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Mitochondrial DNA-Associated Leigh Syndrome Spectrum

Synonym: mtDNA-Associated Leigh Syndrome Spectrum

Megan Ball, MD,^{1,2,3} David R Thorburn, PhD, FHGSA, FFSc(RCPA),^{4,2,3} and Shamima Rahman, PhD, FRCP, FRCPCH⁵

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Summary

Clinical characteristics

Mitochondrial DNA-associated Leigh syndrome spectrum (mtDNA-LSS) is part of a continuum of progressive neurodegenerative disorders caused by abnormalities of mitochondrial energy generation, which includes the overlapping phenotypes mtDNA-associated Leigh syndrome and mtDNA-associated Leigh-like syndrome.

Mitochondrial DNA-LSS is characterized by onset of manifestations typically between ages three and 12 months, often following an intercurrent illness (usually viral) or metabolic challenge (vaccinations, surgery, prolonged fasting). Decompensation (often with elevated lactate levels in blood and/or cerebrospinal fluid) is typically associated with developmental delay and/or regression. Neurologic features include hypotonia, spasticity, seizures, movement disorders, cerebellar ataxia, and peripheral neuropathy. Brain stem dysfunction may manifest with respiratory symptoms, swallowing difficulties, ophthalmoparesis, and abnormalities in thermoregulation. Extraneurologic manifestations may include poor weight gain, cardiomyopathy, and conduction defects. Up to 50% of individuals die by age three years, most often from respiratory or cardiac failure.

Diagnosis/testing

The diagnosis of mtDNA-LSS is established in a proband fulfilling clinical diagnostic criteria for LSS by identification of a heteroplasmic or homoplasmic pathogenic variant in one of the 15 mtDNA genes known to be involved in mtDNA-LSS.

Author Affiliations: 1 Metabolic Fellow, PhD Candidate, Murdoch Children's Research Institute, Victoria, Australia; Email: megan.ball@mcri.edu.au. 2 Royal Children's Hospital, Victoria, Australia; Email: megan.ball@mcri.edu.au; Email: david.thorburn@mcri.edu.au. 3 Department of Paediatrics, University of Melbourne, Melbourne, Australia; Email: megan.ball@mcri.edu.au; Email: david.thorburn@mcri.edu.au. 4 Professorial Fellow, Murdoch Children's Research Institute and Genetic Health Services Victoria, Victoria, Australia; Email: david.thorburn@mcri.edu.au. 5 Professor of Paediatric Metabolic Medicine, Genetics and Genomic Medicine Department, UCL Great Ormond Street Institute of Child Health, London, United Kingdom; Email: shamima.rahman@ucl.ac.uk.

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Management

Treatment of manifestations: Treatment is supportive. Sodium bicarbonate or sodium citrate for significant acidosis (THAM may be used if there is hypernatraemia); anti-seizure medication for seizures; dystonia therapy with benzhexol, baclofen, tetrabenazine, or gabapentin alone or in combination, or by botulinum toxin injection; treatment of respiratory compromise per pulmonologist; caloric and nutritional supplementation and feeding therapy as needed; developmental and educational support; physical therapy and occupational therapy; standard treatment of eye movement disorders; medical management of cardiomyopathy; treatment of constipation as needed; treatment of liver failure per hepatologist; treatment of electrolyte abnormalities per nephrologist; hearing aids or cochlear implants for sensorineural hearing loss; speech therapy and hearing support services as needed; management of diabetes mellitus and adrenal insufficiency per endocrinologist; standard treatments for anxiety and/or depression; psychological support and care coordination for the affected individual and family.

Surveillance: Affected individuals should be followed at regular intervals to monitor for progression of disease and associated complications. Neurologic assessment for ataxia and seizures at each visit along with an assessment of pulmonary issues, growth, nutrition, and gastrointestinal manifestations. Development, educational, and cognitive assessment and assessment of mobility and self-help skills at least annually. Ophthalmology assessment every six to 12 months or as advised by ophthalmologist. Annual blood pressure, EKG, and echocardiogram or as advised by cardiologist. Liver function tests, urinalysis, urine albumin-to-creatinine ratio, urine amino acids, serum electrolytes, blood urea nitrogen, creatinine, complete blood count, and fasting glucose annually. Annual audiology assessment. Assessment of care coordination and family psychosocial needs at each visit.

Agents/circumstances to avoid: Sodium valproate, medications that cause acidosis, and dichloroacetate should be avoided or used with caution; administration of anesthesia requires careful consideration to avoid aggravation of respiratory symptoms and precipitation of respiratory failure.

Genetic counseling

Mitochondrial DNA-LSS is transmitted by maternal inheritance. The mother of a proband may have the mtDNA pathogenic variant and may exhibit mild clinical manifestations of mtDNA-LSS. Many affected individuals have no known family history of mtDNA-LSS or other mitochondrial disorder. The explanation for apparently simplex cases may be absence of a comprehensive and/or reliable family history, a maternal heteroplasmy level below the disease threshold (i.e., the minimum level of heteroplasmy expected to result in mitochondrial disease), or a *de novo* mtDNA pathogenic variant in the proband. If the mother of the proband has the mtDNA pathogenic variant identified in the proband, all sibs of the proband are at risk of inheriting the pathogenic variant. Sibs may inherit the pathogenic variant at varying heteroplasmy levels due to the bottleneck effect and variant-specific segregation patterns. The risk to a sib of developing clinical manifestations is difficult to determine and depends on heteroplasmy level, the variation in heteroplasmy levels among different tissues, and the disease threshold for the specific variant. Recurrence risk assessment and prenatal testing for disorders caused by pathogenic variants in mtDNA is challenging due to the intricacies of mtDNA transmission and the inherent challenge in using prenatal genetic test results to predict clinical outcome. Reproductive options for the family members of a proband with an mtDNA pathogenic variant may include prenatal testing, preimplantation genetic testing, and oocyte donation.

Diagnosis

Diagnostic criteria for Leigh syndrome spectrum (LSS) have been published [McCormick et al 2023].

Suggestive Findings

Mitochondrial DNA (mtDNA)-associated LSS (mtDNA-LSS) should be suspected in probands with several of the following clinical, laboratory, and/or imaging findings.

Clinical findings

- Progressive neurologic disease with developmental delay and neurodevelopmental regression
- Manifestations of brain stem and/or basal ganglia disease (e.g., respiratory abnormalities, nystagmus, ophthalmoparesis, optic atrophy, ataxia, and dystonia)
- Seizures
- Psychiatric disturbance

Laboratory findings

- Elevated blood lactate
 Elevation tends to be more marked in postprandial samples.
- Elevated cerebrospinal fluid (CSF) lactate
 Note: While elevated blood and/or CSF lactate is suggestive of a mitochondrial disorder, a normal lactate does not exclude mtDNA-LSS.
- Plasma amino acids may show increased alanine concentration, reflecting persistent hyperlactatemia. Decreased plasma citrulline concentration and/or elevated 3-hydroxy-isovalerylcarnitine (C5-OH) levels has been reported in individuals with *MT-ATP6* pathogenic variants [Balasubramaniam et al 2017, Larson et al 2019, Peretz et al 2021, Tise et al 2023].
- Urine organic acid analysis results are often nonspecific and may show lactic aciduria, increased Krebs cycle intermediates, and increased dicarboxylic acids.
 - Note: 3-methylglutaconic aciduria may be observed in some nuclear gene-encoded causes of LSS, notably 3-methylglutaconic aciduria with deafness-dystonia, [hepatopathy], encephalomyopathy, and Leigh-like syndrome (MEGD[H]EL syndrome caused by SERAC1 deficiency), and ethylmalonic aciduria is a feature of ethylmalonic encephalopathy caused by biallelic variants in ETHE1. Identification of increased methylmalonic acid or propionic acid is suggestive of other genetic causes of LSS or organic acidemias (e.g., SUCLG1- or SUCLA2-related mtDNA depletion syndrome, methylmalonic acidemia, propionic acidemia) (see Causes of Nuclear Gene-Encoded Leigh Syndrome Spectrum and Differential Diagnosis).

Brain imaging findings

- Characteristic findings include bilateral symmetric hyperintense signal abnormality in the brain stem and/or basal ganglia on T₂-weighted MRI or bilateral symmetric hypodensities in the basal ganglia on CT that may progress/fluctuate with time [Bonfante et al 2016, Alves et al 2020]. Corresponding restricted diffusion may be evident on diffusion-weighted imaging [Alves et al 2020].
- Additional findings can include cerebral atrophy, bilateral hyperintense signal abnormality in the thalamus, cerebellum, cerebral white matter, and/or spinal cord on T₂-weighted MRI [Bonfante et al 2016, Alves et al 2020, Hong et al 2020, Stenton et al 2022].
- Magnetic resonance spectroscopy (MRS) lactate peak (in the absence of acute seizures)

 Note: Characteristic patterns of brain lesions have been described for specific causes of nuclear geneencoded LSS. For example, specific brain lesions affecting the mammillothalamic tracts, substantia nigra, medial lemniscus, medial longitudinal fasciculus, spinothalamic tracts, and cerebellum appear to be

characteristic of *NDUFAF2*-related LSS. SERAC1 deficiency is associated with a distinctive brain MRI pattern affecting the basal ganglia, especially the putamen. Initially there are T₂-weighted signal changes of the pallidum, and later swelling of the putamen and caudate nucleus with an "eye" representing early sparing of the dorsal putamen, followed by progressive involvement of the putamina (see Causes of Nuclear Gene-Encoded Leigh Syndrome Spectrum).

Histopathology of muscle tissue may be normal or may show nonspecific changes such as accumulation of intracytoplasmic neutral lipid droplets or predominance of type II muscle fibers. Ragged red fibers and/or cytochrome *c* oxidase-negative fibers may occasionally be observed.

Since the advent of next-generation sequencing, muscle biopsy is performed less frequently (especially in the pediatric population), as obtaining a muscle biopsy is invasive and often requires general anesthesia, which carries a risk of metabolic decompensation. Muscle biopsy is now generally reserved for critically unwell individuals or (occasionally) for functional validation of genetic findings.

Note: (1) Although muscle histopathology is only occasionally abnormal, when it is abnormal it can be as much of a contributor to diagnostic certainty as respiratory chain enzymes or molecular testing. (2) If muscle biopsy is obtained for respiratory chain enzyme studies, histopathology should also be performed.

