

L'essentielle Maladies Mitochondriales



2^{ème} trimestre 2025



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Ataxie de Friedreich - *Friedreich ataxia*

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Encéphalopathie myo-neuro-gastrointestinale - *Mitochondrial neurogastrointestinal encephalomyopathy (MNGIE)*

Al-Sahlawi, Z., Redha, N. A. et Hasan, H. (2025). **Aseptic Pleocytosis Can Only be Classified as a Phenotypic Manifestation of MNGIE After Exclusion of all Differential Causes.** *J Cent Nerv Syst Dis* 17, 11795735251330595, doi:[10.1177/11795735251330595](https://doi.org/10.1177/11795735251330595).

Maladie de Charcot-Marie-Tooth d'origine mitochondriale – *Mitochondrial Charcot-Marie-Tooth*

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Maladies liées au gène ACO2 - ACO2-related disorders

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Syndrome d'Alpers-Huttenlocher - *Alpers-Huttenlocher syndrome*

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Syndrome de Kearns-Sayre – *Kearns-Sayre syndrome*

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Syndrome de Leigh - Leigh syndrome

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