Opis choroby *

Definicja

A rare hereditary poikiloderma characterized by infantile onset of vesicopustule formation on hands and feet and widespread eczematoid dermatitis (both spontaneously resolving during childhood), as well as gradually developing diffuse poikiloderma with striate and reticulate atrophy (excluding the face, scalp, and ears), and development of keratotic papules on hands, feet, elbows, and knees, beginning in early childhood. There have been no further descriptions in the literature since 1981.

Dane

Klasyfikacja Synonimy

Choroba Weary syndrome

Wrodzona poikilodermia z pęcherzami, typu

Q82.8

Weary

Kod ORPHA Kod OMIM Kod ICD10

Kod ICD11 LD2B

*Źródło

orphanet