Respiratory chain enzyme studies. Biochemical analysis of tissue biopsies or cultured cells often detects deficient activity (<30% of control mean values) of one or more of the respiratory chain enzyme complexes. Isolated defects of complex I or complex IV are the most common enzyme abnormalities observed in individuals with LSS and can help guide subsequent molecular genetic testing of mtDNA or nuclear genes [Sofou et al 2018, Hong et al 2020, Ogawa et al 2020, Ardissone et al 2021]. Combined respiratory chain enzyme defects can also be observed, particularly in disorders of mitochondrial gene expression. Normal respiratory chain enzyme analysis does not exclude mtDNA-LSS and can be seen in individuals with mtDNA pathogenic variants affecting complex V subunits such as the *MT-ATP6* pathogenic variants m.8993T>G, m.8993T>C, and m.9176T>C.

Note: (1) Skeletal muscle (vastus lateralis) is usually the tissue of choice for respiratory chain enzyme studies. (2) Skin fibroblasts can be used, but only about 50% of respiratory chain enzyme defects identified in skeletal muscle are also identified in skin fibroblasts. (3) Approximately 10%-20% of individuals with normal skeletal muscle respiratory chain enzyme analysis may have an enzyme defect detected in liver or cardiac muscle, particularly if those tissues are involved clinically [Frazier et al 2020].

Establishing the Diagnosis

The diagnosis of mtDNA-LSS **is established** in a proband fulfilling criteria for LSS [McCormick et al 2023] in whom a heteroplasmic or homoplasmic pathogenic variant in one of the genes listed in Table 1 has been identified.

Clinical Criteria for mtDNA-LSS

Typical neuroradiologic findings. Bilateral, typically symmetric hyperintensities on T₂-weighted MRI or hypodensities on CT in the brain stem and/or basal ganglia with or without bilateral, T₂-weighted hyperintensities on MRI or hypodensities on CT in the thalamus, cerebellum, subcortical white matter, and/or spinal cord

AND ≥1 of the following **characteristic neurologic clinical features**:

- Developmental regression
- Developmental delay
- Psychiatric features

AND ≥1 of the following **biochemical and/or mitochondrial abnormalities**:

- Elevated lactate in plasma and/or CSF
- MRS lactate peak (in absence of acute seizures)
- Respiratory chain enzyme activity deficiency (<30% enzyme activity) in affected tissues (muscle, liver, fibroblasts)

Note: Historically, the diagnosis of LSS was achieved by demonstrating mitochondrial dysfunction through relevant tissue biopsy to inform a targeted molecular genetic approach. However, the introduction of comprehensive genomic sequencing has removed the need for tissue biopsy in most scenarios.

Molecular Genetic Testing

Molecular genetic testing approaches for mtDNA-LSS can include **comprehensive genomic testing** (genome sequencing or mtDNA sequencing and exome sequencing) or **targeted single-gene testing**. Choice of approach may be dependent on clinical suspicion, age of presentation, and availability of genomic testing.

Option 1

Comprehensive genomic testing includes sequencing of mtDNA and nuclear DNA. Comprehensive genomic testing does not require a phenotype-driven hypothesis and may provide or suggest a diagnosis not previously considered (e.g., variant in a different gene[s] that results in a similar clinical presentation).

- Genome sequencing is preferred (if available), as it allows analysis of the mitochondrial and nuclear genome, and typically has the highest diagnostic rate.
- If genome sequencing is not available, mtDNA sequencing is recommended. Exome sequencing should be considered in conjunction with mtDNA sequencing, especially in the pediatric population, given that approximately 70% of individuals will have nuclear gene-encoded LSS (see Differential Diagnosis). For children, a parent-child trio should be considered to assist in the identification of rare autosomal recessive or *de novo* pathogenic variants. It may be possible to extract mtDNA sequence data from some exome kits, but coverage varies and may not be of clinical quality, hence stand-alone mtDNA sequencing is recommended.

For an introduction to comprehensive genomic testing click here. More detailed information for clinicians ordering genomic testing can be found here.

Option 2

Targeted analysis of common pathogenic variants may be considered if there is a high clinical suspicion based on clinical or biochemical findings or if comprehensive testing is not available (see Table 1).

Note: (1) Most mtDNA pathogenic variants associated with mtDNA-LSS are "heteroplasmic" (i.e., abnormal mtDNA coexists with wild type mtDNA) and for some pathogenic variants, the heteroplasmy level may vary among different tissues and may increase or decrease with age. (2) Mitochondrial DNA pathogenic variants may be lost from the leukocyte population with increasing age [Schon et al 2021, Davis et al 2022]; therefore, in adults with milder manifestations and asymptomatic older maternal relatives, the pathogenic variant may only be detected in hair follicles, urine sediment cells, or skeletal muscle, which is the most reliable source of mtDNA [Davis et al 2022]. (3) Leukocyte testing is recommended in children, particularly when using next-generation sequencing methods, which allow detection of very low heteroplasmy levels. (4) Long-range PCR can be performed to identify large mtDNA rearrangements. (5) Mitochondrial DNA deletions are not usually detectable in leukocyte DNA in adults; in this age group, skeletal muscle (or urinary sediment cells) is the tissue of choice for analysis [Schon et al 2021, Davis et al 2022]. (6) Deletions/duplications of mtDNA are an extremely rare cause of LSS in adults. (7) The authors are not aware of any published reports of children with mtDNA-LSS when the pathogenic variant was present in another tissue but not detected in blood.

Table 1. Molecular Genetics of Mitochondrial DNA-Associated Leigh Syndrome Spectrum

	% of mtDNA-LSS Attributed to	Proportion of Pathogenic Variants ⁵ Identified by Method		
Gene ^{1, 2}	Pathogenic Variants in Gene ^{3, 4}	Sequence analysis ⁶	Mitochondrial DNA deletion/ duplication analysis ⁷	
MT-ATP6	~40% 8			
MT-ND3	~20%			
MT-ND5	~15%			
MT-ND6	~10%			
MT-ND4	~5%	>95%	<5% ⁹	
MT-ND1	<5%			
MT-TL1	<5%			
MT-CO1	Rare			
MT-CO2	Rare			
MT-CO3	Rare			
MT-ND2	Rare			
MT-TI	Rare			
MT-TK	Rare			
MT- TV	Rare			
MT-TW	Rare			

mtDNA-LSS = mitochondrial DNA-associated Leigh syndrome spectrum

- 1. Genes are listed from most frequent to least frequent genetic cause of mtDNA-LSS and then alphabetically.
- 2. See Table A. Genes and Databases for chromosome locus and protein.
- 3. These estimates are based on several cohorts of individuals with mtDNA-LSS, which are likely affected by inclusion criteria, testing strategies, and ascertainment bias [Sofou et al 2014, Bonfante et al 2016, Sofou et al 2018, Wei et al 2018, Yu et al 2018, Alves et al 2020, Hong et al 2020, Lee et al 2020, Ogawa et al 2020, Ardissone et al 2021, Lim et al 2022, Stenton et al 2022, Tinker et al 2022, Ardissone et al 2023, Baldo et al 2023, Kistol et al 2023, Na & Lee 2023].
- 4. Data derived from MITOMAP
- 5. See Molecular Genetics for information on pathogenic variants detected.
- 6. Sequence analysis detects variants that are benign, likely benign, of uncertain significance, likely pathogenic, or pathogenic. Variants may include missense and nonsense variants and small deletions/insertions; typically, larger deletions/duplications are not detected. For issues to consider in interpretation of sequence analysis results, click here.
- 7. Single-gene mtDNA deletions have not been reported. Deletion/duplication analysis of the entire mitochondrial genome may include a range of techniques such as quantitative PCR or long-range PCR.
- 8. The most common *MT-ATP6* pathogenic variants in individuals with mtDNA-LSS include m.8993T>G, m.8993T>C, m.9176T>C, and m.9185T>C [Ganetzky et al 2019, Stendel et al 2020, Na & Lee 2022].
- 9. Large-scale deletions have been reported in individuals with mtDNA-LSS [Yamashita et al 2008, Ardissone et al 2021].

Clinical Characteristics

Clinical Description

Mitochondrial DNA-associated Leigh syndrome spectrum (mtDNA-LSS) is a continuum of progressive neurodegenerative disorders, with typical onset in infancy or early childhood of sudden neurodevelopmental regression. Individuals with LSS may have variable extraneurologic manifestations including gastrointestinal, cardiac, hepatic, renal, and endocrine disease. The clinical features of mtDNA-LSS are similar to those observed in individuals with nuclear gene-encoded Leigh syndrome spectrum (LSS). Clinically, it is difficult to distinguish mtDNA-LSS from nuclear gene-encoded LSS, and there are relatively few publications describing the full range

of clinical features in mtDNA-LSS. The following description of the phenotypic features summarizes the more robust data on features of all individuals with LSS; specific information and references for mtDNA-LSS are included if available.

Table 2. Mitochondrial DNA-Associated Leigh Syndrome Spectrum: Frequency of Select Features

Feature		% of Persons w/Feature		
		LSS ¹	mtDNA-LSS ²	Comment
	Developmental delay	>70%	>75%	
	Regression	~50%	Unknown ³	
	Hypotonia	~60%	~65%	
	Dystonia	~35%	~30%	
	Ataxia	~30%	~50%	
	Spasticity	~40%	~50%	
Neurologic features	Muscle weakness	~30%	~75%	
-	Peripheral neuropathy	~10%	~5%	
	Dysphagia	~25%	~80%	
	Epilepsy	~30%	~40%	
	Respiratory abnormalities	~30%	~35%	Apnea, hypoventilation, hyperventilation, irregular respiration
	Sensorineural hearing loss	15%	15%	
	Ophthalmoplegia	~25%	~30%	
Ophthalmologic features	Optic atrophy	~15%	~20%	
	Retinopathy	~10%	<10%	
	Poor weight gain	~40%	~30%	
	Gastrointestinal manifestations	~25%	~25%	Vomiting, gastroparesis, intestinal dysmotility, constipation
Other	Cardiac manifestations	~15%	~20%	Hypertrophic or dilated cardiomyopathy, conduction defects
	Hepatic manifestations	<10%	~15%	↑ liver transaminases, hepatomegaly, liver failure
	Renal manifestations	<5%	<5%	Tubulopathy
	Endocrine manifestations	<5%	<5%	Diabetes, adrenal insufficiency

mtDNA = mitochondrial DNA; LSS = Leigh syndrome spectrum

^{1.} Estimated percentages are based on several cohorts of individuals with nuclear gene-encoded LSS and mtDNA-LSS (~1,000 individuals); however, each clinical feature was not reported for every individual [Rahman et al 1996, Naess et al 2009, Sofou et al 2014, Sofou et al 2018, Wei et al 2018, Alves et al 2020, Hong et al 2020, Lee et al 2020, Ardissone et al 2021, Lim et al 2022, Stenton et al 2022, Ardissone et al 2023, Kistol et al 2023].

