GENES, CHILDREN AND PEDIATRICIANS

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Herein is reported the clinical case of a nine-year-old female, second child of a non-consanguineous couple, born at term after an uneventful pregnancy and labor. The girl had regular growth and normal development, except for walking delay acquired at 24 months and recent school complaints about attention issues and reading difficulties. Regarding family history, the father had end-stage kidney disease without known cause and had been transplanted, one female cousin deceased at the age of eight years after central nervous system infection, and another female cousin aged 23 years had a neurodegenerative disease with onset after an infection (**Fig. 1**).



The girl was transferred to the Intensive Care Unit with acute respiratory failure after a 24-hour period of cough, sweating, nocturnal agitation, and ill-being evolving to groan, noisy breathing, and perioral cyanosis. She had experienced respiratory arrest at home, requiring mouth-to-mouth resuscitation. At the emergency room, the girl presented polypnea, hypoxemia, tachycardia, and capillary refill time of four seconds with normal blood pressure, without fever. She was agitated, but neurological exam was otherwise normal. Physical exam showed crackles without wheezing. Leukocyte count was 41.000/ul with low C-reactive protein (9.7 mg/L) and chest x-ray displayed right lung pneumonia signs, for which she started empirical antibiotic treatment. She had progressive respiratory distress with respiratory acidosis and hypotension, requiring mechanical ventilation. Cardiac enzymes, creatine-kinase, and echocardiogram were normal. Microbiological exams were sterile and respiratory secretions were positive for Rhinovirus and Enterovirus.

Clinical evolution was favorable and the patient was extubated at hospitalization day three, after which nystagmus at horizontal gaze extremes, ophthalmoparesis with impaired lateral gaze, ataxia, and discrete dysarthria were noted. Careful symptom review identified

episodes of cold extremities, sudoresis and transient skin color alteration, restlessness, irritability, and sleep disturbance in the past month, starting after a sore throat episode.

Brain magnetic resonance revealed symmetrical bilateral T2 hyperintensities on pons and midbrain and basal ganglia involving putamen and caudate (Fig. 2).

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Figure 2 - A and B: Axial T2-weighted and T2 FLAIR basal ganglia images showing circumscribed and hyperintense lesions in the putamen and caudate nuclei. B: Axial T2-weighted brainstem image showing symmetrical involvement of the periaqueductal grey matter. C: Axial T1weighted image showing reduced signal in T2 abnormal areas. D: Axial T1 weighted image with gadolinium showing no significant contrast enhancement.

What is your diagnosis?

DIAGNOSIS

Leigh syndrome, or subacute necrotizing encephalomyelopathy (MIM # 256000)

Leigh syndrome, or subacute necrotizing encephalomyelopathy (MIM # 256000), is a neurodegenerative disease with characteristic neuroradiological features, including symmetrical T2 basal ganglia hyperintensities with or without brainstem involvement. It translates an energy metabolism disease with more than 75 known causing genes, located either on nuclear or mitochondrial genome.¹ It is the most common presentation of mitochondrial disease in childhood, either related with nuclear or mitochondrial genome. Diagnostic criteria include: i) clinical brainstem and/or basal ganglia dysfunction, ii) intellectual and motor developmental delay, and iii) abnormal energy metabolism (severe defect in respiratory chain function or pyruvate dehydrogenase complex activity, molecular diagnosis in a mitochondrial energy-production gene, or elevated serum or CSF lactate).²

Leigh syndrome most often presents after a catabolic event, such as an infection, surgery, or fasting.¹ Neurological involvement – as ataxia (basal ganglia) or acute respiratory insufficiency (brainstem involvement) – following such events should raise clinical suspicion of this diagnosis.¹⁻³

In this patient, proband mitochondrial genetic analysis showed mitochondrial ATP synthase subunit 6 gene (MT-ATP6) m.8993T>G mutation in homoplasmy (>95% of mutated mtDNA copies). MT-ATP6 gene codifies for a respiratory chain complex V subunit.³ Further analyses confirmed the same diagnosis in the patient's cousin.

Acute neurological deterioration with movement disorder or respiratory failure in a patient with a catabolic event (infection with or

without fever, surgery, fasting, trauma, etc.) should raise suspicion of Leigh (-like) syndrome.⁴ In this case, pedigree analysis showing positive family history for individuals interconnected by females raised the possibility of a maternally inherited condition (mitochondrial DNA-related disease). Together with Leigh syndrome diagnosis, a primary mitochondrial disorder related with mitochondrial DNA (mtDNA) was among the first diagnostic options.

Leigh syndrome follow-up includes neurological, ophthalmological (risk of retinopathy), and renal (risk of tubulopathy) evaluation, although other mitochondrial disorder manifestations can occur.⁴ After six years of follow-up, this patient showed emotional lability, hetero-aggressiveness periods, agitated sleep, and discrete ataxia, with no other brainstem dysfunction or development regression episodes besides developmental delay.

Genetic counseling for mitochondrial-DNA related disorders is extremely challenging. First, phenotype results from a combination of factors, including pathogenic variant severity, percentage of mutated mitochondria (mtDNA heteroplasmy), and tissue distribution, besides potential variants in other relevant genes that can modify phenotype (disease-modifying genes). Consequently, there is a wide phenotypic range even within the same family, ranging from mild to severe symptoms with varying age of onset. This makes virtually impossible to interpret asymptomatic at-risk family member test results in order to have phenotypic prediction. Moreover, absence of mtDNA pathogenic variants in one tissue (e.g. blood) does not guarantee its absence from other tissues (e.g. oocytes).⁵ Second, risk of recurrence in sibs depends on the mothers' genetic status, as well as tissue distribution and abnormal mtDNA proportion. Proband mother usually has much lower abnormal mtDNA proportion than the proband and may be asymptomatic.⁵

Genetic counselling regarding reproductive options is also challenging. Prenatal testing (directly performed on biopsy cells, not cultured) can have conflicting results with an intermediate abnormal mtDNA proportion, with nearly impossible outcome prediction. Oocyte donation accompanied by in vitro fertilization using partner's sperm can be an option; however, availability of donor oocytes is limited and personal or cultural issues may arise regarding the use of donor gametes. Preimplantation genetic diagnosis is currently considered the preferred reproductive option and can be used to transfer embryos with lower mutant mtDNA load, with low recurrence risk. Mitochondrial donation, in which nuclear material is transferred as pronucleus or maternal spindle from an unfertilized oocyte or single-cell embryo into an enucleated healthy donor cell (with unaffected mitochondria) could potentially block transmission of mutated mtDNA into the developing embryo.⁵ However, unlike countries such as the United Kingdom, this technique it is not currently available in Portugal.

Overall, genetic counseling for mitochondrial genetic related disease presents considerable challenges. It requires multidisciplinary involvement with an up-to-date experienced team, to ensure that prospective parents are counseled regarding all potential outcomes and limitations of prenatal diagnosis or assisted reproduction.

Keywords: brainstem dysfunction; Leigh syndrome; m.8993T>G; mitochondrial disorder; mtDNA

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