

Opis choroby *

Definicja

Familial progressive hyperpigmentation is a rare, genetic, skin pigmentation anomaly disorder characterized by irregular patches of hyperpigmented skin which present at birth or in early infancy and increase in size, number and confluence with age. Affected areas of the body include the face, neck, trunk and limbs, as well as the palms, soles, oral mucosa and conjunctiva. No hypopigmentation macules are observed and no systemic diseases are associated.

Dane

Klasyfikacja

Choroba

Synonimy

Melanosis diffusa congenita
Dziedziczna melanoza uogólniona
Universal melanosis
Wrodzona melanoza rozsiana
Melanosis universalis hereditaria
Universal melanosis

Kod ORPHA

79146

Kod OMIM

614233

Kod ICD10

L81.4

Kod ICD11

EC23.0

*Źródło

orphanet