was normal. He required ongoing mechanical ventilation for respiratory failure. He received two courses of venoarterial extracorporeal membrane oxygenation for circulatory support and was transitioned to a Berlin left ventricular assist device at 5 months of age. Aside from persistent lactic acidemia (2–16 mmol/L; normal <2.5 mmol/L), extensive biochemical evaluation was unremarkable. He developed multi-organ failure and suffered a left middle cerebral artery stroke with residual neurologic dysfunction and muscular weakness at 6 months of age. Palliative care was initiated and he died at 7 months of age. Post-mortem evaluation demonstrated a markedly enlarged globular heart with biventricular hypertrophy and severe myocyte vacuolization. Negative genetic evaluations included karyotype, single nucleotide polymorphism (SNP) microarray, urine mitochondrial genome analysis and a targeted 89-gene cardiomyopathy panel. The patients’ male twin was diagnosed with Trisomy 18 prenatally and died soon after birth with significant cardiomegaly noted post-mortem. Parental echocardiograms were normal.

**Diagnostic histochemical and biochemical analyses of skeletal and cardiac muscle reveal severe multiple mitochondrial OXPHOS defects**

Histopathologic analysis of skeletal muscle from Patient 1 showed vacuolated fibres with increased lipid. Oxidative enzyme histochemistry showed absent c oxidase (COX) activity in 50% of fibres (Fig. 1A), although an assessment of sequential COX–succinate dehydrogenase (SDH) histochemistry was not made. In Patient 2, oxidative enzyme histochemistry of post-mortem skeletal and cardiac muscle revealed global COX-deficiency (Fig. 1A). Post-mortem histopathologic analysis of the heart noted marked biventricular myocyte vacuolization, myocyte hypertrophy and mild subendocardial fibrosis (Fig. 1Bi), compared to an age-matched control (Fig. 1Bii). Electron microscopy of cardiac muscle showed no ultrastructural evidence for a mitochondrial disorder but demonstrated vacuolar myopathic changes with large membrane bound vesicles containing glycogen.

Biochemical analysis of mitochondrial respiratory chain complex activities (Fig. 1C) revealed markedly decreased complex I and complex IV activities with low complex III activity in Patient 1 and Patient 2 skeletal muscle relative to controls. Similarly, severe complex I and complex IV activities with low complex III activity were also noted in the cardiac muscle from Patient 2, relative to age-matched controls.

**Identification of recessively inherited AARS2 variants**

In Patient 1, analysis of whole exome sequencing (WES) called variants in nuclear genes encoding mitochondriallocalized proteins revealed two heterozygous variants in AARS2 (NM_020745.2): c.1008dupT, (p.Asp337*) and c.1738C>T (p.Arg580Trp), which were confirmed by Sanger sequencing (Fig. 2A). Unfortunately, familial segregation studies were not possible. In Patient 2, whole genome sequencing (WGS) revealed the identical heterozygous c.1738C>T (p.Arg580Trp) AARS2 missense variant that was paternally inherited and confirmed by Sanger sequencing (Fig. 2A). Visual inspection of read alignments and using iGVG (35) also identified a maternally inherited, intragenic 4.1 kb deletion on the short arm of chromosome 6p21.1 encompassing exons 5–7 of AARS2, which was confirmed to segregate with the disease by long-range PCR (Fig. 2B).

None of the identified variants were previously reported as pathogenic and both patients did not harbour the p.Arg592Trp AARS2 founder mutation. In GnomAD, the c.1008dupT(p.Asp337*) variant was present in 3/245814 (minor allele frequency (MAF) = 1.22 × 10−5) alleles and the c.1738C>T (p.Arg580Trp) variant was present in 9/246174 (MAF = 3.656 × 10−5) alleles, all in heterozygous state. The p.Arg580Trp variant was predicted to be damaging by the in silico tools PolyPhen-2 (HumDiv Score 0.996) (36) and SIFT (Score 0.02) (37) but was predicted to be benign by Align GVG (Class 0, GV: 127.27, GD: 46.61) (38).

According to the American College of Medical Genetics guidelines for characterization of sequence variants, the novel AARS2 p.Arg580Trp variant did not meet the criteria for classification as ‘likely pathogenic’ variant. Therefore, we decided to pursue functional analyses to confirm pathogenicity of the identified AARS2 variants.