^{2.} Estimated percentages are based on reports of five cohorts that included clinical features of ~150 individuals with mtDNA-LSS. Estimated percentages of several of the clinical features are based on a small number of individuals (<20) and should be interpreted with caution [Rahman et al 1996, Sofou et al 2018, Wei et al 2018, Ardissone et al 2023, Kistol et al 2023].

^{3.} Data not available for regression, as this was often described in conjunction with developmental delay.

Onset of symptoms is typically between age three and 12 months, often following an intercurrent illness (usually viral) or metabolic challenge (vaccinations, surgery, prolonged fasting). However, disease onset can vary, ranging from prenatal onset to adulthood. Early onset of symptoms within the first 24 months of life has been reported in approximately 75% of individuals with LSS. Later onset (age >2 years) includes presentation in childhood, adolescence, or adulthood and is generally associated with slower progression of disease.

Initial features may be nonspecific and vary with age of onset:

- Prenatally, intrauterine growth restriction, cardiomegaly, and oligohydramnios have been reported in LSS [Sofou et al 2014, Hong et al 2020, Stendel et al 2020, Ardissone et al 2021].
- Early-onset LSS (age <2 years) may present with nonspecific features such as feeding difficulties, poor weight gain, hypotonia, developmental delay, and persistent vomiting.
- Late-onset LSS (age >2 years) may manifest with predominant muscle weakness and movement disorders including ataxia and dystonia.
- Onset in adulthood may manifest with ataxia, peripheral neuropathy, and psychiatric disturbance [Wei et al 2018, Hong et al 2020].

Decompensation (often with increased blood and/or cerebrospinal fluid [CSF] lactate concentrations) during an intercurrent illness is typically associated with sudden loss of developmental skills. A period of recovery may follow the initial decompensation, but the individual rarely returns to the developmental status achieved prior to the presenting illness. Most individuals with LSS will experience at least one acute decompensation that may require hospitalization.

Neurologic features include developmental delay/regression, hypotonia, spasticity, dystonia, muscle weakness, hypo- or hyperreflexia, seizures (myoclonic, generalized tonic-clonic, infantile spasms), movement disorders (including chorea), cerebellar ataxia, and peripheral neuropathy. Adolescents and adults with LSS may exhibit psychiatric disturbance.

Brain stem lesions may cause respiratory difficulty (apnea, central hypo- or hyperventilation, or irregular respiration), bulbar problems (e.g., abnormal swallowing and speech), persistent vomiting, and abnormalities of thermoregulation (hypo- and hyperthermia). Sensorineural hearing loss may be present in approximately 15% of individuals, originating from the cochlea or auditory nerves.

Ophthalmologic findings include optic atrophy, retinitis pigmentosa, strabismus, ptosis, nystagmus, and ophthalmoparesis. Pigmentary retinopathy was reported in 12 out of 13 individuals with an *MT-ATP6* m.8993T>G pathogenic variant in one retrospective cohort [Ng et al 2019].

Gastrointestinal manifestations include poor weight gain, vomiting, dysphagia, gastroparesis, and intestinal dysmotility. One natural history study suggested that gastrointestinal symptoms were the most common extraneurologic manifestation in individuals with LSS, with 25% exhibiting gut dysmotility and nearly half of the cohort requiring gastrostomy or nasogastric tube feeding at follow up [Lim et al 2022].

Cardiac manifestations including hypertrophic or dilated cardiomyopathy and conduction defects (e.g., Wolff-Parkinson-White syndrome and paroxysmal supraventricular tachycardia) are reported in approximately 15% of individuals with LSS. Cardiomyopathy has been reported in up to 70% of individuals with an *MT-ND5* pathogenic variant [Stenton et al 2022, Kistol et al 2023].

Hepatic manifestations including elevated liver transaminases, hepatomegaly, or liver failure have been reported in approximately 10% of individuals with LSS [Naess et al 2009, Van Hove et al 2010, Sofou et al 2014, Duff et al 2015, Alves et al 2020].

Renal manifestations including renal tubulopathy or diffuse glomerulocystic kidney damage have been reported in approximately 5% of individuals with LSS [López et al 2006, Naess et al 2009, Sofou et al 2018, Alves et al 2020, Hong et al 2020, Stenton et al 2022].

Endocrine manifestations including diabetes mellitus and adrenal insufficiency have been reported in a small proportion of individuals with LSS [Alves et al 2020, Ardissone et al 2023].

Prognosis in individuals with LSS is generally poor. Most affected individuals experience episodic deterioration of motor and/or cognitive function interspersed with plateaus in development. Up to 50% of individuals with LSS will die before age three years, most often from respiratory or cardiac failure, neurologic deterioration, or sepsis. However, disease progression can be variable. Some individuals have slower progression of disease, with stable periods lasting years in between decompensations, even into adulthood. In previously undiagnosed individuals, death may appear to be sudden and unexpected.

Poor prognosis has been associated with early-onset disease [Sofou et al 2014, Lee et al 2016, Hong et al 2020, Ogawa et al 2020]. Individuals with pathogenic variants in MT-ND5 [Ogawa et al 2020] and those with MT-ATP6 pathogenic variant m.8993T>G [Stendel et al 2020, Na & Lee 2022] have been reported to be more likely to have a severe disease phenotype.

Intermediate phenotypes. Maternal relatives of individuals with mtDNA-LSS can have any one or a combination of clinical manifestations of mtDNA-LSS, NARP (*n*eurogenic muscle weakness, *a*taxia, and *r*etinitis *p*igmentosa), or other mitochondrial disorders. These include mild learning difficulties, ataxia, muscle weakness, night blindness, deafness, diabetes mellitus, migraine, or sudden unexpected death (see Genetically Related Disorders).

Genotype-Phenotype Correlations

For most mtDNA pathogenic variants, it is difficult to distinguish a correlation between genotype and phenotype because clinical expression of mtDNA pathogenic variants is influenced by pathogenicity of the variant, relative amount of abnormal and wild type mtDNA (heteroplasmy level), the variation in heteroplasmy level in different tissues, and the energy requirements of brain and other tissues, which may vary with age. Intrafamilial and interfamilial clinical variability are seen in individuals with the same genotype [Stendel et al 2020].

Several studies have tried to describe genotype-phenotype correlations in mtDNA-LSS; however, these are generally inconsistent between cohorts, limiting their translation into clinical practice.

MT-ATP6 pathogenic variants, specifically m.8993T>G and m.8993T>C, probably show the strongest genotype-phenotype correlation of any mtDNA pathogenic variants. They show very little tissue-dependent [Ng et al 2019] or age-dependent variation in the proportion of abnormal mtDNA [White et al 1999b] as well as a strong correlation between the heteroplasmy level and disease severity. Logistic regression models were developed that predict the probability of a severe outcome in an individual based on the measured heteroplasmy level of m.8993T>G and m.8993T>C (see Figure 1) [White et al 1999a]. Across all *MT-ATP6* variants, heteroplasmy levels have been shown to have a significant negative association with age of onset of disease [Ganetzky et al 2019]. These findings are supported by Ng et al [2019], who generated logistic regression models to predict the risk of disease manifestation based on the blood heteroplasmy level of *MT-ATP6* pathogenic variants m.8993T>C, m.8993T>G, m.9176T>C, and m.9185T>C (see Figure 2). Note, however, that in such retrospective studies it is not possible to completely avoid ascertainment bias, and the data should be regarded as broadly indicative rather than precise.

MT-ATP6 pathogenic variants m.8993T>G and m.8993T>C tend to be associated with early-onset and severe clinical disease, with m.8993T>G causing a more severe phenotype [Sofou et al 2018, Ganetzky et al 2019,

Stendel et al 2020]. Although the cause of this difference is unknown, it is postulated to result from the instability of the abnormal protein.

- m.8993T>G. Individuals with a heteroplasmy level below 60% are usually asymptomatic or have only mild pigmentary retinopathy or migraine headaches; however, asymptomatic adults with up to 75% abnormal mtDNA have been reported [Ganetzky et al 2019, Ng et al 2019, Stendel et al 2020]. Individuals with moderate heteroplasmy levels (70%-90%) of pathogenic variant m.8993T>G may present with the milder phenotype of NARP, while those with more than 90% abnormal mtDNA typically present with LSS [Ng et al 2019, Stendel et al 2020, Stenton et al 2022].
- m.8993T>C is a less severe variant than m.8993T>G, and generally all symptomatic individuals with m.8993T>C have a heteroplasmy level greater than 90%. However, some asymptomatic individuals with m.8993T>C have also been reported to have a heteroplasmy level greater than 90% [Stendel et al 2020, Stenton et al 2022].

Note: Overlap in *MT-ATP6* pathogenic variant heteroplasmy level is observed between some asymptomatic individuals and others with NARP or mild neurologic manifestations, particularly for variants other than m.8993T>G. There can also be overlap in heteroplasmy levels between some individuals with NARP and others with LSS [Claeys et al 2016, Ganetzky et al 2019, Ng et al 2019, Stendel et al 2020].

Genotype-phenotype correlations are much weaker for other mtDNA pathogenic variants (e.g., m.3243A>G in *MT-TL1*, m.8344A>G in *MT-TK*, m.9176T>C in *MT-ATP6*, m.14459G>A and m.14487T>C in *MT-ND6*, m.10158T>C and m.10191T>C in *MT-ND3*, and m.13513G>A in *MT-ND5*). It is not possible to use heteroplasmy level to predict outcome (e.g., in asymptomatic family members or in prenatal diagnosis) unless the value is near 0% or near 100%. Some individuals with *MT-ND3* and *MT-ND5* pathogenic variants have been reported to have relatively severe phenotypes at low heteroplasmy levels [Wei et al 2018, Ogawa et al 2020, Stenton et al 2022, Na & Lee 2023].

Other genotype-phenotype correlations have been suggested for mtDNA pathogenic variants affecting respiratory chain enzyme complex I, although confirmation of these findings requires further systematic analysis of larger cohorts. Individuals with complex I deficiency are more commonly reported to have cardiac and visual involvement [Sofou et al 2018], with pathogenic variants in *MT-ND5* strongly associated with cardiomyopathy [Stenton et al 2022, Kistol et al 2023]. A retrospective study of neuroimaging in children with mtDNA-LSS suggested that cerebral cortical and white matter involvement is more common in *MT-ND5*- and *MT-ND3*-related LSS [Alves et al 2020]; however, this finding was not supported by a subsequent study [Stenton et al 2022]. Poor prognosis has been associated with *MT-ND5* pathogenic variants in addition to *MT-ATP6* pathogenic variant m.8993T>G [Ogawa et al 2020, Stenton et al 2022].

Individuals with mtDNA-LSS may exhibit overlapping phenotypes with other mitochondrial phenotypes, including MELAS. *MT-ND* pathogenic variants (most commonly involving *MT-ND5* and *MT-ND3*) have been described in multiple individuals with an LSS/MELAS overlapping phenotype in which individuals display clinical and radiologic features of both syndromes [Wei et al 2021].

