



WOMEN'S HEALTH AND GENETICS

# Reveal<sup>®</sup> SNP Microarray Pediatric

Revealing answers early to help shorten  
the journey to diagnosis



SNP microarray  
is a first-line test for  
pediatric patients  
presenting with  
developmental delay,  
intellectual disability  
and/or autism.<sup>1-3</sup>

~1 in 6

~1 in 6 children has a  
developmental delay from  
mild speech delay to serious  
intellectual disability<sup>4</sup>

~1 in 44

~1 in 44 children aged  
8 years are diagnosed with  
autism spectrum disorder<sup>5</sup>





# Labcorp offers a robust test menu and over 35 years of diagnostic experience to meet your pediatric genetic testing needs

Labcorp offers breadth and depth of diagnostic services whether a suspected genetic abnormality requires standard karyotyping, fluorescence in situ hybridization (FISH), single-gene or panel testing, metabolic testing, microarray or whole-exome sequencing.

In some patients, indications for testing may be apparent early in life, but in some cases, the phenotype evolves over time, and making a diagnosis becomes more challenging. In these cases, it may take years to uncover a definitive diagnosis,

resulting in a costly and often frustrating journey for providers, patients and parents alike. Reveal Pediatric SNP Microarray provides a high-resolution, whole-genome analysis to identify copy number and copy neutral abnormalities to help provide diagnostic insights.

Reveal Pediatric can be performed via a cord blood sample or buccal swab in a neonate, or via a buccal swab or venous sample on an older child. Early diagnosis supports appropriate management and the provision of a realistic prognosis.

Reveal Pediatric—Features	
<b>335,000+ arrays analyzed</b>	Extensive experience and database of abnormalities from specimens tested over time providing an exceptional resource to support interpretation
<b>Flexible specimen type</b>	Ability to analyze blood, as well as a buccal swab sample to eliminate the stress of collecting a blood sample on a young child
<b>High success rate</b>	> 99.95% success rate on both blood and buccal samples providing highly reliable results
<b>Low rate of variants of uncertain significance (VUS)</b>	Low VUS rates of ~7.4% due to comprehensive variant database
<b>One microarray utilized for all sample types</b>	One microarray platform for prenatal and pediatric testing, providing a robust database for interpretation and enabling reanalysis of data after delivery using pediatric cut-offs, without the need for a new sample

Prenatal, postnatal, products of conception

Services	
<b>Access to genetic experts</b>	In-house lab genetic counselors, medical geneticists, and lab directors available to support your results interpretation and patient management protocols
<b>2,000+ patient service centers</b>	Convenient specimen draw sites nationwide for patients
<b>400+ managed care plans</b>	Broad in-network coverage and patient-friendly financial programs to increase patient access to genetic testing

# Benefits of Reveal Pediatric

- Easy-to-understand reports provide clinically relevant interpretation support
- Whole genome coverage helps resolve marker chromosome origin and identify unbalanced rearrangements undetectable by routine cytogenetics<sup>1,6,7</sup>
- Detects copy neutral changes which may be associated with increased risk for autosomal recessive conditions, uniparental disomy (UPD) and identity by descent (IBD)
- Reanalysis of a prenatal sample using pediatric cut-offs for deeper interpretation, without requiring a new sample or incurring additional cost

# Why choose Labcorp for your SNP microarrays?

## Depth of experience

- **35+** years' experience performing genetic testing
- **11+** years' experience in microarray testing
- **300+** combined years' cytogenetics experience for array laboratory directors

## Access to experts

- **~140** genetic counselors nationwide
- **~40** genetic counselors dedicated to clinician support
- **15+** lab directors involved in microarray analysis

# Reveal Pediatric is a clinically validated, highly reliable test, supported by professional societies

In a study involving over 21,000 patients referred for the investigation of developmental delay/intellectual disability, multiple congenital anomalies and/or autism spectrum disorder, the diagnostic yield of SNP microarray was 12.2%, about 10% more than standard karyotype alone.<sup>2</sup> In a separate study of over 36,000 patients with developmental delay/intellectual disability, SNP microarray identified a pathogenic finding in 19% of patients.<sup>8</sup>

When a clinical phenotype does not fit a specific syndrome, microarray analysis may identify the etiology.

