Opis choroby *

Definicja

A subtype of autosomal recessive limb-girdle muscular dystrophy characterized by a variable age of onset of progressive weakness and wasting of the proximal skeletal muscles of the shoulder and pelvic girdles, frequently associated with progressive respiratory muscle impairment and cardiomyopathy. Calf hypertrophy, muscle cramps and elevated serum creatine kinase levels are also observed. Neuropsychomotor development is usually normal.

Dane

Klasyfikacja Choroba	Synonimy Autosomal recessive lin dystrophy type 2F Delta-sarkoglikanopatia Dystrofia obręczowo-ko niedoboru delta-sarkog LGMD2F Delta-sarcoglycan-relate Delta-sarcoglycanopath LGMD due to delta-sarco LGMD type 2F LGMD2F Limb-girdle muscular de sarcoglycan deficiency	a pńczynowa z powodu glikanu ed LGMD R6 ny coglycan deficiency ystrophy due to delta-
Kod ORPHA	Kod OMIM	Kod ICD10

219

Kod OMIM 601287 Kod ICD10 G71.0

Kod ICD11 8C70.41

<u>*Źródło</u>

orphanet