

Opis choroby *

Definicja

A rare non-syndromic syndactyly characterized by mesoaxial reduction of fingers, complete syndactyly of the 3rd and 4th fingers with synostoses of the corresponding metacarpals and associated single phalanges, malformed thumbs, and hypoplasia and clinodactyly of the 5th finger. Preaxial webbing of toes with terminal phalangeal hypoplasia of all toes has been reported in association.

Dane

Klasyfikacja	Synonimy
Wada morfologiczna	MSSD
	MSSD
	Syndaktylia typu 9
	Syndaktylia, typ Malika i Percina
	Syndactyly type 9
	Syndactyly, Malik-Percin type

Kod ORPHA
157801

Kod OMIM
609432

Kod ICD10
Q70.2

Kod ICD11
LB79.Y

*Źródło

orphanet