# Beckwith-Wiedemann and Russell-Silver Syndromes

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Beckwith-Weidemann syndrome (BWS) is a congenital overgrowth condition associated with neonatal hypoglycemia, macroglossia, macrosomia, hemihypertrophy and increased risk for embryonal tumors. Russell-Silver syndrome (RSS) is a congenital condition characterized by stunted grow, limb length asymmetry, and developmental delay. Testing can confirm a suspected clinical diagnosis of BWS or RSS.

## Disease Overview

#### Incidence

BWS: ~1/10,000-13,700 newborns

RSS: ~1/100,000 newborns

## Featured ARUP Testing

Beckwith-Wiedemann Syndrome (BWS) and Russell-Silver Syndrome (RSS) by Methylation-Specific MLPA 3001635

**Method**: Multiplex Ligation-Dependent Probe Amplification (MLPA)

Confirm diagnosis of BWS or RSS in individuals with a suspected clinical diagnosis

#### **Symptoms**

BWS (Major Findings)	RSS
Macrosomia	Pre- and postnatal growth deficiency
Visceromegaly	Proportionate short stature
Hemihyperplasia	Limb length asymmetry
Embryonal tumors in childhood (eg, Wilms tumor, hepatoblastoma, neuroblastoma, rhabdomyosarcoma)	Developmental delay and/or learning disabilities
Macroglossia	Triangular facies, broad forehead, narrow chin
Omphalocele	
Renal abnormalities	
Ear creases or pits	

## Genetics

## Etiology

#### Causes of BWS

- 50% have loss of maternal methylation on chromosome 11p15 imprinting center (IC)2
- 20% have paternal uniparental disomy (UPD) for chromosome 11p15
- 5% have gain of methylation in maternal IC1
- Pathogenic sequence variants in CDKN1C
  - 5-10% of nonfamilial cases
  - ~40% of familial cases
  - <1% cytogenetic abnormalities involving 11p15</li>

#### Causes of RSS

- 35-50% have hypomethylation of paternal IC1
- 10% have maternal UPD of chromosome 7
- ~40% have an unknown genetic mechanism

#### Inheritance

- Sporadic in 85% of BWS cases and 60% of RSS cases
- Autosomal dominant in 15% of BWS cases due to parent-of-origin transmission

#### Penetrance

- · Complete for RSS
- Incomplete for BWS due to methylation (eg, individuals with a paternally inherited CDKN1C pathogenic variant will not show features of BWS)

## **Test Interpretation**

Clinical sensitivity/specificity: 75% for BWS; 35-50% for RSS

Analytical sensitivity/specificity: 99%

### Results

Result	BWS	RSS
Positive	IC2 hypomethylation AND normal IC1 methylation IC1 hypermethylation AND hypomethylation of IC2 IC1 hypermethylation AND normal methylation of IC2	IC1 hypomethylation
Negative	Normal methylation patterns:  Risk reduced but not excluded Consider CDKN1C gene sequencing and deletion/duplication Consider chromosome analysis	Normal methylation patterns:  Risk reduced but not excluded Consider UPD analysis of chromosome 7

#### Limitations

Molecular mechanisms causing BWS or RSS that do not affecting methylation patterns are not assessed, including:

- Maternal UPD of chromosome 7
- · Chromosomal translocations, inversions, deletions, or duplications
- Pathogenic CDKN1C sequence variants, deletions/duplications
- Diagnostic errors can occur due to rare sequence variations.

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