Clinical indications for Reveal Pediatric include<sup>1-3</sup>:

- Multiple anomalies not specific to a well-defined syndrome
- Nonsyndromic developmental delay and/or intellectual disability
- Autism spectrum disorders
- Dysmorphic facial features

- Abnormal phenotype with apparently balanced translocation or marker chromosome
- Child with symptoms that resemble, but are not a perfect fit for, a particular phenotype





## Specimen requirements

Direct Testing		
<b>Specimen quantity*</b>	<b>Blood</b> <ul style="list-style-type: none"> <li>• Children: 2–5 mL (less for newborns)</li> <li>• Infants: 2 mL</li> <li>• Blood should be collected in a Sodium Heparin (green top), EDTA (lavender top) or ACD-A (yellow top) tube</li> </ul> <p>*Special studies may call for an additional specimen. Contact the lab to discuss requirements.</p>	<b>Buccal swab</b> <ul style="list-style-type: none"> <li>• Minimum of 2 buccal swabs required</li> <li>• Buccal swab collection kit contains instructions for the use of a buccal swab</li> </ul>
<b>Expected turnaround time*</b>	14–17 days	

\*Turnaround time is calculated from the time the specimen arrives at the Center for Molecular Biology and Pathology (CMBP) in North Carolina.

A completed questionnaire for Reveal Pediatric SNP Microarray should accompany the specimen to aid in interpretation. The form is available on our website, or by calling 800-345-GENE (4363), or through your Labcorp sales representative.

Labcorp is contracted with more than 400 managed care plans and regularly bills third-party payers for testing reimbursement. This could help minimize or eliminate your patient's expense.

## In 2023, 60% of patients paid \$0 for Reveal Pediatric.

\*Based on managed care claim data in 2023. For 45% of patients, insurance paid in full and 15% of patients had no patient responsibility (non-covered and coverage-related).<sup>9</sup>



## Specifications

<b>Probe density</b>	<ul style="list-style-type: none"> <li>• 2.7 million+ copy number and allele-specific genomic markers</li> <li>• 2 million+ structural probes to detect copy number variants</li> <li>• ~ 750,000 SNP probes to detect copy number and copy neutral changes, triploidy and IBD</li> <li>• 100% ISCA constitutional gene and X chromosome coverage</li> </ul>
<b>Specimen types</b>	<ul style="list-style-type: none"> <li>• Whole blood</li> <li>• Buccal swab</li> </ul>
<b>Reporting cut-offs for copy-number changes</b>	<ul style="list-style-type: none"> <li>• <math>\geq 25</math> kb for genes with established clinical significance</li> <li>• Deletions: <math>\geq 200</math> kb with at least one OMIM gene</li> <li>• Duplications: <math>\geq 500</math> kb with at least one OMIM gene</li> </ul>
<b>Identification of copy neutral changes</b>	<ul style="list-style-type: none"> <li>• Regions of homozygosity suggestive of uniparental disomy or identity by descent, associated with an increased risk for recessive disease</li> </ul>
<b>Exon analysis</b>	<ul style="list-style-type: none"> <li>• Specific genes may be analyzed at an exon level upon request</li> </ul>
<b>Susceptibility genes</b>	<ul style="list-style-type: none"> <li>• Reported when associated with a clinical syndrome that has a clear phenotype</li> </ul>
<b>Family studies policy</b>	<ul style="list-style-type: none"> <li>• Familial testing using appropriate technologies is performed at no charge when required to interpret the proband's result</li> </ul>

Women's Health and Genetics client	Labcorp client
476 Reveal® SNP Microarray Pediatric	510002 SNP Microarray — Pediatric (Reveal®)
120 Cytogenetics Blood Chromosome Analysis	052045 Chromosome Analysis with Reflex to SNP Microarray — Pediatric (Reveal®)
	511535 Chromosome Five-Cell Count Plus Microarray (Reveal®) Whole Blood


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#### Call Us

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
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