

# Choroba Lebera 'plus'

Kod Orpha: 99718 Kod OMIM: 500001

## Opis choroby \*

### Definicja

A rare inherited mitochondrial disease characterized by the clinical features of Leber hereditary optic neuropathy in combination with other systemic or neurological abnormalities. These abnormalities include: postural tremor, motor disorder, multiple sclerosis-like syndrome, spinal cord disease, skeletal changes, Parkinsonism with dystonia, anarthria, motor and sensory peripheral neuropathy, spasticity, mild encephalopathy, and cardiac arrhythmias.

### Dane

#### Klasyfikacja

Choroba

#### Synonimy

LHON plus disease

LHON plus disease

#### Kod ORPHA

99718

#### Kod OMIM

500001

#### Kod ICD10

H47.2

#### Kod ICD11

8C73.Y

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\*Źródło

orphanet

## Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.