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# The phenotypic spectrum of germline *YARS2* variants: from isolated sideroblastic anemia to mitochondrial myopathy, lactic acidosis and sideroblastic anemia 2

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## ABSTRACT

**Y***ARS2* variants have previously been described in patients with myopathy, lactic acidosis and sideroblastic anemia 2 (MLASA2). *YARS2* encodes the mitochondrial tyrosyl-tRNA synthetase, which is responsible for conjugating tyrosine to its cognate mt-tRNA for mitochondrial protein synthesis. Here we describe 14 individuals from 11 families presenting with sideroblastic anemia and *YARS2* variants that we identified using a sideroblastic anemia gene panel or exome sequencing. The phenotype of these patients ranged from MLASA to isolated congenital sideroblastic anemia. As in previous cases, inter- and intra-familial phenotypic variability was observed, however, this report

includes the first cases with isolated sideroblastic anemia and patients with biallelic *YARS2* variants that have no clinically ascertainable phenotype. We identified ten novel *YARS2* variants and three previously reported variants. *In vitro* amino-acylation assays of five novel missense variants showed that three had less effect on the catalytic activity of *YARS2* than the most commonly reported variant, p.(Phe52Leu), associated with MLASA2, which may explain the milder phenotypes in patients with these variants. However, the other two missense variants had a more severe effect on *YARS2* catalytic efficiency. Several patients carried the common *YARS2* c.572 G>T, p.(Gly191Val) variant (minor allele frequency = 0.1259) in *trans* with a rare deleterious *YARS2* variant. We have previously shown that the p.(Gly191Val) variant reduces *YARS2* catalytic activity. Consequently, we suggest that biallelic *YARS2* variants, including severe loss-of-function alleles in *trans* of the common p.(Gly191Val) variant, should be considered as a cause of isolated congenital sideroblastic anemia, as well as the MLASA syndromic phenotype.

## Introduction

Sideroblastic anemia is defined by the presence of bone marrow ringed sideroblasts, which are erythroblasts containing pathological intramitochondrial iron deposits.<sup>1</sup> Congenital sideroblastic anemias (CSAs) are caused by a growing list of genetic variants that affect mitochondrial pathways, including heme synthesis, iron-sulfur cluster biogenesis, mitochondrial protein synthesis, and oxidative phosphorylation.<sup>2,3</sup> Variants in *YARS2* have been associated with myopathy, lactic acidosis, and sideroblastic anemia 2 (MLASA2; OMIM #613561),<sup>4,8</sup> and recently cases of *YARS2*-related myopathy in the absence of sideroblastic anemia have been reported.<sup>9</sup> *YARS2* encodes the mitochondrial tyrosyl-tRNA synthetase, *YARS2*, which is responsible for the ATP-dependent conjugation of tyrosine to its cognate tRNA, required to support mitochondrial protein synthesis.<sup>10</sup> *YARS2* catalyses this reaction in a two-step process. In the first step, tyrosine and ATP bind to the catalytic domain to form the tyrosyl-adenylate intermediate. In the second step, cognate tRNA<sup>Tyr</sup> binds the synthetase and the tyrosyl moiety is transferred to the tRNA CCA-end. The resulting tyrosyl-tRNA<sup>Tyr</sup> will be delivered to the ribosome.

The most commonly reported *YARS2* variant, p.(Phe52Leu), prevalent in patients of Lebanese Christian descent, has been shown to reduce *YARS2* amino-acylation catalytic efficiency by approximately 9-fold, and leads to a reduction in mitochondrial protein synthesis in patients with MLASA2.<sup>4</sup> Here we report *YARS2* variants, some of which were associated with milder effects on amino-acylation, in patients with isolated CSA, or CSA with mild myopathy and lactic acidosis. In addition, we describe two pairs of genotypically identical siblings with divergent, affected and unaffected, clinical phenotypes. Importantly, some patients carry a common *YARS2* c.572 G>T, p.(Gly191Val), that we and others have previously shown has a mild effect on amino-acylation activity,<sup>5,11</sup> and suggest that these milder alleles may be the basis of the reduced penetrance and expressivity.

## Methods

### Clinical data

The patients and their immediate family members were referred to MMH, MDF, NS or LA for clinical consultation. Written informed consent was obtained from participants in the study, as approved by the Institutional Review Boards of Boston Children's

Hospital, USA, the Radboud University Medical Center, the Netherlands, and the Hospital Germans Trias i Pujol, Badalona, Spain. In each case, CSA was ascertained by complete blood counts (CBCs), and peripheral blood or bone marrow morphology. Detailed clinical histories are provided in the *Online Supplementary Appendix*.