**Structural modelling of the p.Arg580Trp AARS2 variant**

We first examined the sequence conservation of the Arg580 residue and modelled the p.Arg580Trp variant using the available structural model for mt-AlaRS (22). The Arg580 residue is conserved in this position among mt-AlaRS in mammals and birds but not in the lizard Anolis carolinensis, fish Danio rerio, fly Drosophila melanogaster, worm Caenorhabditis elegans or yeast Saccharomyces cerevisiae (Fig. 3A). However, a bulky aromatic residue in the corresponding position is not found in any of the analysed mt-AlaRS sequences. Arg580 is one of the solvent exposed residues on the surface of the β-barrel subdomain (530–621 aa) of the mt-AlaRS editing domain and is involved in complex electrostatic, hydrophobic and hydrogen interactions with neighbouring residues. This suggests a structural role for Arg580 and an impact on protein folding and stability. Substitution of an arginine to a hydrophobic and bulky tryptophan is predicted to affect folding of the β-barrel subdomain and as a result compromise the stability of the entire protein (Fig. 3B).

mt-AlaRS protein levels are diminished in patient fibroblasts without defective mitochondrial protein synthesis

We assessed steady-state mt-AlaRS and OXPHOS complex subunit protein levels in fibroblast lysates from Patient 1 and two patients harbouring the p.Arg592Trp AARS2 founder mutation on at least one allele (Fig. 4A). Cultured fibroblasts were not available from Patient 2. Quantification of steady-state levels of mt-AlaRS showed a statistically significant decrease in all patients (Fig. 4B). However, there was no change in OXPHOS complex subunit levels (Fig. 4A). Levels of mt-tRNAAla and the presence of uncharged and charged species were also assessed in patient
Figure 1. Histochemical and biochemical studies of AARS2 patient skeletal and cardiac muscle. (A) Diagnostic skeletal and cardiac muscle were subjected to COX, SDH and sequential COX-SDH histochemical reactions. Skeletal muscle from Patient 1 was not subjected to sequential COX-SDH histochemistry. (B) Photomicrograph of (i) Patient 2 cardiac muscle sampled at autopsy and (ii) healthy heart of a child of similar age. (C) Measurement of mitochondrial OXPHOS activities (CI-CV) normalized to CS in skeletal (Patient 1 and 2) and cardiac muscle (Patient 2), as a percentage of residual controls. Controls are denoted in blue, Patient 1 in red and Patient 2 in orange. Decreased OXPHOS activities are denoted by asterisks (**). (D) Echocardiographic images in the parasternal short axis view of Patient 2’s heart (i and ii; annotated in panel ii) and the normal heart of a child approximately the same age (iii). The patient’s heart demonstrates severe concentric hypertrophy of the left ventricle (LV) involving both the interventricular septum (IVS) and the posterior wall. Also identified is the right ventricle (RV).

Patient skeletal and cardiac muscle have decreased mt-AlaRS protein levels and mitochondrial protein synthesis defects

Next, we examined steady-state mt-AlaRS and OXPHOS subunit protein levels in skeletal and cardiac muscle homogenates from Patient 2 (Fig. 5A). No skeletal or cardiac muscle was available from Patient 1. Quantification of steady-state mt-AlaRS protein levels showed a statistically significant decrease in both skeletal and cardiac muscle homogenate from Patient 2; mt-AlaRS levels were undetectable in cardiac homogenate (Fig. 5B). There was marked loss of MT-COI (complex IV) and NDUFB8 (complex I) subunits with a mild reduction of UQCRC2 (complex III). This was consistent with the decreased biochemical activities for complexes I, III and IV in skeletal and cardiac muscle (Fig. 1C). We also quantified AARS2 mRNA levels in cardiac muscle, which confirmed a 50% decrease in AARS2 mRNA relative to controls (data not shown). We then assessed mt-tRNAAla levels in cardiac muscle from Patient 2 as well as an unrelated patient who was homozygous for the p.Arg592Trp founder mutation. Northern blot analysis showed decreased levels of uncharged mt-tRNAAla in both patients (Fig. 5C). Unfortunately, the presence of charged and uncharged mt-tRNAAla could not be assessed.

