

## Opis choroby \*

### Definicja

A rare, genetic, hyperpigmentation of the skin disease characterized by adulthood-onset of reticular, reddish-brown to dark-brown, macular and/or comedone-like, hyperkeratotic papules with hypopigmented macules, predominantly affecting flexural areas and, on occasion, progressing to involve trunk and acral regions. Histologically, epidermal acanthosis, thin, branch-like, rete ridges, and a tendency for acantholysis and pigmentary incontinence is observed.

### Dane

| Klasyfikacja | Synonimy   |
|--------------|--|
| Choroba      | Reticular pigment anomaly of flexures<br>Siateczkowata barwnikowa anomalia powierzchni zgięciowych |

| Kod ORPHA | Kod OMIM | Kod ICD10 |
|-----------|----------|-----------|
| 79145     | 615696   | L81.8     |

| Kod ICD11 |
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| EC23.0    |

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\*Źródło

orphanet