### Variant detection

Targeted sequencing of nuclear encoded CSA genes,<sup>12</sup> and the mitochondrial genome as well as mitochondrial DNA deletion analysis was performed on the probands of families 1-3 and 5-9. Genomic DNA was isolated from peripheral blood or skin fibroblasts, using the Puregene DNA Purification Kit (Qiagen, Valencia, CA, USA). DNA templates for sequencing were amplified from genomic DNA by PCR, enzymatically cleaned, bidirectionally sequenced using fluorescent dye termination sequencing chemistry, and analyzed with the Sequencher 5.3 DNA sequence assembly software (Gene Codes, Ann Arbor, MI, USA), as previously described.<sup>12</sup>

Exome sequencing for Patient 4 was performed on genomic DNA isolated from whole blood. The experimental workflow was performed at BGI Europe (Beijing Genome Institute Europe, Copenhagen, Denmark) using an Illumina HiSeq (Illumina, CA, USA) platform. Variants in genes previously associated with Mendelian diseases (OMIM), including CSAs, were analyzed bioinformatically.

Patient 10 DNA was analyzed using a targeted gene panel for congenital and acquired sideroblastic anemias, including *ABCB7*, *ALAS2*, *GLRX5*, *PUS1*, *SF3B1*, *SLC19A2*, *SLC25A38*, *STEAP3*, *TRNT1* and *YARS2*. The library was constructed using the Custom HaloPlex™ Target Enrichment System (Agilent, Santa Clara, CA, USA) and sequenced on a MiSeq platform (Illumina, San Diego, CA, USA). Data were analyzed with SureCall software (Agilent, Santa Clara, CA, USA).

Patient 11 DNA was analyzed using a targeted gene panel for sideroblastic anemia (*ABCB7*, *ALAS2*, *GLRX5*, *HSCB*, *HSPA9*, *PUS1*, *SLC25A38*, *STEAP3*, *YARS2*) and ion semiconductor sequencing as developed by Ion Torrent systems.<sup>13</sup>

*In silico* predictions of variant pathogenicity were performed using the Alamut Visual suite of genetic analysis software (Interactive Biosoftware, Rouen, France), and linking externally to the PolyPhen2 and SIFT analytical tools.<sup>14,15</sup> Minor allele frequencies are reported as in gnomAD (*gnomad.broadinstitute.org*) current as of September 2017.<sup>16</sup>

### Amino-acylation assays

Recombinant wild-type and the p.(Leu61Val), p.(Met195Ile), p.(Ser203Ile), p.(Tyr236Cys) and p.(Gly244Ala) *YARS2* variants were expressed in *E. coli*, purified to homogeneity and assayed for

tyrosylation activity as previously described.<sup>10</sup> Apparent kinetic parameters were determined from Lineweaver-Burk plots in the presence of 4.8 to 6.5 nM YARS2 and 0.28 to 1.12  $\mu$ M native *E. coli* tRNA<sup>Tyr</sup> (Sigma, St. Louis, MO, USA). Experimental errors on  $k_{cat}$  and  $K_m$  varied at most by 20%. Numerical values are averages of at least two independent experiments.

## Results

### Phenotypic spectrum

Eleven probands with CSA were identified with potentially pathogenic *YARS2* variants by targeted gene

sequencing panels or exome sequencing (Table 1A and 1B). The majority of these families were derived from a group of more than 200 probands with CSA referred to SSB, MDF and MMH, in which approximately 4% of cases were attributed to *YARS2* variants. *YARS2* variants have previously been identified in patients with myopathy, lactic acidosis and sideroblastic anemia 2 (MLASA2);<sup>4</sup> however, some patients in this study did not have overt clinical features of MLASA2 other than CSA, and several individuals with biallelic variants had no phenotype whatsoever. In two families, the proband had moderate sideroblastic anemia (P8a and P9a), while a sibling with the same *YARS2* genotype was not anemic and was otherwise

