Angelman Syndrome and Prader-Willi Syndrome by Methylation-Specific MLPA

Angelman syndrome (AS) and Prader-Willi syndrome (PWS) are complex neurodevelopmental disorders characterized by developmental delay and cognitive disability, as well as symptoms unique to each disorder (e.g., unique happy demeanor in AS, excessive eating in PWS). Both conditions are linked to loss of function of genes in the 15q11.2-q13 region.

Disease Overview

Prevalence
- **AS:** 1 in 12,000-24,000
- **PWS:** 1 in 10,000-30,000

Age of Onset
- **AS:** 6-12 months of age
- **PWS:** Neonatal

Genetics

Genes
15q11.2-q13 region

Etiologies
- Deletion of 15q11.2-q13 (AS: maternal; PWS: paternal)
- Uniparental disomy (UPD) for chromosome 15 (AS: paternal; PWS: maternal)
- Imprinting center defect
- Unbalanced chromosome translocation
- UBE3A gene mutation (AS only)
- Unidentified (AS only)

For more information about the underlying mechanisms of AS and PWS, refer to the ARUP Consult Angelman Syndrome and Prader-Willi Syndrome topic.

Prenatal Screening
- Prenatal testing is recommended for subsequent pregnancies of couples who have a previous child with AS or PWS.
- Parental testing does not exclude somatic and/or germline mosaicism.
- Testing of chorionic villus samples is not recommended as methylation may be incomplete in early embryonic development.

Test Interpretation

Clinical Sensitivity
- >99% for PWS
- 80% for AS

Analytic Sensitivity
- 99% for PWS and AS
### Results

#### Positive Result

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<th>Interpretation</th>
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| Maternally contributed AS/PWS critical region only, with normal copy number | • Confirms a diagnosis of PWS  
• Order DNA polymorphism analysis to distinguish between UPD and imprinting defect |
| Maternally contributed AS/PWS critical region only, with abnormal copy number consistent with deletion | • Confirms a diagnosis of PWS  
• Consider chromosome analysis for proband to exclude rare rearrangement and to determine the need for paternal/maternal karyotyping[^a] |
| Paternally contributed AS/PWS critical region only, with normal copy number | • Confirms a diagnosis of AS  
• Order DNA polymorphism analysis to distinguish between UPD and imprinting defect |
| Paternally contributed AS/PWS critical region only, with abnormal copy number consistent with deletion | • Confirms a diagnosis of AS  
• Consider both chromosome analysis and fluorescence in situ hybridization (FISH) in mother to exclude rare rearrangement[^a] |
| Paternally and maternally contributed AS/PWS critical regions detected, with abnormal copy number consistent with duplication | This assay is not validated to detect increased copy number of 15q11.2-q13 or determine parent of origin for duplications |

[^a]: Alters recurrence risk. Refer to the ARUP Consult Angelman Syndrome and Prader-Willi Syndrome topic for more information.

#### Negative Result

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| Normal methylation pattern of both maternally and paternally contributed AS/PWS critical regions with normal copy number | • Greatly reduces the probability of a PWS diagnosis; <1% of individuals with PWS have normal methylation patterns  
• Reduces, but does not exclude, the probability of an AS diagnosis; approximately 20% of individuals with AS have normal methylation patterns |

### Limitations

- Disease mechanisms causing AS that do not alter methylation patterns will not be detected.
- The specific molecular mechanism responsible for abnormal methylation results cannot be determined via this test alone.
- Diagnostic errors can occur due to rare sequence variations.
- This assay is not validated to detect increased copy number of 15q11.2-q13 or determine parent of origin for duplications.
- This assay cannot distinguish between UPD and imprinting defects causative of PWS and AS.
- AS and PWS mosaicism will not be assessed by this assay.
- Interpretation of this test result may be impacted if the proband has had an allogeneic stem cell transplantation.
- Methylation patterns may not be fully established in early gestation; thus, diagnostic testing on chorionic villus samples is not recommended.

### References


### Additional Resources


Related Information

**Angelman Syndrome and Prader-Willi Syndrome**

Cytogenomic SNP Microarray

Laboratory Testing for Developmental Delay, Intellectual Disability, and Autism Spectrum Disorder

Testing for Genetic Syndromes Related to Developmental Delay (DD), Intellectual Disability (ID), and Autism Spectrum Disorder (ASD)