

Zespół wieloukładowy związany z LAMA5

Kod Orpha: 521450 Kod OMIM:

Opis choroby *

Definicja

A rare genetic systemic or rheumatologic disease characterized by infantile onset of skin anomalies (such as delayed wound healing with atrophic scars and mild alopecia with dry and brittle hair), retinal rod degeneration with night blindness, degenerative myopathy with muscle weakness, myalgia, and cramps, osteoarthritis, joint laxity, prolapse of internal organs, floating kidney syndrome, malabsorption syndrome, and hypothyroidism. The phenotype has been reported to be more severe in women than in men.

Dane

Klasyfikacja

Choroba

Kod ORPHA
521450

Kod OMIM
-

Kod ICD10
M79.8

Kod ICD11

[*Źródło](#)

[orphanet](#)

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.