**Table 1A. Clinical data.**

Participant ID	P1	P2a	P2b	P3	P4	P5	P6
YARS2 variant 1 (NM_001040436.2)	c.156C>G p.(Phe52Leu)	c.156C>G p.(Phe52Leu)	c.156C>G p.(Phe52Leu)	c.156C>G p.(Phe52Leu)	c.181C>G p.(Leu61Val)	c.585G>A p.(Met195Ile)	c.572G>T p.(Gly191Val)
YARS2 variant 2	c.156C>G p.(Phe52Leu)	c.156C>G p.(Phe52Leu)	c.156C>G p.(Phe52Leu)	c.156C>G p.(Phe52Leu)	c.181C>G p.(Leu61Val)	c.1165_1166insG p.(Leu389Cysfs*6)	c.590_625del p.(Thr197_Leu208del)
Year of birth	1988	2007	2009	2007	1986	2001	1998
Gender	Female	Male	Female	Male	Male	Female	Female
Ethnicity	Lebanese/ American	Lebanese	Lebanese	Lebanese	Caucasian/ Dutch	Caucasian/ American	African American
Consanguinity	No	Yes	Yes	No	Yes	No	No
Age at presentation	14 years	6 years	4 years	9 years	19 years	2 years	20 months
Sideroblastic anemia	Severe, transfusion dependent from 27 years	Severe, transfusion dependent from 6 years	Mild	Moderate	Severe, transfusion dependent intermittently from 20 years	Severe, transfusion dependent intermittently from 2 years	Severe, transfusion dependent from 20 months
Hemoglobin, g/dL	9.9	10.5	11.5	9.5	6.6	3	2
MCV, fL	84.5	111	102	92.4	81	82.6	101
Abs Retic, M/ $\mu$ L	0.101	0.035	0.13	0.132	ND	0.0175	0.016
Retic, %	3.1	1	2.9	2	1.1	1.7	2.3
WBC x10 <sup>9</sup> /L	2.41	4.35	6.6	5.2	5.4	3.63	6.2
ANC x10 <sup>9</sup> /L	780	2960	3480	2012	ND	617	861-2070
Platelets x10 <sup>9</sup> /L	294	195	305	216	374	163	324
RS, % of BM erythroblasts	10	ND	ND	30	56	ND	>15
Transferrin saturation, %	60 (2002)	97 (2016)	80 (2016)	45	90 (2014)	53	91 (2000)
Ferritin, ng/mL	34.4 (2002)	825 (2016)	296 (2016)	61	683 (2014)	93	256 (2000)
Chelation (year started)	No	Yes (2016)	Yes (2017)	No	Yes (2012)	Yes (2011)	Yes (2004)
Lactic acidosis	Severe 9.1 mmol/L	ND	ND	Severe 9.5 mmol/L	Exercise induced only	Premortem only	Mild
Myopathy	Severe	None	None	Mild	None	Mild	None
Other clinical features	Sinus tachycardia, pericardial effusion, neutropenia thrombocytopenia, primary ovarian failure	Atrial septal defect	None	Diarrhea, hepatosplenomegaly	Successful stem cell transplant at 28 years	Mild cardiomyopathy	Thrombocytopenia, intermittent neutropenia
Vital status	Deceased at 28 y	Alive	Alive	Alive	Alive	Deceased at 12 y	Alive

MCV: mean corpuscular volume; retic: reticulocytes; WBC: white blood cell count; ANC: absolute neutrophil count; RS: ringed sideroblasts; BM: bone marrow; ND: not determined; y: years.

asymptomatic (P8b and P9b) (Table 1B). In a third family (P2a and P2b) (Table 1A), the proband was identified with a severe, new onset anemia at six years of age, and subsequent to her brother's diagnosis, the younger sibling was found to be anemic. Four of the probands presented within the first two years of life (P5, P6, P7, P9a), and 4 presented in adolescence (P1, P4, P8a, P11). Two patients have died (P1, P5), both from multi-organ failure, one of these following two unsuccessful hematopoietic stem cell transplantations (HSCTs). One patient (P4) has undergone successful HSCT.

The 11 probands all had moderate to severe normocytic to macrocytic anemia. In nine probands, the presence of ringed sideroblasts, ranging from 10% to over 50% of bone marrow erythroblasts, was documented on bone marrow aspiration; marrows were not examined in 3

