## Opis choroby \*

## Definicja

A rare genetic, neurodevelopmental syndrome characterized by hypothalamic-pituitary dysfunction with severe hypotonia and feeding deficits during the neonatal period followed by an excessive weight gain period with hyperphagia with a risk of severe obesity during childhood and adulthood, learning difficulties, deficits of social skills and behavioral problems or severe psychiatric problems.

Dane

Klasyfikacja Synonimy

Choroba Prader-Labhart-Willi syndrome

Zespół Pradera, Labharta i Williego

Zespół Williego i Pradera

**Kod ORPHA** 

739

**Kod OMIM** 

**Kod ICD10** 

615547

Q87.1

Kod ICD11 LD90.3

## \*Źródło

orphanet