## Opis choroby \*

## Definicja

A very rare acrofacialdyosotosis characterized by short stature, acrocephaly, ocular hypertelorism, ptosis of eyelids, ocular proptosis, downslanting palpebral fissures, high nasal bridge, anteverted nostrils, short philtrum, cleft palate, micrognathia, abnormal external ears, preauricular pits, mixed hearing loss, bulbous digits, metatarsus varus, pectus excavatum and various radiological abnormalities. Features of this syndrome were reported to overlap with otopalatodigital syndrome types 1 and 2. There have been no further descriptions in the literature since 1988.

Dane

Klasyfikacja Synonimy

Zespół wad wrodzonych Kaplan-Plauchu-Fitch syndrome

Zespół Kaplana, plauchu i Fitcha

 Kod ORPHA
 Kod OMIM
 Kod ICD10

 949
 201050
 Q87.0

Kod ICD11

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## \*Źródło

orphanet