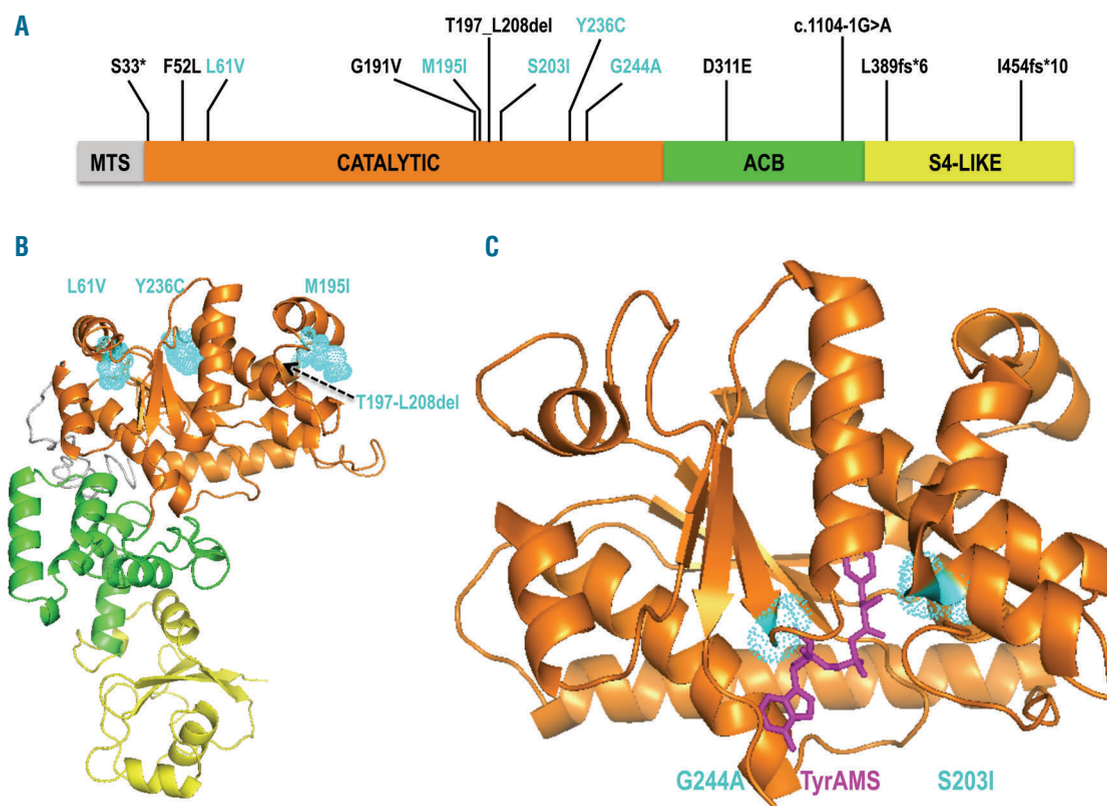
other patients and 2 clinically unaffected siblings (Table 1A and B). Eight patients required transfusion; however, one patient spontaneously became transfusion independent at 16 months of age (P7), and 3 patients had periods of hematologic remission (P4, P5, P9a), transiently becoming RBC transfusion independent. In addition to anemia, 3 probands had variable neutropenia and/or thrombocytopenia (P1, P6, P8a). Four patients were treated with pyridoxine with no improvement in their anemia (P4, P5, P6, P11).

Two patients had severe lactic acidosis (P1, P3), but the remaining cases in which it was studied had mild or no lactic acidosis (Table 1A and B). Two patients had elevated blood lactate upon light exercise (P4, P8a); those with mild lactic acidosis also tended to have mild myopathy, although one patient with no reported lactic acidosis had