Penetrance

Penetrance is reduced (see Genotype-Phenotype Correlations).

Nomenclature

Leigh syndrome was originally described in an affected infant as "subacute necrotizing encephalomyelopathy" [Leigh 1951]. Leigh syndrome was historically defined by characteristic neuropathologic features including multiple focal symmetric necrotic lesions in the basal ganglia, thalamus, brain stem, dentate nuclei, and optic

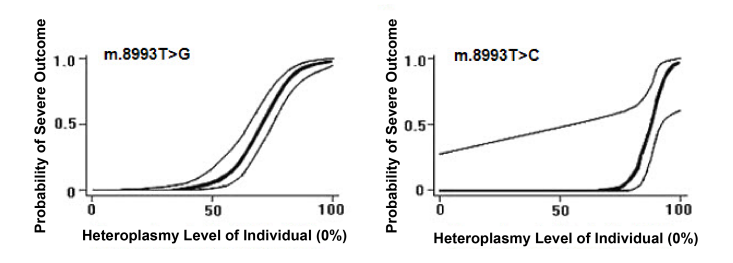


Figure 1. Estimated probability of a severe outcome (95% CI) for an individual with *MT-ATP6* pathogenic variants m.8993T>G or m.8993T>C, based on the heteroplasmy level (i.e., proportion of abnormal mitochondrial DNA) in the individual. A severe outcome is defined as severe manifestations of Leigh syndrome spectrum and probability is calculated using a logistic regression model. Note that the risk of developing severe manifestations is minimal below the approximate threshold value of 50% for m.8993T>G and 75% for m.8993T>C [White et al 1999a].

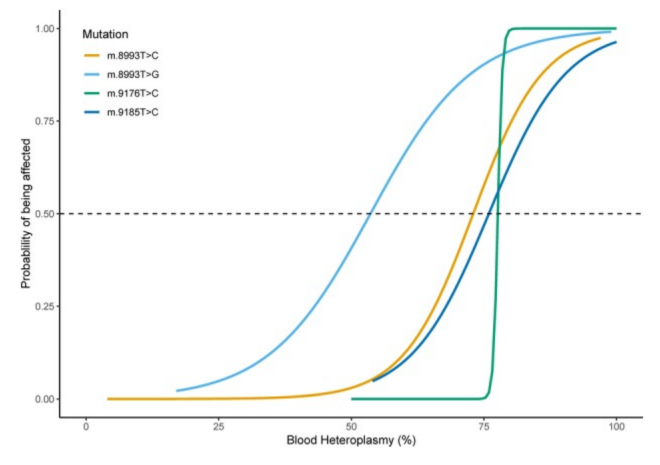


Figure 2. Estimated probability of having clinical manifestations of Leigh syndrome spectrum based on the heteroplasmy level detected in blood for *MT-ATP6* pathogenic variants m.8993T>C, m.8993T>C, m.9176T>C, and m.9185T>C [Ng et al 2019].

nerves that histologically demonstrate a spongiform appearance and are characterized by demyelination, gliosis, and vascular proliferation. The advent of MRI and next-generation sequencing has made it possible to establish the diagnosis without postmortem examination. Individuals with mtDNA-LSS are often referred to as having "maternally inherited Leigh syndrome" (MILS) [Ciafaloni et al 1993]. This term can cause confusion given perhaps one quarter of probands with mtDNA-LSS have *de novo* pathogenic variants (see Genetic Counseling).

The term **Leigh-like syndrome** was used previously when clinical and other features were strongly suggestive of Leigh syndrome but did not fulfill the stringent diagnostic criteria because of atypical neuropathology (i.e., variation in the distribution or character of lesions or the presence of unusual features such as extensive cortical destruction), neuroimaging inconsistent with Leigh syndrome, normal blood and CSF lactate levels, and/or incomplete evaluation.

Leigh syndrome spectrum encompasses both Leigh syndrome and Leigh-like syndrome. Leigh syndrome and Leigh-like syndrome represent a continuum of overlapping clinical and biochemical features and result from pathogenic variants in the same group of mitochondrial and nuclear genes.

Prevalence

LSS is the most frequently observed phenotype of pediatric-onset mitochondrial disorders. Mitochondrial DNA-associated LSS accounts for approximately 30% of LSS.

The following prevalence data are for all genetic causes of LSS. In southeastern Australia, Leigh syndrome occurs in 1:77,000 infants, and the combined birth prevalence of LSS was estimated to be at least 1:40,000 [Rahman et al 1996]. In western Sweden, the prevalence of Leigh syndrome in preschool children was 1:34,000 [Darin et al 2001]. Thus, the prevalence of LSS is likely to be 1:30,000-40,000.

Analyses of 67 individuals with LSS reported by Rahman et al [1996] identified mtDNA pathogenic variants in 34% [Authors, unpublished data]. Hence, the prevalence of mtDNA-LSS is likely to be 1:100,000-140,000.

Genetically Related Disorders

Pathogenic variants in mtDNA genes known to be associated with mitochondrial DNA-associated Leigh syndrome spectrum (mtDNA-LSS) can also be associated with a variety of other mitochondrial disorders. For example, the pathogenic variants most often identified in individuals with Leber hereditary optic neuropathy (m.11778A>G in *MT-ND4* and m.14484T>C in *MT-ND6*) have also been reported in individuals with mtDNA-LSS [Lim et al 2022, Stenton et al 2022]. See Table 3 and Primary Mitochondrial Disorders Overview.

Table 3. Selected Allelic Disorders

Gene	Phenotype ¹
MT-ATP6	Charcot-Marie-Tooth-like peripheral neuropathy ²
	Spinocerebellar ataxia w/upper motor neuron signs ³
	Bilateral striatal necrosis (OMIM 500003)
	Myopathy, lactic acidosis, & sideroblastic anemia (OMIM 500011)
MT-CO1 Cytochrome c oxidase (COX) deficiency (OMIM 516030)	
MT-CO2	COX deficiency (OMIM 516040)
W11-CO2	MELAS

Table 3. continued from previous page.

,			
Gene	Phenotype ¹		
	COX deficiency (OMIM 516050)		
MT-CO3	Leber hereditary optic neuropathy		
	MELAS		
MT-ND1	Leber hereditary optic neuropathy		
WII-IVDI	MELAS		
MT-ND2	Leber hereditary optic neuropathy		
MT-ND3	Leber hereditary optic neuropathy		
1711-17103	MELAS ⁴		
MT-ND4	Leber hereditary optic neuropathy		
WII-IND4	MELAS ⁵		
MT-ND5	Leber hereditary optic neuropathy		
WII-ND3	MELAS		
MT-ND6	Leber hereditary optic neuropathy		
WIT-INDO	MELAS		
MT-TI	MERRF		
MT-TK	MELAS		
WII-IK	MERRF		
	Progressive external ophthalmoplegia (OMIM 590050)		
	Maternally inherited diabetes mellitus w/ or w/o deafness (OMIM 590050)		
MT-TL1	Cardiomyopathy (OMIM 590050)		
IVI I - I L.I	Nonsyndromic hearing loss and deafness		
	MELAS		
	MERRF		
MT-TV	MELAS		
MT-TW	MELAS		

- 1. See hyperlinked *GeneReview*, OMIM phenotype entry, or cited reference for more information.
- 2. Pitceathly et al [2012], Ganetzky et al [2019], Ng et al [2019]
- 3. Pfeffer et al [2012a], Ganetzky et al [2019], Ng et al [2019]
- 4. Mezuki et al [2017]
- 5. Lin et al [2020]

Differential Diagnosis

Leigh syndrome spectrum (LSS) is most often caused by a pathogenic variant in a nuclear gene. Pathogenic variants in 98 nuclear genes have been associated with autosomal recessive, autosomal dominant, and X-linked nuclear gene-encoded LSS [McCormick et al 2023] (see Causes of Nuclear Gene-Encoded Leigh Syndrome Spectrum).

Other monogenic disorders that cause or resemble LSS are listed in Table 4.

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Table 4. Other Monogenic Disorders That Cause or Resemble Leigh Syndrome Spectrum

Key Feature	Gene(s)	Disorder	MOI
Information below and the standard and t	ADAR	ADAR-related Aicardi-Goutières syndrome	AR AD ¹
Infantile bilateral striatal necrosis	NUP62	<i>NUP62</i> -related infantile bilateral striatal necrosis (OMIM 605815)	AR
Infection-induced acute encephalopathy	RANBP2	<i>RANBP2</i> -related susceptibility to infection-induced acute encephalopathy 3 (OMIM 608033)	AD
	ATP7B	Wilson disease	AR
	FTL	Neuroferritinopathy	AD
	GAMT	GAMT deficiency (See Creatine Deficiency Disorders.)	AR
	GCDH	Glutaric acidemia type 1	AR
Neurodegenerative &/or neurodevelopmental disorders w/similar changes on neuroimaging	MCEE MMAA MMAB MMADHC MMUT	Isolated methylmalonic acidemia	AR
	MORC2	Developmental delay, impaired growth, dysmorphic facies, & axonal neuropathy (OMIM 619090)	AD
	PANK2	Pantothenate kinase-associated neurodegeneration	AR
	PCCA PCCB	Propionic acidemia	AR

AD = autosomal dominant; AR = autosomal recessive; MOI = mode of inheritance

1. ADAR-related Aicardi-Goutières syndrome can be inherited in an autosomal recessive or autosomal dominant manner depending on the specific pathogenic variant.

Acquired conditions that cause or resemble LSS with similar clinical features and/or similar neuroimaging findings include the following:

- Acute necrotizing encephalopathy (may be triggered by viral infections)
- Viral encephalopathies
- Hypoxic-ischemic encephalopathy
- Wernicke encephalopathy (thiamine deficiency)

Management

Evaluation Following Initial Diagnosis

To establish the extent of disease and needs in an individual diagnosed with mitochondrial DNA-associated Leigh syndrome spectrum (mtDNA-LSS), the evaluations summarized in Table 5 (if not performed as part of the evaluation that led to the diagnosis) are recommended.

Table 5. Mitochondrial DNA-Associated Leigh Syndrome Spectrum: Recommended Evaluations Following Initial Diagnosis

System/Concern	Evaluation	Comment
Neurologic	 Neurologic eval Brain MRI & MRS Plasma & CSF lactate & pyruvate Urine organic acids 	 Consider EEG if seizures are suspected. Consider nerve conduction studies if neuropathy is suspected (due to history or √/absent reflexes on clinical exam).

Table 5. continued from previous page.

