

**DISEASE:**  
**15q11.2 microdeletion syndrome**

<b>NAME:</b>	15q11.2 microdeletion syndrome
<b>DESCRIPTION:</b>	15q11.2 microdeletion syndrome is a rare partial autosomal monosomy with a variable phenotypic expression and reduced penetrance associated with an increased susceptibility to neuropsychiatric or neurodevelopmental disorders including delayed psychomotor development, speech delay, autism spectrum disorder, attention deficit-hyperactivity disorder, obsessive-compulsive disorder, epilepsy or seizures. It may also include mild non-specific dysmorphic features (such as dysplastic ears, broad forehead, hypertelorism), cleft palate, neurological and neuroimaging abnormalities (such as ataxia and muscular hypotonia).
<b>ORPHACODE:</b>	261183
<b>SYNONYMS:</b>	15q11.2 BP1-BP2 microdeletion syndrome Del(15)(q11.2) Monosomy 15q11.2
<b>XREF(S):</b>	<a href="#">Orphanet</a> <a href="#">OMIM</a> <a href="#">ICD-10</a>
<b>ANALYTE(S):</b>	<a href="#">NIPA1</a> <a href="#">NIPA2</a> <a href="#">TUBG1</a>
<b>CREATED:</b>	17 May 2022 - 02:46

<b>CHANGED:</b>	22 Jun 2023 - 16:14
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- [NIPA magnesium transporter 2](#)
- [tubulin gamma 1](#)

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