The p.Arg580Trp AARS2 variant likely has a minimal effect on mitochondrial protein synthesis

We assessed the aminoacylation and editing activities of mt-AlaRS with the p.Arg580Trp variant in vitro. ATP-Pi exchange reaction assay showed that the p.Arg580Trp variant exhibited the same amino acid activation activity compared to wild-type mt-AlaRS, suggesting that there is no direct impact on the synthetic active site (Fig. 6A). Next, an aminoacylation assay showed that the p.Arg580Trp variant had comparable tRNAAla charging activity compared to wild-type human mt-AlaRS, suggest-
Figure 2. Genetic analysis of identified AARS2 variants. (A) Family pedigrees showing Sanger sequencing confirmation of the c.1008dupT (p.Asp337*) and c.1738C>T (p.Arg580Trp) AARS2 variants for Patient 1 and segregation of the c.1738C>T (p.Arg580Trp) variant for Patient 2. (B) Long-range PCR confirmation of a maternally inherited, heterozygous intragenic 4.1 kb deletion on the short arm of chromosome 6p21.1 encompassing exons 5–7 of AARS2. A non-specific product at 2 kb does not affect segregation analysis. The wild-type allele (5 kb) and the deleted allele (1 kb) are denoted by a solid red arrow.

Discussion

Autosomal recessive AARS2 variants were first described in patients with fatal infantile cardiomyopathy (19). Recently, the clinical spectrum has expanded to include childhood and adult-onset leukodystrophy (with POF in females) (9), retinopathy and optic atrophy (29) and fatal non-immune hydrops fetalis (30). However, cardiomyopathy is conspicuously absent in more recently described phenotypes. Currently, there are 14 reported patients presenting with AARS2-related fatal infantile cardiomyopathy (19,20,22–24) and one patient who died in utero with myopathy, hypotonia and multiple fractures (21). These reported patients all harbour at least one copy of the recurrent p.Arg592Trp missense founder mutation.

In contrast, we report two unrelated patients presenting with fatal infantile cardiomyopathy, lactic acidosis and respiratory failure, with severe multiple OXPHOS defects and who harboured biallelic AARS2 variants but not the recurrent founder allele. Instead, our patients both harboured a novel p.Arg580Trp missense variant that was compound heterozygous with a second, loss-of-function AARS2 variant. Nonetheless, the clinical presentation of our patients was severe and broadly comparable to the previously reported patients harbouring the p.Arg592Trp founder mutation. In contrast, we report two unrelated patients presenting with fatal infantile cardiomyopathy, lactic acidosis and respiratory failure, with severe multiple OXPHOS defects and who harboured biallelic AARS2 variants but not the recurrent founder allele. Instead, our patients both harboured a novel p.Arg580Trp missense variant that was compound heterozygous with a second, loss-of-function AARS2 variant. Nonetheless, the clinical presentation of our patients was severe and broadly comparable to the previously reported patients harbouring the p.Arg592Trp founder mutation on at least one allele. For all patients, the clinical course was fatal before 1 year of life with onset of cardiac features either before or shortly after birth.

Mt-AlaRS is one of two reported mitochondrial synthetases that have editing activities to prevent the formation of mischarged mt-tRNAs (31,32,34). The conserved editing domain of mt-AlaRS is essential since the aminoacylation domain cannot discriminate alanine with serine and glycine, thus avoiding the misincorporation of serine at alanine codons and clearing mischarged Ser-tRNAAla (31,32). Both p.Arg580Trp and p.Arg592Trp variants are located in the β-barrel subdomain of the mt-AlaRS editing domain and involve an arginine to...
tryptophan substitution, while a β-barrel subdomain variant (c.1616A>G, p.Tyr539Cys) has also been reported in trans with the p.Arg592Trp founder mutation (24). However, the mutated residues are not located in the editing core of mt-AlaRS. Previous in silico modelling showed that the mt-AlaRS Arg592 residue is surface exposed and forms one salt bridge with Glu567, which is invariant in this position among cytosolic, mitochondrial and bacterial homologues of the enzyme. The p.Arg592Trp founder mutation was predicted to have a deleterious effect on tRNA binding and a severe reduction in aminoacylation activity of mt-AlaRS but with no effect on editing activity or protein stability (22). Similarly, Arg580 is also solvent exposed but in contrast to Arg592, it forms a number of different non-covalent bonds with neighbouring residues that suggests importance of arginine in this position for protein stability (Fig. 3). Hence, substitution of the Arg580 residue with tryptophan is predicted to compromise mt-AlaRS protein folding and stability.

