

## Opis choroby \*

### Definicja

A rare genetic disease associated with mild to severe intellectual deficit that may be associated with behavioral disorders and characteristic physical features including a high forehead, prominent and large ears, hyperextensible finger joints, flat feet with pronation and, in adolescent and adult males, macroorchidism.

### Dane

<b>Klasyfikacja</b>	<b>Synonimy</b>
Zespół wad wrodzonych	FRAXA syndrome
	FXS
	Zespół FraX
	Zespół FRAXA
	Zespół Martina i Bell
	FXS
	FraX syndrome
	Martin-Bell syndrome

**Kod ORPHA**  
908

**Kod OMIM**  
311360

**Kod ICD10**  
Q99.2

**Kod ICD11**  
LD55

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### \*Źródło

orphanet