

Test Patient

Sex Assigned at Birth: Female
Date of Birth: 01/01/1970
Sample ID: SM09123
Sample Type: BLOOD
Collection Date: 02/01/2024
Received Date: 02/03/2024

**Clinic: Medical Genetics
Center**

Physician: Test Doctor, M.D.
Phone: 510-555-0000
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Requisition ID:
RQ-0000059

Report Number:
RP828

Report Date:
09/30/2025

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TEST INFORMATION

MyOme Rare Disease, Whole Genome Analysis, Trio

Indication for testing: Developmental delay, autism, microcephaly, seizures

Positive

Clinically relevant variant(s) detected

LOCATION	CONDITION (MODE OF INHERITANCE)	VARIANT	ZYGOSITY	INHERITED FROM	CLASSIFICATION
chr15	Angelman syndrome (Imprinting)	UPD	N/A	Paternal	Pathogenic

INTERPRETATION

Paternal UPD of chromosome 15 was detected.

A single long contiguous stretch of homozygosity was detected on chromosome 15. No copy number gain or loss was identified in this region. Trio analysis demonstrated that all chromosome 15 sequences are paternally inherited, consistent with paternal uniparental disomy of chromosome 15 (patUPD15). PatUPD15 is associated with Angelman syndrome and is consistent with the reported clinical features in this individual.

NEXT STEPS

- These results should be interpreted in the context of this individual's clinical findings, family history, and other laboratory data.
- Genetic counseling is recommended to discuss the significance of these results.

VARIANT SUMMARY

seq[GRCh38]15x2upat, Pathogenic

EVIDENCE

Paternal UPD of chromosome 15 is associated with Angelman syndrome. This condition is characterized by severe developmental delay, intellectual disability, speech impairment, gait ataxia, tremulousness of the limbs, and unique behavior (frequent laughing, smiling, excitability, and hand flapping). Seizures and microcephaly are also common. Developmental delay is often the first symptom and presents around 6 months of age ([PMID: 16470747](#), [24876791](#), [27615419](#), [29696750](#)). There are published consensus guidelines for Angelman syndrome ([PMID: 35150089](#)).

TEST METHODS

- Specimen receipt, accessioning, data analysis, and interpretation is performed by MyOme Inc., 1505 Adams Drive, Suite B1, Menlo Park, CA 94025, CLIA# 05D2203070. Whole Genome Sequencing, excluding data analysis and interpretation, is performed by Broad Clinical Labs LLC, 27 Blue Sky Dr, Burlington, MA 01803, CLIA#22D2055652.
- Genomic DNA obtained from submitted samples is sequenced using Illumina technology. Reads are aligned to the NCBI GRCh38 reference assembly.
- Information about the patient's phenotype is used to prioritize variants across a large number of genes. Variants are interpreted and reported based on the standards and guidelines set forth by the American College of Medical Genetics and Genomics (ACMG). Classification categories include pathogenic (P), likely pathogenic (LP), variants of unknown significance (VUS), likely benign (LB) and benign (B). Reported variants only include those which are classified as P, LP, or VUS, overlap with the tested individual's indication for testing and are consistent with the expected pattern of inheritance (when parental samples are submitted). For mitochondrial genome variants, only known P and LP variants are reported.
- Tandem repeat expansions (TREs) in the following genes are reported when they overlap the clinical indication for testing and fall within the reportable range: AFF2, AR, ATN1, ATXN1, ATXN2, ATXN3, ATXN7, ATXN8OS, C9orf72, CACNA1A, DIP2B, DMPK, FMR1, FXN, HTT, JPH3, LRP12, PABPN1, PPP2R2B, and TBP. This test does not report on the status of repeat expansion interruptions.
- All reported variants are confirmed, if necessary, by a secondary technology: SNVs are confirmed using Sanger sequencing; CNVs are confirmed using arrays, MLPA, PCR, or long-read sequencing, depending on the nature of the copy number variant; TREs are confirmed using long-read sequencing.
- Methylation analysis is performed using Oxford Nanopore long-read sequencing on genomic DNA from whole blood. Reads are aligned to the NCBI GRCh38 reference assembly, and methylation data is extracted via modified basecalling. Methylation analysis is performed as a reflex test when a VUS is identified in a gene for which a genome-wide methylation signature has been validated, or when a copy number variant or region of homozygosity associated with an imprinting condition, currently limited to Angelman syndrome and Prader-Willi syndrome, is identified. Methylation findings are incorporated as evidence towards the assessment and classification of variants identified by whole genome sequencing. The results of methylation analysis are used to provide functional epigenetic information to further support variant classification.
- For the mitochondrial genome, the following quality control metrics are generally achieved: an average read depth of >3,000x and a minimum acceptable read depth of 1000x. Heteroplasmy is estimated using variant allele fraction.
- Nuclear genome mean depth of coverage: 33.8X; 92.7% of bases with coverage of at least 10X.