Indeed, quantification of mt-AlaRS protein levels confirmed a statistically significant decrease in fibroblasts, skeletal and cardiac muscle from both patients harbouring the p.Arg580Trp variant and in fibroblasts from two patients harbouring at least one copy of the p.Arg580Trp allele (Fig. 4A and B). Furthermore, mt-AlaRS protein levels appeared to be decreased more in fibroblasts from Patient 1, who harboured the p.Arg580Trp allele, compared to two patients with at least one copy of the p.Arg580Trp founder mutation. Hence, this supports our in silico modelling of the p.Arg580Trp missense change. Decreased mt-AlaRS protein levels were accompanied by severe multiple OXPHOS defects in patient skeletal and cardiac muscle (Fig. 5A) but were absent in patient fibroblasts (Fig. 4A). This suggests that skeletal and cardiac muscle are more susceptible to mitochondrial protein synthesis defects, while mt-AlaRS activity is essential in cardiac cells during early development. Our in vitro studies of mutant p.Arg580Trp mt-AlaRS also showed decreased protein stability when overexpressed in HEK293T cells (Fig. 6F). Decreased mt-tRNA\textsubscript{Ala} levels were also observed in patient cardiac muscle (Fig. 5C), confirming a deleterious effect on mitochondrial protein synthesis. However, our in vitro data clearly showed that the p.Arg580Trp variant had little effect on amino acid activation and aminoacylation activities (Fig. 6), which was expected since it is not located near the aminoacylation active site. These data also showed that the p.Arg580Trp variant did not accumulate more mischarged tRNA\textsubscript{Ala} in misaminoacylation when compared to wild-type human mt-AlaRS, since the residue...
Figure 4. Western blot, northern blot and aminoacylation analysis in AARS2 patient fibroblast lysates. (A) Steady-state mt-AlaRS and OXPHOS subunit protein levels in Patient 1 (P1) and two control fibroblast lysates. Also shown are two reported AARS2 patients in the literature who were homozygous or compound heterozygous for the p.Arg592Trp founder mutation; Patient x (homozygous p.Arg592Trp) corresponds to Patient 11, and Patient y (p.Arg592Trp and c.2882C>T (p.Ala961Val)) corresponds to Patient 7 in (24). Antibodies against mt-AlaRS, NDUF88 (C1), SDHA (CII), UQCRCC2 (CIII), MT-COI (CIV) and ATP5B (CV) were used, with β-actin as a loading control. (B) Graph of relative mt-AlaRS protein levels (n = 3) in controls, Patient 1 (P1), Patient x (x) and Patient y (y) fibroblasts. All data were normalized to SDHA and represented as mean ± standard error of the mean. Significant difference between controls and patient fibroblasts is indicated by asterisks above the columns (*, P-value < 0.05; **, P-value < 0.01 by two-tailed paired students t-test). (C) Northern blot analysis of mt-tRNAAla levels in patient fibroblasts. (D) Aminoacylation assay showing aminoacylated (‘charged’) and deacylated (‘uncharged’) mt-tRNAAla in patient fibroblasts. Mitochondrial tRNAArg was used as a loading control. Lower bands in the dAc lanes denote fully deacylated control tRNA species. Patient z was homozygous for the p.Arg592Trp founder mutation and has been previously reported, corresponding to Patient 1 in (19).

