

Wielohormonalna niedoczynność przysadki

Kod Orpha: 90695 Kod OMIM: 312000

Opis choroby *

Definicja

A rare genetic pituitary disease characterized by variable deficiency of all hormones produced in the anterior lobe of the pituitary gland. Clinical manifestations include hypothyroidism, hypogonadism, growth retardation and short stature, and secondary adrenal insufficiency. Age of onset is variable. Signs and symptoms usually develop gradually, and loss of the different hormones is often sequential.

Dane

Klasyfikacja	Synonimy	
Choroba	Genetic panhypopituitarism Genetic panhypopituitarism	
Kod ORPHA	Kod OMIM	Kod ICD10
90695	312000	E23.0
Kod ICD11		
5A61.0		

*[Źródło](#)

orphanet

Rozszerzony opis choroby

Brak opisu rozszerzonego dla tej choroby. Opracowanie w toku.