

Zespół Stolla, Alembika i Fincka

Kod Orpha: 3200 Kod OMIM: 601701

Opis choroby *

Definicja

A rare, genetic developmental defect during embryogenesis syndrome characterized by camptodactyly, joint contractures with amyotrophy, and ectodermal anomalies (oligodontia, enamel abnormalities, longitudinally broken nails, hypohidrotic skin with tendency to excessive bruising and scarring after injuries and scratching), as well as growth retardation, kyphoscoliosis, mild facial dysmorphism, and microcephaly. There have been no further descriptions in the literature since 1992.

Dane

Klasyfikacja	Synonimy
Zespół wad wrodzonych	Stoll-Alembik-Finck syndrome Artrogrypoza - dysplazja ektodermalna - inne wady
Kod ORPHA	Kod OMIM
3200	601701
Kod ICD10	Kod ICD10
-	Q68.8
Kod ICD11	
-	

*Źródło

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.