

Rybia łuska z keratynopatią

Kod Orpha: 281103 Kod OMIM:

Opis choroby *

Definicja

A group of rare inherited non-syndromic ichthyoses characterized by mutations in keratin genes. Mutations in *KRT1* and *KRT10* cause most cases of epidermolytic ichthyosis (EI), as well as congenital reticular ichthyosiform erythroderma (CRIE). EI manifests at birth with generalized blistering, which later transforms into hyperkeratosis. Severe palmoplantar involvement is suggestive of the presence of a *KRT1* mutation. CRIE patients present at birth with erythroderma and scaling, often with a collodion membrane, and gradually develop confetti-like clear areas of normal skin. *KRT2* mutations are associated with superficial epidermolytic ichthyosis (SEI), which is clinically similar to EI, but generally milder and more localized.

Dane

Klasyfikacja

Grupa fenomenów

Synonimy

KPI

KPI

Kod ORPHA

281103

Kod OMIM

-

Kod ICD10

-

Kod ICD11

EC20.03

[*Źródło](#)

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.

Orphanet - internetowa baza danych dotyczących rzadkich chorób i sierochych leków. ©INSERM 1999 -
Dostępna na stronie www.orphanet.pl