

# Zespół FG typu 1

Kod Orpha: 93932 Kod OMIM: 305450

## Opis choroby \*

### Definicja

A rare X-linked syndromic intellectual disability characterized by developmental delay and intellectual disability, early hypotonia, constipation, feeding problems, imperforate anus, characteristic behavior (affable, eager to please), and dysmorphic craniofacial features (such as relative macrocephaly, prominent forehead with frontal hair upsweep, hypertelorism, downslanting palpebral fissures, and open mouth). Additional manifestations are partial agenesis of the corpus callosum, sensorineural hearing loss, joint laxity, cardiac anomalies, and abnormalities of the fingers and toes, among others.

### Dane

#### Klasyfikacja

Choroba

#### Synonimy

Opitz-Kaveggia syndrome

Zespół Opitza i Kaveggia

#### Kod ORPHA

93932

#### Kod OMIM

305450

#### Kod ICD10

Q87.8

#### Kod ICD11

LD2F.1Y

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#### [\\*Źródło](#)

orphanet

## Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.