**Table 1B. Clinical data.**

Participant ID	P7	P8a	P8b	P9a	P9b	P10	P11
YARS2 variant 1 (NM_001040436.2)	c.[572G>T;731G>C] p.(Gly191Val); (Gly244Ala)	c.572G>T p.(Gly191Val)	c.572G>T p.(Gly191Val)	c.98C>A p.(Ser33*)	c.98C>A p.(Ser33*)	c.608G>T p.(Ser203Ile)	c.933C>G p.(Asp311Glu)
YARS2 variant 2	c.933C>G p.(Asp311Glu)	c.1360_1361insG p.(Ile454Serfs*10)	c.1360_1361insG p.(Ile454Serfs*10)	c.707A>G p.(Tyr236Cys)	c.707A>G p.(Tyr236Cys)	c.1104-1G>A p.?	c.933C>G p.(Asp311Glu)
Year of birth	1999	1963	1965	2010	2010	1992	2003
Gender	Female	Female	Female	Male	Male	Female	Male
Ethnicity	Caucasian/ American	Caucasian/ American	Caucasian/ American	Caucasian/ American	Caucasian/ American	Caucasian/ Spanish	Caucasian / Dutch
Consanguinity	No	No	No	No	No	Unknown	No
Age at presentation	3 months	18 years	49 years (Asymptomatic)	3 months	3 months (Asymptomatic)	23 years	13 years
Sideroblastic anemia	Severe, transfusion dependent until 16 months	Moderate	None	Severe, transfused intermittently from 3 months	None	Moderate	Severe transfusion dependent from 13 years
Hemoglobin, g/dL	5.8	9.9	13.9	2.4	12.8	9.6	6.6
MCV, fL	94.6	111.9	82	113.8	94.1	86	95
Abs Retic, M/ $\mu$ L	0.037	0.059	0.106	0.015	0.053	0.088	0.018
Retic, %	1.8	2.3	2.1	2.4	1.3	2.38	0.8
WBC x10 <sup>9</sup> /L	8.01	6	6.8	10.1	9.8	7.65	4.9
ANC x10 <sup>9</sup> /L	1201	3600	4340	1919	3180	4280	1700
Platelets x10 <sup>9</sup> /L	337	149	182	537	414	243	257
RS, % of BM erythroblasts	47	40	ND	>50	ND	32	81
Transferrin saturation, %	ND	66.7 (2015)	Unknown	45 (2016)	51 (2015)	79.4 (2015)	62 (2016)
Ferritin, ng/mL	ND	387 (2015)	Unknown	225 (2016)	42 (2015)	295 (2015)	180 (2016)
Chelation (Year started)	No	No	No	No	No	No	Yes (2017)
Lactic acidosis	None	Exercise induced only	None	None	ND	None	Mild
Myopathy	Moderate	Mild	None	None	None	None	Mild
Other clinical features	Intermittent diarrhea and abdominal pain	Dependent edema, leukopenia, thrombocytopenia, atypical pulmonary carcinoid tumor (age 53)	None	Facial dysmorphism	Facial dysmorphism	Asthenia	None
Vital status	Alive	Alive	Alive	Alive	Alive	Alive	Alive

MCV: mean corpuscular volume; retic: reticulocytes; WBC: white blood cell count; ANC: absolute neutrophil count; RS: ringed sideroblasts; BM: bone marrow; ND: not determined.



**Figure 1. Representation of mutated YARS2 proteins.** (A) Schematic view of YARS2 domains: MTS: mitochondrial targeting sequence; ACB: anticodon binding domain; S4-Like: S4 ribosomal protein-like domain. Amongst all the variants identified, only those tested in this study are shown in cyan. Note that the recombinant YARS2 used in the amino-acylation assays is deprived of the MTS. (B) Model of YARS2 p.(Thr197-Leu208del), built with I-TASSER.<sup>28</sup> The structural domains from (A) are shown with the same color code. The locations of the variants, which have the weakest effects on amino-acylation [p.(Leu61Val), p.(Met195Ile), p.(Tyr236Cys)] are shown in cyan. (C) Crystal structure of YARS2 catalytic domain<sup>19</sup> with the tyrosyl-adenylate analog (TyrAMS, magenta) bound to the active site. The locations of variants p.(Ser203Ile) and p.(Gly244Ala), characterized by the strongest effects on amino-acylation, are indicated in cyan.

**Table 2. *In silico* predictions of pathogenicity for YARS2 missense variants.**

YARS2 variant	SIFT prediction	SIFT score	PolyPhen2 prediction	PolyPhen2 score	gnomAD frequency (%)
p.(Leu61Val)	Deleterious	0.03	Benign	0.001	0.0016*
p.(Met195Ile)	Tolerated	0.17	Possibly damaging	0.827	0
p.(Ser203Ile)	Deleterious	0.02	Probably damaging	0.989	0
p.(Tyr236Cys)	Tolerated	0.09	Probably damaging	1.000	0.0008*
p.(Gly244Ala)	Deleterious	0.00	Probably damaging	0.995	0.0047*

\*No homozygotes reported.

moderate myopathy (P7). Patient 1 (P1) with severe lactic acidosis and myopathy had combined respiratory chain deficiency in skeletal muscle, and the muscle biopsy showed histopathological features typical of a mitochondrial myopathy, including ragged red fibers on trichrome stain and “parking lot” inclusions and whorled arrays of mitochondrial cristae by transmission electron microscopy (*data not shown*). In one family, the proband (P9a) and his clinically unaffected, but genotypically identical sibling (P9b), had distinctive “triangular” faces, unlike their parents or genotypically normal sibling, which has not previously been reported in association with YARS2 variants, but has been described in mitochondrial myopathy with lactic acidosis and sideroblastic anemia 1 (MLASA1; OMIM #600462) due to pseudouridine synthase 1 (*PUS1*) variants.<sup>17</sup>

