

L'essentielle Maladies Mitochondriales
MMITO-2024-3 du 1^{er} juillet au 30 septembre 2024

L'essentielle Maladies Mitochondriales



3^{ème} trimestre 2024



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Cette veille, effectuée à partir de requêtes sur PubMed® sans prétendre à l'exhaustivité, présente une sélection de références d'articles médico-scientifiques concernant le domaine des maladies mitochondriales.



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Ataxie de Friedreich - *Friedreich ataxia*

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Atrophie optique autosomique dominante – ADOA

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Encéphalopathie myo-neuro-gastrointestinale - MNGIE

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

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Maladies liées au gène ACO2 - ACO2-related disorders

Pas de résultat ce trimestre

Maladie liée au gène GFM1 – GFM1-related disorder

Pas de résultat ce trimestre

Maladies liées au gène POLG (inclus SANDO, SCAE) - POLG-related disorders

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