

GENETIC TEST: **Angelman / Prader Willi Syndrome**

FULL NAME:	Angelman / Prader Willi Syndrome
TEST TYPE:	Clinical
TEST SPECIALTY:	Molecular Genetics
TEST PURPOSE:	Post-natal Diagnosis, Prenatal diagnosis
SPECIMEN:	Peripheral (whole) blood on EDTA, Amniotic fluid, Chorionic villi
METHOD CATEGORY:	Methylation analysis Deletion/duplication analysis
METHOD TECHNIQUE:	MLPA based techniques
RIZIV CODE:	565456-565460
ACCREDITATION (ISO 15189):	2022-10-07 / 2027-10-06

EQA:	<ul style="list-style-type: none">• Prader-Willi and Angelman Syndromes,• Prader-Willi and Angelman Syndromes ,• Prader-Willi and Angelman Syndromes
TURNAROUND TIME (MAXIMUM):	15 days
CREATED:	24 Jul 2019 - 11:08
CHANGED:	23 Jan 2023 - 15:54

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RELATED CONTENT

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- [Angelman syndrome due to imprinting defect in 15q11-q13](#)
- [Angelman syndrome due to maternal 15q11q13 deletion](#)
- [Angelman syndrome due to paternal uniparental disomy of chromosome 15](#)
- [Prader-Willi syndrome due to imprinting mutation](#)
- [Prader-Willi syndrome due to maternal uniparental disomy of chromosome 15](#)
- [Prader-Willi syndrome due to paternal deletion of 15q11q13 type 2](#)

Related Laboratories

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

Related Analytes

- [chromosome 15 - 15q11-q13](#)
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