### YARS2 variants in patients with congenital sideroblastic anemia

We identified three previously described YARS2 variants and ten novel variants in patients with CSA: the Lebanese Christian founder variant, p.(Phe52Leu),<sup>4</sup> was in the homozygous state in 4 patients; the p.(Asp311Glu) variant<sup>8</sup> homozygous in one patient; and a novel variant, p.(Leu61Val) homozygous in one patient. The remaining six families had compound heterozygous variants including four novel missense variants: p.(Met195Ile), p.(Ser203Ile), p.(Tyr236Cys), p.(Gly244Ala); a novel nonsense variant p.(Ser33\*); three novel indels, p.(Thr197\_Leu208del), p.(Leu389Cysfs\*6), p.(Ile454Serfs\*10); one novel splicing variant, c.1104-1G>A; and two previously reported missense variants, p.(Gly191Val) and p.(Asp311Glu).<sup>5,8</sup> No patient had two indel or splicing variants.

The five novel missense variants all lie in the catalytic domain of YARS2 (Figure 1A) and are rare in the gnomAD database ([gnomad.broadinstitute.org](http://gnomad.broadinstitute.org)) (Table 2). *In silico* predictions of pathogenicity for p.(Leu61Val), p.(Met195Ile) and p.(Tyr236Cys) vary between the SIFT and PolyPhen2 prediction programs while p.(Ser203Ile) and p.(Gly244Ala) are predicted to be damaging to the YARS2 protein by both algorithms (Table 2 and Figure 1B). Conservation among species for each missense variant is shown in *Online Supplementary Figure S1*.

The nonsense variant, the splicing variant and three novel indels are likely to be deleterious. The splicing variant c.1104-1G>A alters a canonical position in the 3' splice acceptor site of intron 3 and it is predicted to result in skipping of exon 4. The YARS2 c.98C>A, p.(Ser33\*) nonsense variant and the c.1165\_1166insG, p.(Leu389Cysfs\*6) frameshift variant both lie greater than 55 nucleotides upstream of the last exon-exon junction and are most likely targeted for nonsense mediated decay.<sup>18</sup> The p.(Thr197\_Leu208) in frame deletion results in loss of 12 residues in  $\alpha$ -helical regions of the catalytic domain, and more precisely of cluster 1, which is important for tRNA acceptor end recognition<sup>19</sup> (Figure 1B). The c.1360\_1361insG, p.(Ile454Serfs\*10) variant lies in the last exon of YARS2 and is not predicted to be targeted for nonsense mediated decay.<sup>18</sup> This variant would cause a frameshift at position 454 in the S4-like domain, which is found in all prokaryotic and organellar tyrosyl-tRNA synthetases, and is thought to stabilize the interaction between the tRNA and YARS2.<sup>19,20</sup>

### Amino-acylation activity of YARS2 missense variants

Amino-acylation assays are commonly used to evaluate the effect of variants on aminoacyl-tRNA synthetase activity, with reduced activity being a strong predictor of pathogenicity.<sup>21</sup> Consequently, the effect of the five missense variants, p.(Leu61Val), p.(Met195Ile), p.(Ser203Ile), p.(Tyr236Cys) and p.(Gly244Ala) on amino-acylation activity was measured by the incorporation of [<sup>14</sup>C]-tyrosine into an *E. coli* tRNA<sup>Tyr</sup> substrate and compared to wild-type YARS2 activity. *In vitro* studies of the YARS2 variants revealed that amino-acylation efficiency was mildly reduced for p.(Leu61Val) and p.(Met195Ile), while p.(Tyr236Cys) was not affected as compared to the wild-type enzyme (Table 3). YARS2 p.(Ser203Ile) and p.(Gly244Ala) demonstrated a 17-fold loss in catalytic efficiency. The reduced activity of YARS2 p.(Ser203Ile) is a consequence of an increased  $K_m$ , indicating that its affinity for tRNA<sup>Tyr</sup> was reduced. On the other hand, the YARS2 p.(Gly244Ala) is characterized by a 13-fold lower  $k_{cat}$ , suggesting that the variant hinders efficient transfer of the tyrosyl moiety from the active site to the tRNA.

### Discussion

Here we expand the clinical spectrum associated with YARS2 variants and describe patients with milder phenotypes who do not display all the features of MLASA2. Rather, most of the patients we describe presented principally with a normo- or macro-cytic CSA; they are mostly non-syndromic and unlike the most common forms of non-syndromic sideroblastic anemia (e.g. ALAS2 or SLC25A38 deficiency), the anemia is not microcytic. Nevertheless, in addition to ringed sideroblasts, some of

**Table 3.** Kinetic parameters for tyrosylation of *E. coli* tRNA<sup>Tyr</sup> by YARS2 wild-type and novel missense variant recombinant proteins.