System/Concern	Evaluation	Comment
Pulmonology	Assess for apnea, hyperventilation, or irregular respiration.	Consider referral for sleep study.
Feeding/ Nutrition	Measurement of growth parametersNutrition / feeding team eval	To incl eval of nutritional status & aspiration risk w/ consideration of radiographic assessment of swallow
Development/ Cognition	Developmental &/or cognitive assessment	Eval for early intervention / special education in children
Musculoskeletal	PT & OT eval	 To incl assessment of: Gross motor & fine motor skills Mobility, ADL, & need for adaptive devices Need for PT (to improve gross motor skills) &/or OT (to improve fine motor skills)
Eyes	Ophthalmologic eval	To assess for reduced vision, abnormal ocular movement, strabismus, & more complex findings (e.g., retinal dystrophy) that may require referral for subspecialty care &/or low vision services
Cardiology	Blood pressureEKGEchocardiogram	 Referral to cardiologist to screen for conduction defects, arrythmias, or cardiomyopathy. Consider Holter monitoring in high-risk individuals.
Liver	 Liver function tests incl transaminases, bilirubin, albumin, & coagulation studies Consider abdominal ultrasound. 	To assess for liver dysfunction
Renal	 Urinalysis, urine albumin-to- creatinine ratio, urine amino acids Serum electrolytes, BUN, creatinine 	To assess for renal dysfunction
Hearing	Audiologic eval	To assess for hearing loss
Hematology	Complete blood count	To assess for anemia
Endocrine	Random glucose	Consider hemoglobin A1c or oral glucose tolerance test depending on clinical history to screen for diabetes mellitus.
Psychosocial	Assess for anxiety & other psychosocial manifestations	
Genetic counseling	By genetics professionals ¹	To inform affected persons & their families re nature, MOI, & implications of mtDNA-LSS to facilitate medical & personal decision making
Family support & resources	By clinicians, wider care team, & family support organizations	Assessment of family & social structure to determine need for: • Community or online resources such as Parent to Parent • Social work involvement for parental support • Home nursing referral

ADL = activities of daily living; BUN = blood urea nitrogen; CSF = cerebrospinal fluid; MOI = mode of inheritance; MRS = magnetic resonance spectroscopy; mtDNA-LSS = mitochondrial DNA-associated Leigh syndrome spectrum; OT = occupational therapy; PT = physical therapy

1. Medical geneticist, certified genetic counselor, certified advanced genetic nurse

Treatment of Manifestations

As for most mitochondrial disorders, no specific curative treatment for mtDNA-LSS exists [Pitceathly et al 2021], and currently there are no licensed or approved treatments for mtDNA-LSS. Supportive care to improve quality of life, maximize function, and reduce complications is recommended. This ideally involves multidisciplinary care by specialists in relevant fields (see Table 6).

Table 6. Mitochondrial DNA-Associated Leigh Syndrome Spectrum: Treatment of Manifestations

Manifestation/Concern	Treatment	Considerations/Other
Acute acidosis	 Sodium bicarbonate or sodium citrate for significant acidosis Consider THAM if accompanying hypernatremia. 	Avoid lactate containing solutions (e.g., Ringer lactate).
Epilepsy	Treatment w/ASM by experienced neurologist	 Sodium valproate should be avoided because of its inhibitory effects on mitochondrial respiratory chain. ¹ Education of parents/caregivers ²
	Benzhexol, baclofen, tetrabenazine, & gabapentin may be useful, alone or in various combinations.	An initial low dose should be started & gradually \(^1\) until symptom control is achieved, or intolerable side effects occur.
Dystonia	Botulinum toxin injection has been used in persons w/LSS & severe intractable dystonia.	Use w/caution given potential side effects of exacerbating muscle weakness; & do not use in head & neck because of risk of adverse events (e.g., impaired swallow).
Pulmonology	 Referral to pulmonologist as needed Ventilatory support for persons w/respiratory compromise 	
Feeding/Nutrition	 Caloric & nutritional supplementation as needed (incl micronutrients) Feeding therapy as needed Gastrostomy tube placement may improve nutritional intake for those w/persistent feeding issues, choking, or aspiration risk due to dysphagia. 	Low threshold for clinical feeding eval &/or radiographic swallowing study when showing clinical signs or symptoms of dysphagia
Development/Cognition	See Developmental Delay / Intellectual Disability Management Issues.	
Myopathy &/or ataxia	PT & OT	Equipment to prevent falling & maintain independent mobility as appropriate
Eyes	Standard treatments for eye movement disorders as advised by ophthalmologist	
Cardiomyopathy	Anticongestive therapy or antiarrhythmic therapy may be required & treatment should be managed by cardiologist.	
Gastrointestinal	Stool softeners, laxatives for constipation as needed	
Liver	Treatments for liver failure as advised by hepatologist	
Renal	Electrolyte monitoring & replacement for renal tubular losses per nephrologist	
Hearing	 Hearing aids or cochlear implants for sensorineural hearing loss Referral to speech therapy Referral to hearing support services 	Counsel to avoid excessive noise exposure, which can exacerbate hearing loss.

Table 6. continued from previous page.

Manifestation/Concern	Treatment	Considerations/Other
Endocrine	Treatment for diabetes mellitus or adrenal insufficiency managed by endocrinologist	
Psychosocial	Standard treatments of anxiety &/or depression	
Family/Community	 Ensure appropriate social work involvement to connect families w/local resources, respite, & support. Coordinate care to manage multiple subspecialty appointments, equipment, medications, & supplies. 	Ongoing assessment of need for palliative care involvement &/or home nursing

 $ASM = anti-seizure \ medication; \ OT = occupational \ therapy; \ PT = physical \ therapy; \ THAM = tris-hydroxymethyl \ aminomethane$

Surveillance

To monitor existing manifestations, the individual's response to supportive care, and the emergence of new manifestations, the evaluations summarized in Table 7 are recommended.

Table 7. Mitochondrial DNA-Associated Leigh Syndrome Spectrum: Recommended Surveillance

System/Concern	Evaluation	Frequency	
Neurologic	Neurology assessment for ataxia/movement disorders, neuropathy, & seizures		
Pulmonary Monitor for evidence of aspiration, nocturnal hypoventilation, or sleep-disordered breathing.		At each visit	
Feeding/Nutrition	 Measurement of growth parameters 		
Gastroenterology	Assess for nausea, constipation, or dysmotility.		
Development/ Cognition	Monitor developmental progress, educational needs, & cognitive issues.	At least annually	
Musculoskeletal	Physical medicine, OT/PT assessment of mobility, self-help skills		
Ophthalmology	Ophthalmologic eval	Every 6-12 mos or as advised by ophthalmologist	
Cardiac	Blood pressureEKGEchocardiogram	Annually or as advised by cardiologist	

^{1.} De Vries et al [2020]

^{2.} Education of parents/caregivers regarding common seizure presentations is appropriate. For information on non-medical interventions and coping strategies for children diagnosed with epilepsy, see Epilepsy Foundation Toolbox.

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Table 7. continued from previous page.

System/Concern	Evaluation	Frequency
Liver	Liver function tests incl transaminases, albumin, bilirubin, & coagulation studies	
Renal	 Urinalysis, urine albumin-to-creatinine ratio, urine amino acids Serum electrolytes, BUN, creatinine 	
Hearing	Audiologic eval	. "
Hematology	Complete blood count	Annually
Endocrine	 Fasting glucose Consider hemoglobin A1c or OGTT if clinical features consistent w/diabetes mellitus. Consider early-morning cortisol & short synacthen test if clinical symptoms of adrenal insufficiency. 	
Family/Community	Assess family need for social work support (e.g., palliative/respite care, home nursing, other local resources), care coordination, or follow-up genetic counseling if new questions arise (e.g., family planning).	At each visit

BUN = blood urea nitrogen; OGTT = oral glucose tolerance test; OT = occupational therapy; PT = physical therapy

Agents/Circumstances to Avoid

An international Delphi-based consensus on the safety of medication use in individuals with primary mitochondrial disorders was published in 2020 to assist clinical decision making. However, it is important to tailor medication recommendations to the individual [De Vries et al 2020].

Sodium valproate should be avoided if possible, because of its inhibitory effect on respiratory chain enzymes.

Medications that can cause acidosis should be avoided or, if necessary, prescribed with caution, with frequent monitoring of blood acid-base levels.

Anesthesia can potentially aggravate respiratory symptoms and precipitate respiratory failure, so careful consideration should be given to its use and to monitoring of the individual prior to, during, and after anesthetic procedures [Parikh et al 2017]. Catabolism should be prevented by minimizing preoperative fasting and considering intravenous glucose perioperatively during prolonged anesthesia (unless the individual is on a ketogenic diet). Prolonged propofol use during maintenance anesthesia may increase the risk of lactic acidosis. Neuromuscular blocking drugs should be avoided in individuals with muscle disease, or if necessary, under strict monitoring. Avoid lactate-containing agents (e.g., Ringer lactate) in those with risk of lactic acidosis [Parikh et al 2017, De Vries et al 2020].

Dichloroacetate (DCA) reduces blood lactate by activating the pyruvate dehydrogenase complex. Mixed clinical outcomes have been observed following the use of DCA in individuals with mitochondrial disorders. DCA was reported to worsen peripheral neuropathy in a randomized clinical trial in individuals with MELAS [Kaufmann et al 2006]. It is therefore recommended that DCA should be used with caution in individuals with LSS, particularly in individuals with established peripheral neuropathy or with a genetic cause associated with increased risk of peripheral neuropathy.

Evaluation of Relatives at Risk

See Genetic Counseling for issues related to testing of at-risk relatives for genetic counseling purposes.

Therapies Under Investigation

A number of vitamins, cofactors, and other compounds, including riboflavin, thiamine, creatine, biotin, coenzyme Q_{10} , carnitine, and alpha-lipoic acid, have been prescribed in individuals with mtDNA-LSS. To date, there is no consistency in prescribing practice between international centers. Despite the theoretic potential, there is no established evidence base for any of these compounds in mtDNA-LSS [Pfeffer et al 2012b]. Complex phenotypic and genetic heterogeneity, as well as a lack of robust clinical outcome measures, are some of the obstacles to developing effective therapies for mtDNA-LSS. Evidence for some of these therapies has been shown in specific vitamin/cofactor metabolism and transporter defects in nuclear gene-encoded Leigh syndrome spectrum. In recent years, there has been an increase in clinical trials for pharmacologic and non-pharmacologic therapies for mtDNA-LSS.

Antioxidants such as coenzyme Q_{10} and its analogs have demonstrated potential antioxidant activities against toxic metabolites and accumulated reactive oxygen species. Apart from disorders of coenzyme Q_{10} biosynthesis, coenzyme Q_{10} has limited evidence for efficacy in the treatment of LSS.

