

**DISEASE:**  
**16q24.3 microdeletion syndrome**

<b>NAME:</b>	16q24.3 microdeletion syndrome
<b>DESCRIPTION:</b>	16q24.3 microdeletion syndrome is a recently described syndrome associated with variable developmental delay, facial dysmorphism, seizures and autistic spectrum disorder.
<b>ORPHACODE:</b>	261250
<b>SYNOMYS:</b>	Del(16)(q24.3) Monosomy 16q24.3
<b>XREF(S):</b>	<a href="#">Orphanet</a> <a href="#">ICD-10</a>
<b>ANALYTE(S):</b>	<a href="#">ANKRD11</a>
<b>CREATED:</b>	13 May 2019 - 01:02
<b>CHANGED:</b>	22 Jun 2023 - 16:14

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### Related Genetic Tests

- [Short Stature \(gene panel\)](#)

### Related Laboratories

- [Centre de Génétique-Institut de Pathologie et de Génétique \(IPG\)](#)

### Related Analytes

- [ankyrin repeat domain containing 11](#)

### Related Gene Panels

- [Short Stature \(46 genes\) - IPG](#)

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