

# Dermatoosteoliza typu Kirghiziana

## Kod Orpha: 1657 Kod OMIM: 221810

### Opis choroby \*

#### Definicja

A rare genetic disease characterized by infantile onset of recurrent skin ulcerations, arthralgias, fever, peri-articular fistulous osteolysis, oligodontia, nail dystrophy, and keratitis. The disease takes a self-limiting course in childhood but results in severe cicatrization, chronic arthroses, pseudoacromegalic appearance of hands and feet, secondary scoliosis, and visual impairment. There have been no further descriptions in the literature since 1983.

#### Dane

#### Klasyfikacja

Zespół wad  
wrodzonych

**Kod ORPHA**  
1657

**Kod OMIM**  
221810

**Kod ICD10**  
Q82.8

**Kod ICD11**  
LD27.0Y

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\*Źródło

orphanet

### Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.