

Niski wzrost typu Brussels

Kod Orpha: 2867 Kod OMIM: 601350

Opis choroby *

Definicja

A rare primary bone dysplasia characterized by severe intrauterine and postnatal growth retardation and short stature in association with craniofacial dysmorphism (such as large forehead, triangular face, low-set ears, and micro-retrognathism) and osteochondrodysplastic lesions. Radiographic findings include epiphyseal maturation delay, abnormal metaphyses, a narrow thorax, small pelvis, and short and broad metacarpal bones and phalanges. There have been no further descriptions in the literature since 1996.

Dane

Klasyfikacja

Zespół wad
wrodzonych

Synonimy

Mievis-Verellen-Dumoulin syndrome
Zespół Mievisa, Verellena i Dumoulina

Kod ORPHA

2867

Kod OMIM

601350

Kod ICD10

Q87.1

Kod ICD11

LD24.2Y

*Źródło

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.