Vatiquinone (PTC-743; previously known as EPI-743) is a structurally modified variant of coenzyme Q₁₀ that may help to replenish intracellular glutathione levels [Shrader et al 2011]. An open-label clinical trial in an end-of-life setting in 14 individuals with primary mitochondrial disorders (including mtDNA-LSS) showed clinical improvement in 11 of the 14 individuals, which corresponded to increased regional and whole-brain technetium-99m hexamethylpropylene oxime uptake on single-photon emission computed tomography [Enns et al 2012]. An open-label Phase IIA single-center clinical trial of vatiquinone therapy showed stabilization of disease progression in ten children with mtDNA-LSS or nuclear gene-encoded LSS [Martinelli et al 2012]. The unpredictable natural history of mtDNA-LSS causes difficulty in interpretation of open-label studies. A Phase IIB randomized placebo-controlled double-blind crossover clinical trial of vatiquinone in children with mtDNA-LSS or nuclear gene-encoded LSS was completed in May 2015 (NCT01721733), and a long-term safety and efficacy evaluation of vatiquinone in children with mtDNA-LSS or nuclear gene-encoded LSS was completed in late 2023 (NCT02352896). The findings of these studies have not yet been published. A recent clinical trial of vatiquinone in individuals with mitochondrial disorders with associated epilepsy phenotype (NCT04378075) failed to meet its primary end point (see PTC Therapeutics press release).

KH176 (sonlicromanol) is a redox-modulating agent that targets the thioredoxin/peroxiredoxin system to reduce cellular reactive oxygen species. Promising preclinical studies in a mouse model for LSS [de Haas et al 2017] led to a Phase I study, which reported that KH176 was tolerated but at higher doses was associated with changes in cardiac electrophysiology (QTc prolongation and T wave morphologic changes) [Koene et al 2017]. A Phase II randomized, double-blind, placebo-controlled, parallel group clinical trial in children with genetically confirmed primary mitochondrial disorders with motor symptoms is currently recruiting (NCT04846036).

Gene therapy. Gene-editing techniques with mitochondrial targeted restriction endonucleases (mitoREs) or programmable nucleases, such as mitochondrial targeted zinc finger nucleases (mtZFN) and transcription activator-like effector nucleases (mito-TALEN), are postulated to shift heteroplasmy toward wild type mtDNA by inducing mtDNA double-stranded breaks, leading to rapid degradation of mutated mtDNA. Gene-editing techniques that introduce specific *de novo* nucleotide changes in mtDNA include bacterial cytidine deaminases. However, to date, gene therapy has not progressed to clinical trials in humans.

Search ClinicalTrials.gov in the US and EU Clinical Trials Register in Europe for access to information on clinical studies for a wide range of diseases and conditions.

Other

Ketogenic diet (KD) has been proposed as a therapy for mitochondrial disorders, in particular complex I deficiency [Paleologou et al 2017]. Although this high-fat, low-carbohydrate diet has proven efficacy for

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refractory epilepsies, high-quality evidence for its use in the treatment of primary mitochondrial disorders, including mtDNA-LSS, is currently lacking. Studies in patient-derived fibroblasts have postulated that KD may be beneficial in mitochondrial disorders [Santra et al 2004]. A systematic review on the use of KD in mitochondrial disorders highlighted the paucity of high-quality studies but suggested that KD can be considered in individuals with mitochondrial disorders (including mtDNA-LSS) and therapy-refractory epilepsy (except mtDNA deletion-related myopathy and pyruvate carboxylase deficiency) [Zweers et al 2021]. A prospective controlled study in individuals with mitochondrial disorders (including one individual with LSS) found that KD reduced seizure frequency in 76%, with a high efficacy in individuals with MELAS [Huang et al 2022]. Further clinical trials are required to determine the efficacy of KD in mtDNA-LSS. Another proposed option is supplementation with the fatty acid decanoic acid, which is thought to be the active component of the KD and appears to stimulate mitochondrial biogenesis in cell models [Hughes et al 2014], including fibroblasts from individuals with complex I-deficient nuclear gene-encoded LSS [Kanabus et al 2016]. A prospective open-label feasibility study of K.Vita, a medical food containing a unique ratio of decanoic acid to octanoic acid, had a beneficial effect on frequency of seizures, including in individuals with primary mitochondrial disorders [Schoeler et al 2021]. Further clinical trials are needed.

Mitochondrial biogenesis is the process in which cells increase the mitochondrial mass. Several studies have investigated whether upregulation of mitochondrial biogenesis may provide an effective therapeutic approach for mitochondrial respiratory chain disorders. This approach involves using agonists such as bezafibrate or resveratrol to stimulate the peroxisome proliferator-activated receptor gamma (PPARgamma) coactivator alpha (PGC-1a) pathway [Bastin et al 2008, Steele et al 2020]. Alternatively, nicotinamide analogs such as nicotinamide riboside or nicotinamide mononucleotide have been used to boost reduced nicotinamide adenine dinucleotide (NAD+) levels and induce mitochondrial biogenesis via the same PGC-1a pathway [Cerutti et al 2014, Khan et al 2014, Felici et al 2015, Long et al 2015]. KL1333, a novel NAD+ modulator, interacts with NAD(P)H:dehydrogenase 1 (NQO1) to increase intracellular NAD+ levels via NADH oxidation [Seo et al 2018]. Results of a Phase Ia/Ib trial assessing the safety and tolerability in individuals with primary mitochondrial disorders are yet to be published (NCT03888716). Further studies are needed to investigate the safety and efficacy of these therapies in mtDNA-LSS.

Genetic Counseling

Genetic counseling is the process of providing individuals and families with information on the nature, mode(s) of inheritance, and implications of genetic disorders to help them make informed medical and personal decisions. The following section deals with genetic risk assessment and the use of family history and genetic testing to clarify genetic status for family members; it is not meant to address all personal, cultural, or ethical issues that may arise or to substitute for consultation with a genetics professional. —ED.

Mode of Inheritance

Mitochondrial DNA-associated Leigh syndrome spectrum (mtDNA-LSS) is transmitted by mitochondrial (maternal) inheritance.

Risk to Family Members

Parents of a proband

- The father of a proband is not at risk of having the mtDNA pathogenic variant.
- The mother of a proband may have the mtDNA pathogenic variant and may exhibit mild clinical manifestations of mtDNA-LSS.

- In most instances, the mother has a much lower heteroplasmy level (i.e., proportion of abnormal mtDNA) than the proband and usually remains asymptomatic or develops only mild manifestations such as gait disturbance or retinopathy.
- Occasionally the mother has a substantial heteroplasmy level and develops severe manifestations of mtDNA-LSS in adulthood [de Vries et al 1993, Wei et al 2018].
- With the exception of the *MT-ATP6* pathogenic variants m.8993T>G and m.8993T>C, a lower heteroplasmy level in maternal blood does not exclude a higher heteroplasmy level in tissues such as brain or muscle.
- Many affected individuals have no known family history of mtDNA-LSS or other mitochondrial disorders. The explanation for apparently simplex cases may be absence of a comprehensive and/or reliable family history, a maternal heteroplasmy level below the threshold level that causes disease, or a *de novo* mtDNA pathogenic variant in the proband. A study of 105 probands with childhood-onset mitochondrial disease, including mtDNA-LSS, suggested that approximately 25% of individuals had a *de novo* pathogenic variant [Sallevelt et al 2017].
- Molecular genetic testing of the mother for the mtDNA pathogenic variant identified in the proband is recommended. Assessment in at least two different maternal tissues (e.g., blood, urine sediment cells, buccal cells) is suggested [Mavraki et al 2023]. Testing of maternal relatives can be considered to provide reassurance or appropriate surveillance and genetic counselling. (Note: Mitochondrial DNA pathogenic variants may be lost from the leukocyte population with increasing age [Schon et al 2021, Davis et al 2022]; therefore, in adults with milder manifestations and for asymptomatic older maternal relatives, the pathogenic variant may only be detected in tissues such as hair follicles, urine sediment cells, or skeletal muscle.)
- If the proband represents a simplex case (i.e., the only family member known to be affected) and the mtDNA pathogenic variant identified in the proband cannot be identified in maternal tissues, possibilities to consider include a *de novo* pathogenic variant in the proband and germline (gonadal) mosaicism in the mother. Care should be used when determining if a pathogenic variant is *de novo*. For some pathogenic variants, homoplasmic (or apparently homoplasmic levels) in an affected child may result from very low heteroplasmy levels in the mother.

Sibs of a proband. The risk to the sibs depends on the genetic status of the mother.

- If the mother of the proband has the mtDNA pathogenic variant identified in the proband, all sibs are at risk of inheriting the pathogenic variant. Sibs may inherit the pathogenic variant at varying heteroplasmy levels due to the bottleneck effect and variant-specific segregation patterns.
 - For the *MT-ATP6* pathogenic variants m.8993T>G and m.8993T>C, a strong positive relationship exists between the mother's heteroplasmy level and the predicted recurrence risk [White et al 1999a] (see Figure 3). However, the 95% confidence interval of the risk estimates are wide, and these data may be of limited use for genetic counseling.
 - Due to positive selection, the m.8993T>G pathogenic variant exhibits skewed segregation [Otten et al 2018], with the heteroplasmy levels in oocytes/embryos generally displaying a distribution to the extremes, either 0% or approximately 100%.
- The risk to a sib of developing clinical manifestations is difficult to determine and depends on heteroplasmy level, the variation in heteroplasmy levels among different tissues, and the disease threshold (i.e., the minimum level of heteroplasmy expected to result in mitochondrial disease) for the specific variant (see Genotype-Phenotype Correlations).
 - In mtDNA-LSS, the *MT-ATP6* pathogenic variants m.8993T>G and m.8993T>C probably exhibit the strongest genotype-phenotype correlation of any mtDNA pathogenic variants [Ng et al 2019]; they show very little tissue-dependent or age-dependent variation in heteroplasmy level as well as a strong correlation between heteroplasmy level and clinical outcome.

- It is difficult to use heteroplasmy level to predict clinical outcome (e.g., in asymptomatic family members or in prenatal diagnosis) for other mtDNA pathogenic variants detected in individuals known to be associated with mtDNA-LSS unless the value is near 0% or near 100%. Retrospective studies for some of the more common mtDNA pathogenic variants can be used to indicate an approximate (empiric) recurrence risk. See Chinnery et al [1998] and Pickett et al [2019] for *MT-TL1* pathogenic variant m.3243A>G and *MT-TK* pathogenic variant m.8344A>G and Sallevelt et al [2017], and Steffann et al [2021], and Vachin et al [2018] for other mtDNA pathogenic variants.
- Variation in clinical manifestations is seen among family members with the same genotype [Stendel et al 2020].
- If the proband is presumed to have a *de novo* mtDNA pathogenic variant (i.e., the pathogenic variant is absent from the blood and at least one other sample from the mother and any maternal relatives who were tested), sibs are at low risk [Sallevelt et al 2017]. However, the possibility of maternal germline (gonadal) mosaicism cannot be excluded and therefore the recurrence risk is not 0%.