YARS2 variant	$K_m$ ( $\mu$ M)	$k_{cat}$ ( $min^{-1}$ )	$k_{cat}/K_m$ (efficiency)	Loss of efficiency* (fold change)
WT	0.75	20.0	26.7	1
p.(Leu61Val)	1.45	9.1	6.3	4.2
p.(Met195Ile)	1.90	25.5	13.4	2.0
p.(Ser203Ile)	18.60	28.6	1.5	17.3
p.(Tyr236Cys)	0.70	16.5	23.6	0.9
p.(Gly244Ala)	1.00	1.5	1.5	17.8

\*Loss of efficiency is calculated relative to wild-type (WT) YARS2.

these patients had vacuolization of marrow precursors and/or other cytopenias that are often seen in the syndromic sideroblastic anemias (e.g. Pearson syndrome), which may be a diagnostic clue.

Patients 1 and 3 had all the typical features of MLASA2, whereas Patients 2a, and 2b, who share homozygosity for the YARS2 Lebanese founder allele, p.(Phe52Leu), had only anemia. Patient 1 also had other features not typically associated with MLASA2, including neutropenia, thrombocytopenia, pericardial effusion, and premature ovarian failure. Neutropenia and pericardial effusion have each been reported in one other patient homozygous for the p.(Phe52Leu) variant.<sup>5,22</sup> Two other patients in the current series with other genotypes also had mild or intermittent neutropenia. Premature ovarian failure is associated with variants in several mitochondrial aminoacyl-tRNA synthetase-encoding genes including *HARS2*, *LARS2* and *AARS2*,<sup>23-25</sup> and thus may be a feature common to mitochondrial protein synthesis defects. There are now 10 reported individuals homozygous for the YARS2 p.(Phe52Leu) variant<sup>5,22</sup> and all have been symptomatic, supporting complete penetrance of this allele. However, the great range of phenotypic severity strongly suggests the presence of other genetic and environmental influences that can modify the effects of YARS2 deficiency.

Patient 4 presented in late adolescence with sideroblastic anemia without myopathy and has a homozygous p.(Leu61Val) variant that diminished the amino-acylation catalytic efficiency 4-fold. Leu61 is located in a region of the catalytic domain specific to mitochondrial YARSs that was proposed to contact the tRNA<sup>Tyr</sup> acceptor helix (Figure 1B).<sup>19</sup> In this case, HSCT appeared to be an effective treatment, restoring the patient's hemoglobin levels to normal.

Patient 5 presented in infancy with CSA and was transfusion dependent other than a remission occurring between three and six years of age; she had no myopathy until her post-HSCT terminal illness. This patient had a YARS2 c.1165\_1166insG variant predicted to result in a null allele, and a novel p.(Met195Ile) variant which lies within cluster 1, in a region involved in recognition of the tyrosine accepting arm of tRNA<sup>Tyr</sup> (Figure 1B).<sup>19</sup> Some YARS proteins (e.g. yeast) have an isoleucine (Ile) at this position, suggesting that it might be a milder allele. Indeed, *in vitro* this mutant had little effect on YARS2 catalytic efficiency.

Patient 10 is a compound heterozygote for a splicing mutation (c.1104-1G>A) predicted to cause skipping of exon 4, and a missense variant p.(Ser203Ile), also located

in cluster 1. YARS2 (p.Ser203Ile) led to a reduced affinity for tRNA<sup>Tyr</sup>, resulting in a 17-fold loss in catalytic efficiency (Figure 1C). Patient 10 has no lactic acidosis or myopathy, and presented with isolated normocytic anemia and asthenia, and has not required transfusion.

Patient 7 presented with anemia in infancy requiring two transfusions within the first 16 months of life and then became transfusion independent. She has moderate myopathy and no lactic acidosis and a compound heterozygous genotype: a missense variant, p.(Gly244Ala), occurring *in cis* with p.(Gly191Val) and *in trans* with the p.(Asp311Glu) variant. Gly244 is a critical residue for tyrosyl-adenylate binding.<sup>19</sup> YARS2 p.(Gly244Ala) only affected the  $k_{cat}$  indicating that, as predicted, this variant hinders binding of the tyrosyl-adenylate in the active site (Figure 1C). YARS2 Asp311 is involved in the recognition of anticodon residue G34 of tRNA<sup>Tyr</sup>.<sup>19</sup> The p.(Asp311Glu) variant is respiratory deficient in a yeast model, and patients homozygous for this allele also have transfusion-dependent sideroblastic anemia in the first year of life; however, in contrast to patient 7, they have lactic acidosis but no myopathy.<sup>8</sup> Further phenotypic variability for the p.(Asp311Glu) variant was observed in Patient 11 who was homozygous for p.(Asp311Glu), with transfusion-dependent MLASA2.

