## Opis choroby \*

## Definicja

Weismann-Netter syndrome is a rare, genetic, primary, bent bone dysplasia characterized by anterior diaphyseal bowing of the tibia and fibula, broadening of the fibula, posterior cortical thickening of both bones and short stature. Additional skeletal abnormalities include scoliosis with marked lumbar lordosis, horizontal sacrum and square iliac wings and/or, less frequently, vertebral malformations, abnormal shape of the clavicles and ribs, calvarial hyperostosis and delayed eruption of permanent teeth. Delayed ambulation is also frequently associated.

Dane

Klasyfikacja Synonimy

Zespół wad wrodzonych Anterior bowing of legs with dwarfism

**WNS** 

Weismann-Netter-Stuhl syndrome

 Kod ORPHA
 Kod OMIM
 Kod ICD10

 3344
 112350
 077.8

Kod ICD11 LD24.C

## \*Źródło

orphanet