Offspring of a proband

- Offspring of a male proband with an mtDNA pathogenic variant are not at risk.
- All offspring of a female proband with an mtDNA pathogenic variant are at risk of inheriting the pathogenic variant. Offspring may inherit the pathogenic variant at varying heteroplasmy levels due to the bottleneck effect and variant-specific segregation patterns.
- The risk to offspring of a female proband of developing clinical manifestations is difficult to determine and depends on the maternal heteroplasmy level, the variation in heteroplasmy levels among different tissues in the proband, and the specific mtDNA pathogenic variant. Retrospective studies for some of the most common mtDNA pathogenic variants can be used to indicate an approximate (empiric) recurrence risk for the offspring of females. See White et al [1999a] for the *MT-ATP6* pathogenic variants m.8993T>G and m.8993T>C, Chinnery et al [1998] and Pickett et al [2019] for the *MT-TL1* pathogenic variant m.3243A>G and *MT-TK* pathogenic variant m.8344A>G, and Sallevelt et al [2017] and Steffann et al [2021] for other mtDNA pathogenic variants.

Other family members. The risk to other family members depends on the genetic status of the proband's mother: if the proband's mother has an mtDNA pathogenic variant, her sibs and mother are also at risk.

Related Genetic Counseling Issues

Family planning

- The optimal time for determination of genetic risk and discussion of the availability of prenatal/ preimplantation genetic testing is before pregnancy. Similarly, decisions regarding testing to determine the genetic status of at-risk asymptomatic family members are best made before pregnancy.
- It is appropriate to offer genetic counseling (including general discussion of potential risks to offspring and reproductive options) to young adults who are affected or at risk. It should be emphasized in genetic counseling for mtDNA-LSS that it is not possible to make specific predictions about clinical outcome in individuals or their offspring.

Phenotypic variability. The phenotype of an individual with an mtDNA pathogenic variant results from a combination of factors, including the severity of the pathogenic variant, the percentage of abnormal mitochondria (mtDNA heteroplasmy), and the organs and tissues in which the pathogenic variant is found (tissue distribution). Family members often inherit different percentages of abnormal mtDNA and therefore can have a wide range of clinical manifestations.

Testing in asymptomatic at-risk family members. Testing to evaluate the genetic status of an apparently asymptomatic family member is technically possible once the mtDNA pathogenic variant has been identified in

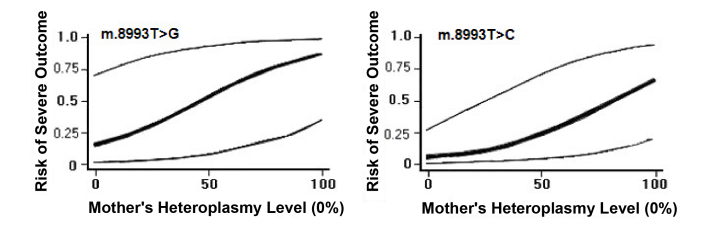


Figure 3. Predicted risk to offspring (95% CI) to develop clinical manifestations of Leigh syndrome spectrum due to *MT-ATP6* pathogenic variants m.8993T>G or m.8993T>C based on the heteroplasmy level detected in blood in the mother [White et al 1999a]

the proband. However, interpretation of test results of asymptomatic at-risk family members is extremely difficult. Prediction of phenotype based on test results is not possible. Furthermore, absence of the mtDNA pathogenic variant in one tissue (e.g., blood) does not guarantee that the pathogenic variant is absent in other tissues.

Prenatal Testing and Preimplantation Genetic Testing

Recurrence risk assessment and prenatal testing for disorders caused by pathogenic variants in mtDNA are challenging due to the intricacies of mtDNA transmission, such as the mtDNA bottleneck effect (resulting in variation in heteroplasmy levels between generations), pathogenic variant-specific selection, the threshold effect (the heteroplasmy level associated with clinical manifestations of mitochondrial disease), and the inherent challenge in using prenatal genetic test results to predict clinical outcome.

The European Neuromuscular Disease Centre International Consensus Workshops have been instrumental in developing guidelines for the counseling of reproductive options for families with mitochondrial disease [Poulton & Turnbull 2000, Poulton & Bredenoord 2010, Poulton et al 2019].

Reproductive options for the family members of a proband with an mtDNA pathogenic variant may include prenatal testing, preimplantation genetic testing (PGT), and oocyte donation. Prenatal testing appears to be the preferred reproductive testing option for mtDNA pathogenic variants that appear to have occurred *de novo* in the proband and are associated with low recurrence risk [Sallevelt et al 2017], as there is a high likelihood that the pathogenic variant will not be present in prenatal samples. Prenatal testing may also be suitable for women thought to have low recurrence risk due to having non-zero but low levels of the pathogenic mtDNA variant, such as <20% heteroplasmy. PGT is currently considered an appropriate reproductive option for females with familial heteroplasmic mtDNA pathogenic variants and is likely the best option available, at present, for those with moderate recurrence risk. However, PGT may not be successful for women at moderate risk and is not recommended for females containing homoplasmic or near homoplasmic levels of an mtDNA pathogenic variant [Poulton et al 2019]. For families in which prenatal testing and/or PGT are unlikely to successfully reduce recurrence risk, oocyte donation may be considered.

Prenatal testing involves quantification of the heteroplasmy level in fetal tissue obtained by chorionic villus biopsy (CVS) (usually performed between 11-12 weeks' gestation) or amniocentesis (usually performed at 15-18 weeks' gestation).

- Available evidence suggests that the proportion of abnormal mtDNA in extraembryonic and embryonic tissues is similar and does not change substantially during pregnancy. For a number of mtDNA pathogenic variants, including m.8993T>G, m.8993T>C, and m.9176T>C in *MT-ATP6*, m.3243A>G in *MT-TL1*, m.10191T>C in *MT-ND3*, and m.13513G>A in *MT-ND5*, the proportion of abnormal mtDNA has been reported to remain relatively stable throughout embryofetal development and in postnatal tissue [White et al 1999a, Dahl et al 2000, Thorburn & Dahl 2001, Jacobs et al 2005, Steffann et al 2007, Monnot et al 2011, Steffann et al 2021]. However, variation in mtDNA heteroplasmy levels between the placenta and embryonic/postnatal tissue or between multiple placental samples has also been reported [Marchington et al 2006, Monnot et al 2011, Vachin et al 2018, Steffann et al 2021].
- There is no consensus between centers regarding the preference for CVS or amniocentesis for prenatal testing for mtDNA disease, although some studies support the use of amniocentesis given reports of intraplacental variation with CVS [Steffann et al 2021].
- Analysis should be done on direct CVS/amniocytes, not on cultured cells, as the heteroplasmy level of mtDNA pathogenic variants can change during cell culture.

The major limitation of prenatal testing is the potential difficulty in predicting clinical outcome based on fetal genetic testing results. For prediction of clinical outcome to be reliable, there must be a close correlation between the heteroplasmy level and disease severity; uniform distribution of heteroplasmy levels in all tissues; and stability of the heteroplasmy level over time [Poulton & Turnbull 2000]. See Genotype-Phenotype Correlations for pathogenic variant-specific information.

Clinical practice varies in determining which thresholds are used to determine the likelihood a fetus will be affected or unaffected based on prenatal test results. A systematic review suggested a generic threshold of 18% after it was found that individuals with muscle heteroplasmy levels below approximately 18% had a 95% chance or higher of being unaffected, independent of the mtDNA variant [Hellebrekers et al 2012]. Prenatal heteroplasmy risk groups of low (<30%), intermediate (30%-60%) and high (>60%) have been applied in different studies [Nesbitt et al 2014, Vachin et al 2018, Steffann et al 2021]. An intermediate proportion of abnormal DNA would represent a "gray zone" in which interpretation is difficult or impossible. For some pathogenic variants, using this risk stratification is feasible; however, it may not be appropriate for other pathogenic variants, such as m.3243A>G in *MT-TL1* [Hellebrekers et al 2012].

- While the *MT-ATP6* pathogenic variant m.8993T>G is known to exhibit skewed segregation in embryos or oocytes (either 0% or approximately 100%) [Blok et al 1997, Otten et al 2018], prenatal testing is still considered a suitable option for females with low heteroplasmy levels given that a reliable prediction is possible for most heteroplasmy levels [Smeets et al 2015].
- A model has been developed for the *MT-TL1* m.3243A>G pathogenic variant to facilitate predictions of fetal heteroplasmy (below a specified threshold of 18%) based on maternal heteroplasmy levels [Pickett et al 2019].
- For other mtDNA pathogenic variants, prenatal diagnosis may be offered to females with a low proportion of abnormal mtDNA. However, weaker genotype-phenotype correlation or lack of data would mean that accurate prediction of the phenotype may not possible. Couples would require careful counseling before embarking on these procedures.

Preimplantation genetic testing (PGT) involves quantification of the heteroplasmic variant load to facilitate selection of the most appropriate embryo for transfer. PGT is traditionally performed by sampling one or two blastomeres on day three or four (cleavage stage) after fertilization. At the cleavage stage, there are robust data supporting the correlation between the heteroplasmic variant load in individual blastomeres and the whole

embryo [Steffann et al 2006, Monnot et al 2011, Sallevelt et al 2013]. Most centers offering PGT for inherited diseases now use trophectoderm biopsy rather than cleavage stage biopsy. There are limited data using trophectoderm biopsy for mtDNA PGT, with one study suggesting it may be less suitable for estimating heteroplasmy in the embryo [Mitalipov et al 2014], although this remains unclear [Poulton et al 2019].

Embryos should only be regarded as suitable for implantation if they have a very low proportion of abnormal mtDNA, preferably 0%, although threshold levels of 18% may imply low recurrence risk [Hellebrekers et al 2012, Smeets et al 2015]. Nevertheless, as disease thresholds and patterns of transmission vary with different mtDNA pathogenic variants, it is important to assess this on an individual basis, exploring the couple's own views on what level of residual risk they are willing to accept. Heteroplasmy levels of clinically affected and unaffected individuals within a family should also be considered [Poulton et al 2019].

In some females, particularly those with a high proportion of abnormal mtDNA in blood or urine sediment cells, a large proportion of oocytes may have a high level of abnormal mtDNA, and multiple cycles of ovarian stimulation may not result in an embryo suitable for implantation. However, PGT for mtDNA pathogenic variants may provide valuable information for future reproductive planning even if a successful unaffected conception is not achieved.