is distant from the editing active site (∼40 Å). Moreover, it is also not in the potential tRNA entrance pathway into the editing active site during tRNA 3′-end translation. Taken together, our data suggest that the p.Arg580Trp variant impacts on stability of mt-AlaRS protein but not aminoacylation or editing activities and that the β-barrel subdomain has a critical role in protein folding and stability. The combination of the p.Arg580Trp variant in trans with a second, heterozygous loss-of-function allele as detected in our patients, suggests that marked loss of mt-AlaRS is sufficient to cause a severe defect of mitochondrial protein synthesis that manifests as fatal infantile cardiomyopathy. Although skeletal and cardiac muscle appear more susceptible to mitochondrial protein synthesis defects than other cell types such as fibroblasts, data suggest that specific loss of mt-AlaRS during early development has a highly detrimental effect on cardiac cells. By contrast, loss of other mt-aaRS does not affect cardiac function during early life. For example, a loss of mt-GluRS protein due to recessively inherited EARS2 variants leads to a severe, lethal neonatal leukoencephalopathy with thalamus and brainstem involvement and high lactate (39), but without cardiac involvement. Our findings continue to support the hypothesis that there is an enhanced requirement for mitochondrial protein synthesis in the heart in early life and that a 50% reduction of aminoacylation activity is sufficient to maintain mitochondrial translation (22). On the other hand, recessively inherited variants causing other AARS2-related phenotypes are predicted to cause only a partial loss of mt-AlaRS protein level or aminoacylation activity, manifesting in childhood or adulthood phenotypes but not in the heart (22). However, steady-state mt-AlaRS protein levels have not yet been assessed in tissue from patients with other AARS2-related disorders. Furthermore, additional in vitro studies of the p.Arg592Trp founder mutation are necessary to confirm any impact on amino acid activation, aminoacylation and misaminoacylation activities. This would determine whether p.Arg580Trp and p.Arg592Trp have a shared biological mechanism or if disturbed tRNA binding with a severe loss of aminoacylation activities or marked instability of mt-AlaRS are two mechanisms that manifest clinically with fatal cardiomyopathy, concurring with in silico models.

Overall, our data strengthen the importance of mt-AlaRS in cardiac muscle during early embryonic development and the relationship between β-barrel subdomain variants (p.Arg580Trp and p.Arg592Trp) and the manifesting AARS2-related mitochondrial disease phenotypes.

Materials and Methods

Ethical compliance and informed consent

Informed consent for diagnostic and research-based studies was obtained for all subjects in accordance with the Declaration of Helsinki protocols and approved by local institutional review boards.
Histopathology, biochemical and molecular studies

Diagnostic skeletal muscle biopsies from both patients, endomyocardial biopsy and post-mortem cardiac muscle from Patient 2 were processed and mounted on glass slides according to standard procedures. Skeletal muscle biopsies (Patient 1 and 2) and post-mortem cardiac muscle (Patient 2) were subjected to cytochrome COX, SDH and sequential COX-SDH histochemical reactions (40). Mitochondrial OXPHOS activities (complexes I–IV) relative to citrate synthase (CS) were measured in skeletal (Patient 1 and 2) and cardiac muscle (Patient 2) homogenates as previously described (41). Whole mitochondrial genome sequencing of both patients was performed to exclude pathogenic variants. Quantitative real-time PCR assay of skeletal (Patient 1 and 2) and cardiac muscle DNA (Patient 2) was performed to assess mtDNA copy number, according to standard protocols.

Next-generation sequencing and genetic investigations

WES, filtering and candidate variant analysis was performed for Patient 1 as described previously (39). In Patient 2, bidirectional sequence from WGS was prepared using the Kapa Hyper library prep omitting PCR, sequenced using the Illumina HiSeq 2500 system utilizing paired end 2 × 125 base pair reads with v4 Chemistry, aligned to reference gene sequences based on human genome build GRCh37/UCSC hg19 and variants were analysed using custom-developed software; RUNES and VIKING (42,43). WES was performed on unaffected parents of Patient 2. Patient 2 was sequenced to a depth of 115.52 Gb for a mean coverage of ∼37x. Variants were filtered with a MAF less than 1% and then prioritized by the American College of Medical Genetics categorization. Align GVGD (http://agvgd.hci.utah.edu/agvgd_input.php) (38), SIFT (http://sift.jcvi.org/) (37) and PolyPhen-2 (http://genetics.bwh.harvard.edu/pph2/) (36) were used to assess pathogenicity of missense variants. Identified candidate variants were confirmed by Sanger sequencing. Long-range PCR of the AARS2 (NM_020745.2) gene was performed using forward (5′-GTGGGGTCAGCCCTGTTCCT-3′) and reverse (5′-CAGGAAGGTTGCTGTTGCT-3′) primers.

