

{#patient}

Whole Exome Sequencing Report

Patient Information

- **Name:**
- **Sex/Age:**
- **Date Received:**
- **Indication:**

Sequencing Platform

Whole Exome Sequencing on the Illumina Novaseq 6000 NGS Platform

Clinical Indication:

Likely Pathogenic Variant Detected **Related** to the Clinical Phenotype

Key Findings

Gene&Transcript	Location	Variant	Zygosity	Inhertance	ClinicalSignificance
{gene.geneAndTranscript}					{d.clinicalSignificance}

Genetic test results are reported based on the recommendations of American College of Medical Genetics

Variant Interpretation & Clinical Correlation

****:

The submitted sample showsvariant inof gene().

Variant Characteristics:

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Variant Evidence

Gene Impact

- **Gene:**
- **Effect:**

- ****Exon:****
- ****Transcript:****
- ****Protein:****
- ****Coding:****

Based on above evidence, this variant (:) is classified as.

OMIM Phenotype:

Recommendations

- 1.
- 2.
- 3.

Methodology

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Limitations/Disclaimer

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Computer
performance