
DISEASE:
7q31 microdeletion syndrome

NAME:	7q31 microdeletion syndrome
DESCRIPTION:	7q31 microdeletion syndrome is a rare chromosomal anomaly characterized by speech and language disorder, predominantly presenting as an apraxia of speech, sometimes associated with oral motor dyspraxia, dysarthria, receptive and expressive language disorder, and hearing loss. Individuals with larger deletions in this region have also been reported to display intellectual disability and autism.
ORPHACODE:	251061
SYNONYMS:	Del(7)(q31) Monosomy 7q31
XREF(S):	Orphanet
ANALYTE(S):	FOXP2
CREATED:	13 May 2019 - 01:02
CHANGED:	22 Jun 2023 - 16:14

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