

Opis choroby *

Definicja

Sanjad-Sakati syndrome (SSS), also known as hypoparathyroidism - intellectual disability-dysmorphism, is a rare multiple congenital anomaly syndrome, mainly occurring in the Middle East and the Arabian Gulf countries, characterized by intrauterine growth restriction at birth, microcephaly, congenital hypoparathyroidism (that can cause hypocalcemic tetany or seizures in infancy), severe growth retardation, typical facial features (long narrow face, deep-set eyes, beaked nose, floppy and large ears, long philtrum, thin lips and micrognathia), and mild to moderate intellectual deficiency. Ocular findings (i.e. nanophthalmos, retinal vascular tortuosity and corneal opacification/clouding) and superior mesenteric artery syndrome have also been reported. Although SSS shares the same locus with the autosomal recessive form of Kenny-Caffey syndrome (see this term), the latter differs from SSS by its normal intelligence and skeletal features.

Dane

Klasyfikacja	Synonimy
Zespół wad wrodzonych HRD syndrome	
	SSS
	Zespół HRD
	Zespół niedoczynności przytarczyc, niepełnosprawności intelektualnej i dysmorfii
	Zespół niedoczynności przytarczyc, niskiego wzrostu, niepełnosprawności intelektualnej i napadów padaczkowych
	Zespół Richardsona i Kirka
	Hypoparathyroidism-intellectual disability-dysmorphism syndrome
	Hypoparathyroidism-short stature-intellectual disability-seizures syndrome
	Richardson-Kirk syndrome
	SSS

Kod ORPHA
2323

Kod OMIM
241410

Kod ICD10
Q87.1

Kod ICD11
LD24.D

*[Źródło](#)

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