- If most of the embryos tested have a substantial proportion of abnormal mtDNA, oocyte donation is likely to be the only current option for ensuring an unaffected embryo.
- In contrast, if most of the embryos tested have undetectable abnormal mtDNA, the parents may opt for prenatal testing in subsequent unassisted (natural) pregnancies.

Other Reproductive Options

Oocyte donation accompanied by IVF using the partner's sperm. Use of a maternal relative as the oocyte donor should be avoided, since the relative may have oocytes with a high proportion of abnormal mtDNA even though her leukocytes may lack detectable abnormal mtDNA.

Mitochondrial replacement therapy (MRT) involves the transfer of a nucleus removed from either a zygote (pronuclear transfer) or an oocyte (maternal spindle transfer) into an enucleated cell at the same embryonic stage from an unaffected donor. Clinical trials are currently under way to investigate whether this reproductive option would be suitable for females with pathogenic mtDNA variants where PGT is not likely to be (or has not been) an effective option (e.g., homoplasmic variants and/or heteroplasmic variants at high heteroplasmy levels).

The first child born after maternal spindle transfer was reported in 2017, born to a female with *MT-ATP6* pathogenic variant m.8993T>G [Zhang et al 2017]. The boy was reported to be well at age seven months but long-term neurodevelopmental follow up has yet to be published.

After extensive scientific, ethical, and public consultation, in 2016, the UK government gave permission for MRT to prevent the transmission of severe mitochondrial disease caused by specific mtDNA pathogenic variants [Herbert & Turnbull 2017]. The use of MRT in the UK is regulated by the Human Fertilisation and Embryology Authority (HFEA) and is limited to families who are at substantial risk for transmitting severe mtDNA disease in a situation in which PGT is deemed inappropriate or likely to be unsuccessful. A five-year clinical trial is being carried out at the Newcastle Fertility Clinic, and outcomes are yet to be published. In 2022, the Australian government passed the Mitochondrial Donation Law Reform Bill (Maeve's Law) regarding the clinical implementation of MRT. Under the bill, MRT may be introduced in a staged manner, with current legislation allowing for the commencement of a clinical trial of the use of MRT techniques (see National Health and Medical Research Council).

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Resources

GeneReviews staff has selected the following disease-specific and/or umbrella support organizations and/or registries for the benefit of individuals with this disorder and their families. GeneReviews is not responsible for the information provided by other organizations. For information on selection criteria, click here.

MedlinePlus

Leigh syndrome

• United Mitochondrial Disease Foundation

Phone: 888-317-UMDF (8633)

Email: info@umdf.org

umdf.org

• Metabolic Support UK

United Kingdom **Phone:** 0845 241 2173 metabolicsupportuk.org

Mito Foundation

Australia

Phone: 61-1-300-977-180 **Email:** info@mito.org.au

mito.org.au

Mitochondrial Disease Registry and Tissue Bank

Massachusetts General Hospital

Phone: 617-726-5718 **Fax:** 617-724-9620

Email: nslate@partners.org

• Muscular Dystrophy Association (MDA) - USA

Phone: 833-275-6321

Email: ResourceCenter@mdausa.org

mda.org

• People Against Leigh Syndrome (PALS)

peopleagainstleighs.org

• Retina International

Ireland

retina-international.org

• The Lily Foundation

United Kingdom

Email: liz@thelilyfoundation.org.uk

thelilyfoundation.org.uk

• RDCRN Patient Contact Registry: North American Mitochondrial Disease Consortium

Patient Contact Registry

Molecular Genetics

Information in the Molecular Genetics and OMIM tables may differ from that elsewhere in the GeneReview: tables may contain more recent information. —ED.

Table A. Mitochondrial DNA-Associated Leigh Syndrome Spectrum: Genes and Databases

Gene	Chromosome Locus	Protein	ClinVar
MT-ATP6	Mitochondrion	ATP synthase F(0) complex subunit a	MT-ATP6
MT-CO1	Mitochondrion	Cytochrome c oxidase subunit 1	MT-CO1
MT-CO2	Mitochondrion	Cytochrome c oxidase subunit 2	MT-CO2
MT-CO3	Mitochondrion	Cytochrome c oxidase subunit 3	MT-CO3
MT-ND1	Mitochondrion	NADH-ubiquinone oxidoreductase chain 1	MT-ND1
MT-ND2	Mitochondrion	NADH-ubiquinone oxidoreductase chain 2	MT-ND2
MT-ND3	Mitochondrion	NADH-ubiquinone oxidoreductase chain 3	MT-ND3
MT-ND4	Mitochondrion	NADH-ubiquinone oxidoreductase chain 4	MT-ND4
MT-ND5	Mitochondrion	NADH-ubiquinone oxidoreductase chain 5	MT-ND5
MT-ND6	Mitochondrion	NADH-ubiquinone oxidoreductase chain 6	MT-ND6
MT-TI	Mitochondrion	Not applicable	MT-TI
MT-TK	Mitochondrion	Not applicable	MT-TK
MT-TL1	Mitochondrion	Not applicable	MT-TL1
MT-TV	Mitochondrion	Not applicable	MT-TV
MT-TW	Mitochondrion	Not applicable	MT-TW

Data are compiled from the following standard references: gene from HGNC; chromosome locus from OMIM; protein from UniProt. For a description of databases (Locus Specific, HGMD, ClinVar) to which links are provided, click here.

Table B. OMIM Entries for Mitochondrial DNA-Associated Leigh Syndrome Spectrum (View All in OMIM)

161700	NECROTIZING ENCEPHALOMYELOPATHY, SUBACUTE, OF LEIGH, ADULT			
500017	LEIGH SYNDROME, MITOCHONDRIAL; MILS			
516000	COMPLEX I, SUBUNIT ND1; MTND1			
516001	COMPLEX I, SUBUNIT ND2; MTND2			
516002	COMPLEX I, SUBUNIT ND3; MTND3			
516003	COMPLEX I, SUBUNIT ND4; MTND4			
516005	COMPLEX I, SUBUNIT ND5; MTND5			
516006	COMPLEX I, SUBUNIT ND6; MTND6			
516030	COMPLEX IV, CYTOCHROME c OXIDASE SUBUNIT I; MTCO1			
516040	COMPLEX IV, CYTOCHROME c OXIDASE SUBUNIT II; MTCO2			
516050	CYTOCHROME c OXIDASE III; MTCO3			
516060	ATP SYNTHASE 6; MTATP6			
590045	TRANSFER RNA, MITOCHONDRIAL, ISOLEUCINE; MTTI			
590050	TRANSFER RNA, MITOCHONDRIAL, LEUCINE, 1; MTTL1			

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Table B. continued from previous page.

590060	TRANSFER RNA, MITOCHONDRIAL, LYSINE; MTTK
590095	TRANSFER RNA, MITOCHONDRIAL, TRYPTOPHAN; MTTW
590105	TRANSFER RNA, MITOCHONDRIAL, VALINE; MTTV

Molecular Pathogenesis

Human mitochondrial DNA (mtDNA) encodes 37 genes, including 13 genes encoding protein subunits of the mitochondrial respiratory chain and oxidative phosphorylation system, 22 transfer RNA (tRNA) genes, and two ribosomal RNA (rRNA) genes. All mtDNA genes lack introns and are transcribed as large polycistronic transcripts that are processed into monocistronic mRNAs. Protein-coding genes are then translated by the mitochondrial-specific translational machinery. The mitochondrial-specific translational machinery is required because translation of mtDNA-encoded genes is physically separated from the cytosolic translational machinery and because the mtDNA genetic code differs from the universal genetic code in several codons.

For some mtDNA pathogenic variants associated with Leigh syndrome spectrum (LSS), a strong correlation exists between the proportion of abnormal-to-wild type mtDNA and severity of the biochemical phenotype in cultured cells. For some variants, such as *MT-ATP6* pathogenic variants m.8993T>G and m.8993T>C, a strong correlation also exists between the proportion of abnormal-to-wild type mtDNA and clinical severity. However, the mechanism by which some mtDNA pathogenic variants cause a phenotype such as LSS, while others cause myopathy, deafness, or diabetes mellitus, is unknown.

Pathogenic mtDNA variants causing LSS fall into two major classes, namely, those in tRNA genes and those in protein-coding genes.

- Transfer RNA pathogenic variants cause decreased mitochondrial protein synthesis by abnormalities in base modification and aminoacylation of the tRNA.
- Pathogenic variants in protein-coding mtDNA genes typically cause decreased activity of the respiratory chain complex of which that subunit is a part.

Mechanism of disease causation. Loss of function

Table 8. Pathogenic Variants Referenced in This GeneReview by Gene

Reference Sequence	Gene	Mitochondrial DNA Nucleotide Change	Predicted Protein Change	Comment [Reference]
	MT-TL1	m.3243A>G		See Genetic Counseling.
	MT-TK	m.8344A>G		
	MT-ATP6	m.8993T>G	p.Leu156Arg	Most common pathogenic variant in persons w/mtDNA-LSS; see Genotype-Phenotype Correlations.
		m.8993T>C	p.Leu156Pro	See Genotype-Phenotype Correlations.
		m.9176T>C	p.Leu217Pro	
NC_012920.1		m.9185T>C	p.Leu220Pro	One of the most common pathogenic variants in persons w/mtDNA-LSS
	MT-ND3	m.10158T>C	p.Ser34Pro	
		m.10191T>C	p.Ser45Pro	
	MT-ND5	m.13513G>A	p.Asp393Asn	See Genotype-Phenotype Correlations.
	MT-ND6	m.14459G>A	p.Ala72Val	
		m.14487T>C	p.Met63Val	

mtDNA-LSS = mitochondrial DNA-associated Leigh syndrome spectrum

Variants listed in the table have been provided by the authors. *GeneReviews* staff have not independently verified the classification of variants.

GeneReviews follows the standard naming conventions of the Human Genome Variation Society (varnomen.hgvs.org). See Quick Reference for an explanation of nomenclature.

See MITOMAP for details of mitochondrial variants.

Chapter Notes

Author Notes

Prof Rahman, Prof Thorburn, and Dr Ball are actively involved in clinical research regarding individuals with mitochondrial DNA-associated Leigh syndrome spectrum (mtDNA-LSS). Prof Thorburn is a member of the mitoHOPE Program team performing an Australian clinical trial of mitochondrial replacement therapy.

The authors would be happy to communicate with persons who have any questions regarding diagnosis of mtDNA-LSS or other considerations.

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Author History

Megan Ball, MD (2024-present)

Joyeeta Rahman, BSc; UCL Great Ormond Street Institute of Child Health (2017-2024)

Shamima Rahman, PhD, FRCP, FRCPCH (2003-present)

David R Thorburn, PhD, FHGSA, FFSc(RCPA) (2003-present)

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