In two families in this study (Families 6 and 8), affected patients have the common p.(Gly191Val) allele (MAF = 0.1259) *in trans* of a predicted null allele. Importantly, all of the unaffected carriers of predicted null alleles in these and other families, where probands had the ancestral p.Gly191 variant *in trans*, were asymptomatic (*data not shown*). Patient 6 presented in infancy with CSA requiring transfusions every three weeks. She has mild lactic acidosis, no myopathy and intermittent neutropenia. She has a c.590\_645del variant resulting in a 12 amino acid deletion in the catalytic domain (Figure 1A), which would almost certainly lead to a completely dysfunctional protein, *in trans* with p.(Gly191Val). Individuals 8a and 8b also carry p.(Gly191Val) *in trans* with a predicted null or severe loss-of-function allele, c.1360\_1360insG, p.(Ile454Serfs\*10). This variant truncates the S4-like domain which is thought to stabilize the interaction with tRNA<sup>Tyr</sup>, and the deletion of the YARS2 S4-like domain leads to a 100-fold reduced amino-acylation activity *in vitro*.<sup>20</sup> Patient 8a had sideroblastic anemia and edema. Lactate was elevated only on exertion and the patient did not have myopathy. Her sister (P8b) is asymptomatic. Patient 8a also had a somatic mutation in SF3B1 p.(Lys700Glu) that is strongly associated with myelodysplastic syndromes with ringed sideroblasts.<sup>26</sup> Based on the childhood presentation of her anemia and exercise intolerance that was exacerbated significantly decades later, and the fact that a mutation in SF3B1 would be exceptional in a patient under 30 years of age, we infer the YARS2 mutations to be the primary cause of her anemia with the SF3B1 mutation occurring as a secondary somatic event, which exacerbated her anemia, bringing her to clinical attention. In addition to the reduced activity *in vitro*,<sup>5</sup> support of the notion that YARS2 p.(Gly191Val) contributes to the disease phenotype in these patients comes from the observation that this variant is a disease modifier in Leber Hereditary Optic Neuropathy (LHON); the three common LHON

mitochondrial DNA mutations have incomplete penetrance. However, all patients who carry both the LHON m.11778G>A mtDNA disease-associated variant in combination with a homozygous YARS2 p.(Gly191Val) genotype were symptomatic.<sup>11</sup>

Patients 9a and 9b carried the YARS2 c.98C>A, p.(Ser33\*) nonsense variant, which would result in a null allele, and the p.(Tyr236Cys) variant (Figure 1A and B) that did not alter amino-acylation activity *in vitro*. In addition, *in silico* analysis using Alamut did not predict that this variant would lead to alteration of an exonic splicing enhancer site. Patient 9a presented in infancy with sideroblastic anemia that has come and gone throughout his life. He has no lactic acidosis or myopathy. He and his unaffected brother have some dysmorphic features, which have not previously been reported in association with YARS2 variants, but are typical of MLASA1 patients with pseudouridine synthase 1 (*PUS1*) mutations.<sup>17,27</sup> His genotypically concordant fraternal twin (P9b) has only mild anemia and similar facial dysmorphism, once again highlighting the potential for decreased penetrance and/or expressivity of the disorder.

Interestingly, some YARS2 patients with myopathy, but no sideroblastic anemia, have recently been reported by Sommerville *et al.*<sup>9</sup> They report siblings with a homozygous YARS2 p.(Leu392Ser) variant who had MLASA2, while another individual homozygous for the same variant had myopathy without sideroblastic anemia or lactic acidosis.

To summarize, the inter- and intra-familial phenotypic variability, intermittent transfusion dependence of some YARS2 cases, and the association of a common variant with disease, suggest that all MLASA2 phenotypes may be susceptible to subtle changes in YARS2 function, which may in turn be influenced by genetic and/or environmental modifiers. This study shows that YARS2 variants can result in CSA in the absence of clinically significant myopathy or lactic acidosis. Thus, we recommend that YARS2 variants be considered as a cause of isolated sideroblastic anemia as well as MLASA2 or mitochondrial myopathy.

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