

## **Opis choroby \***

### Definicja

A rare congenital anomalies syndrome characterized by a variable spectrum of ulnar defects, mammary and apocrine gland hypoplasia and genital anomalies. The most frequent signs include fifth finger and dental anomalies, delayed puberty and mammary hypoplasia. Short stature and obesity are common.

### Dane

#### **Klasyfikacja**

Zespół wad wrodzonych Pallister ulnar-mammary syndrome

UMS

Zespół łokciowo-sutkowy Pallistera

Zespół Schinzela

Schinzel syndrome

UMS

#### **Kod ORPHA**

3138

#### **Kod OMIM**

181450

#### **Kod ICD10**

Q71.8

#### **Kod ICD11**

LD2F.1Y

---

#### \*Źródło

orphanet