Structural modelling

Structure prediction for human mt-AlaRS with bound tRNA\textsubscript{Ala} and alanyl-adenylate in the aminoacylation site was done as earlier described (22). Briefly, a multiple sequence alignment of different cytoplasmic, mitochondrial and bacterial homologues of human mt-AlaRS was done using Promals3D server. The resulting alignment was submitted to SWISS-MODEL server. As a template for human mt-AlaRS structure modelling, the full-length AlaRS from Archaeoglobus fulgidus was used (PDB id 3WQY chain A) (44). Docking of the tRNA and alanyl-adenylate into the
Figure 6. In vitro studies of the human mt-AlaRS p.Arg580Trp mutant. (A) ATP-Pi exchange determination of human mt-AlaRS (●) and the p.Arg580Trp mutant (■). A reaction at the absence of Ala was included for a control (▲).

(B) Aminoacylation activity of human mt-AlaRS (●) and the p.Arg580Trp mutant (■). (C) A representative TLC showing the mischarging of mt-tRNAAla by human mt-AlaRS and the p.Arg580Trp mutant. (D) Graph of the mischarging activity of human mt-AlaRS (●) and the p.Arg580Trp mutant (■).

Northern blotting
Total RNA was extracted from cultured fibroblasts and cardiac muscle using Trizol reagent (ThermoFisher Scientific) according to the manufacturer's instructions. To preserve the aminoacetylation state the final RNA pellet was re-suspended in 10 mM NaOAc at pH 5.0. To investigate the aminoacylation status of mt-tRNAs, RNA (4 μg) was separated on long (16 cm length) 6.5% polyacrylamide gel (19:1 acrylamide:bis-acrylamide) containing 8 M urea in 0.1 M NaOAc, pH 5.0. The control of fully decacylated tRNA (dAc) was obtained by incubation of control RNA at 75°C (pH 9.0) for 15 min. To determine mt-tRNAAla steady-state levels the samples were run on 10 cm gel. Northern hybridization was performed with γ-32P labelled oligonucleotide probes: 5′-GTGGCTGATATTGCGTTCAGT-3′ for the mt-tRNAAla, 5′-GAGTCGAAATCATTCGTTTTG-3′ for the mt-tRNAArg and 5′-GTTGTTAGACATGGGGGCAT-3′ for mt-tRNASer(AGY). Radioactive signal was detected by PhosphorImager plate using Typhoon scanner and quantified with the Image-Quant v5.0 software (GE Healthcare).

Cloning, gene expression and protein purification of human mt-AlaRS
Gene expression and protein purification of human mt-AlaRS and the p.Arg580Trp mutant were performed as earlier described (33). The host strain used was Escherichia coli Rosetta (DE3).

Transcription of human mt-tRNAAla
Transcription of human mt-tRNAAla and 32P-labelling of human mt-tRNAAla were performed as earlier described (33).

ATP-Pi exchange assay
ATP-Pi exchange assay of human mt-AlaRS and the p.Arg580Trp mutant was performed as earlier described (33). The reaction buffer contained 50 mM Tris-HCl (pH 8.0), 20 mM KCl, 10 mM MgCl2, 2 mM DTT, 4 mM ATP, 5 mM Ala, 2 mM tetrasodium [32P]pyrophosphate and 200 nM enzyme at 37°C.

Aminoacylation activity
Aminoacylation assay of human mt-tRNAAla and the p.Arg580Trp mutant was performed as earlier described (33). The reaction buffer contained 50 mM Tris-HCl (pH 8.0), 20 mM KCl, 10 mM MgCl2, 2 mM DTT, 4 mM ATP, 1 M Ser, 4 μM human mt-tRNAAla, 0.455 μM [32P]tRNAAla (54 000 cpm) and 2 μM human mt-AlaRS and the p.Arg580Trp mutant at 37°C.
Misaminoacylation assay

Misaminoacylation assay was performed in a reaction buffer containing 50 mM Tris-HCl (pH 8.0), 20 mM KCl, 10 mM MgCl₂, 2 mM DTT, 4 mM ATP, 1 mM Ser, 4 μM human mt-tRNAAla, 0.455 μM [³²P]tRNAAla (54 000 cpm) and 2 μM human mt-AlaRS and the relative level of charged tRNAAla in the aliquots: [Ser-[³²P]AMP/(Ser-[³²P]AMP + [³²P]AMP)].

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