

Opis choroby *

Definicja

A rare, genetic, epidermal disorder characterized by intermittent (remitting and recurring), annular, polycyclic, target-like (or 'en cocardes') plaques with concentric rings of scaling erythema occurring on the extremities, flexural areas, and trunk. Concurrent erythrokeratoderma variabilis-like scaly plaques are commonly found in other parts of the body.

Dane

Klasyfikacja

Choroba

Synonimy

Degos genodermatosis "en cocardes"

Degos genodermatosis "en cocardes"

Kod ORPHA

315

Kod OMIM

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Kod ICD10

Q82.8

Kod ICD11

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*Źródło

orphanet