

# **Leukoencefalopatia - chondrodysplazja przynasadowa**

## **Kod Orpha: 83629 Kod OMIM: 300232**

### **Opis choroby \***

#### **Definicja**

A rare genetic neurological disorder characterized by the association of hypomyelinating leukodystrophy with spondylometaphyseal dysplasia. Patients present in infancy with absent or delayed ability to walk independently, slowly progressive motor deterioration, spasticity, ataxia, proximal weakness, and joint contractures. Additional manifestations include mild cognitive impairment, short stature, scoliosis, enlarged and deformed joints, dysarthria, nystagmus, visual defects, and mildly dysmorphic features, among others. Mode of inheritance is X-linked recessive.

#### **Dane**

<b>Klasyfikacja</b>	<b>Synonimy</b>
Choroba	H-SMD
	Hypomyelination-spondyloepimetaphyseal dysplasia syndrome
	Leukoencephalopathy-SEMD syndrome
	Leukoencephalopathy-metaphyseal chondrodysplasia syndrome
	H-SMD
	Hypomyelination-spondyloepimetaphyseal dysplasia syndrome
	Leukoencephalopathy-SEMD syndrome
	Leukoencephalopathy-metaphyseal chondrodysplasia syndrome

**Kod ORPHA**  
83629

**Kod OMIM**  
300232

**Kod ICD10**  
G37.8

**Kod ICD11**

\*[Źródło](#)

## Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

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