## Opis choroby \*

Definicja

A rare syndrome characterized by gingival fibromatosis associated with progressive sensorineural hearing loss. It has been described in two families (with at least 16 affected members spanning five generations in one of the families, and five affected members spanning three generations in the other family). It is transmitted as an autosomal dominant trait.

Dane

Klasyfikacja Synonimy

Zespół wad wrodzonych Gingival fibromatosis-progressive hearing loss

syndrome Zespół Jonesa Jones syndrome

Kod ORPHA

HA Kod OMIM

**Kod ICD10** 

2027

135550

H90.3

Kod ICD11 LD2H.Y

## \*Źródło

orphanet