TEST LIMITATIONS

- This test is designed to detect clinically relevant single-nucleotide variants, small insertions and deletions, and copy number variants (CNVs) across the genome, single-nucleotide variants and small insertions and deletions in the mitochondrial genome, and tandem repeat expansion (TRE) in a select set of genes.
- There are certain regions that are not well covered and will not be analyzed such as segmentally duplicated regions.
- The sensitivity of this test to detect deletions and duplications may vary depending on the depth of coverage, the size of the variant or other inherent sequence properties. For example, sensitivity to detect all CNVs 50-100 bp in size and duplications < 1kb is reduced.
- This analysis does not detect balanced alterations (reciprocal translocations, Robertsonian translocations, inversions, and balanced insertions) or other complex structural variants, methylation abnormalities, genomic imbalances in segmentally duplicated regions, and mosaicism of the nuclear genome; possible cases of mosaicism may be investigated at the discretion of the laboratory director. Sensitivity to detect variants may be reduced in low complexity regions such as homopolymer regions.
- This analysis does not detect deletions, duplications, or complex rearrangements of the mitochondrial genome. Although sensitivity for detection of >5% heteroplasmic single-nucleotide variants is expected to be high based on validation studies, we cannot guarantee that these low-level heteroplasmic variants will always be identified due to paralogy with the nuclear genome.
- This analysis does not detect variants in the following regions in the mitochondrial genome: chrM:1-576, chrM:16024-16569.
- Methylation analysis is a reflex test and is only performed when a qualifying variant, copy number variant, or region of homozygosity is identified that meets the criteria described in the Methods section. Epimutations will not be detected.
- A history of stem cell or bone marrow transplantation, or recent blood transfusion may impact the accuracy of the results.
- Like most tests, this test carries a risk of false negative or false positive results, which may be caused by, without limitation, sample contamination from biological or non-biological sources, specimen marking issues, rare genetic variants interfering with analysis, and other technical issues and limitations.

DISCLAIMERS

- This test was developed, and its performance characteristics were determined, by MyOme, Inc., a clinical laboratory certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA) and College of American Pathologist (CAP) accredited to perform high complexity clinical laboratory testing. This test has not been cleared or approved by the U.S. Food and Drug Administration (FDA).
- Like most tests, this test carries a risk of false negative or false positive results. Testing is unavailable for samples damaged by human error, lost/destroyed due to weather, transit issues or other problems beyond the control of MyOme. Test results should always be interpreted by a clinician in the context of clinical and familial data with the availability of genetic counseling when appropriate. MyOme is not responsible for the content of third-party websites referenced in this report.
- The interpretation of variants is based on our current understanding of the genome. These interpretations may change over time as more information about these alterations becomes available. Possible diagnostic errors include variant call errors, sample misidentification, and other sources.

REVIEWED BY



MyOme Example Lab Director

09/30/2025

Date