

Client: Example Client ABC123 123 Test Drive Salt Lake City, UT 84108 UNITED STATES

Physician: Doctor, Example

Patient: Patient, Example

DOB 6/23/2020 Gender: Female

Patient Identifiers: 01234567890ABCD, 012345

Visit Number (FIN): 01234567890ABCD **Collection Date:** 00/00/0000 00:00

Cytogenomic Molecular Inversion Probe Array FFPE Tissue - Oncology

ARUP test code 3004275

Cytogenomic MIP Array, FFPE

Normal (Ref Interval: Normal)

Test Performed:CytogenomicMolecular Inversion Probe Array, FFPE Tissue - Oncology(FFPEARRAY) Specimen Type:TumorRight Kidney EstimatedTumorContent: 100 percent Indication for Testing:Post treatment Wilm's tumor

RESULT SUMMARY

Normal Microarray Result (Female)

RESULT DESCRIPTION

No clinically significant copy number changes or regions of homozygosity were detected.

INTERPRETATION

This analysis showed normal result.

Health care providers with questions may contact an ARUP genetic counselor at (800) 242-2787 ext. 2141.

Cytogenomic Nomenclature (ISCN) arr(X,1-22)x2

TechnicalInformation

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-This assay was performed using theOncoScan(TM)CNV
Assay(ThermoFisher Scientific) according to validated protocols
within the Genomic Microarray Laboratory at ARUP Laboratories
-This assay is designed to detect alterations to DNA copy number
state (gains and losses) as well as copy-neutral alterations
(regions of homozygosity; ROH) that indicate an absence- or
loss-of-heterozygosity (AOH or LOH)
- Copy-neutral LOH (CN-LOH) may be present due to acquired UPD
(segmental or whole chromosome)

(segmental or whole chromosome)
- AOH may be present due to parental relatedness (consanguinity)
or uniparentaldisomy(UPD)

-The detection sensitivity (resolution) for anyparticular genomicregion may vary dependent upontumor burden, the number of probes (markers), probe spacing, and thresholds for copy number and ROH determination

-TheOncoScanCNVarray containsover220,000SNP probeswitha medianprobedensity (kb/probe)of16-19kb
- Genome-wide resolution varies from approximately300-400kb for

copy number changes and approximately5Mb for ROH for samples with high tumor contenttoseveral Mbfor samples with lower tumor content(greater than 50 percent tumor content is recommended for this assay)

- The limit of detection forclonality(mosaicism)varies dependent upon the size and type of genomic imbalance. In general, genotype mixture due to mosaicism (distinct cell lines from the

H=High, L=Low, *=Abnormal, C=Critical

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same individual) or chimerism (cell lines from different individuals) will be detected when present at greater than30percent in thesample
-Genomic coordinates correspond to the Genome Reference Consortium human genome build 37/human genome issue 19 (GRCh37/hg19)

Variant Classification and Reporting Criteria - Variant analysis is performed in accordance with recommendations by the American College of Medical Genetics and Genomics (ACMG), using tiered classification terminology
- Acquired/somatic or constitutional/germline cancer-associated - Acquired/Somatic or constitutional/germine cancer-associated copy number variants (CNVs) and ROH are classified and reported using the following clinical significance categories: Clinically Significant CNVs and/or ROH (Tier 1 and Tier 2 Variants) and Other Clonal Variants (Tier 3) Constitutional/germline CNVs not associated with cancer are classified according to the ACMG recommended 5-tier classification system: pathogenic, likely pathogenic, variant of uncertain significance (VUS), likely benign, and benign—In general, only constitutional CNVs classified as pathogenic or likely pathogenic will be reported using the following clinical significance category: Other Variants (Likely

Constitutional) - Constitutional CNVs conferring non-cancer recessive disease risk will generally not be reported
-CNVsclassified asTier 4,likely benign or benignthat aredevoid of relevant gene content or reported as common findings in the

general population, are generally notreported

- ROH are generally reported when known or suspected to be mosaic and representative of CN-LOH - Total autosomal homozygosity (only autosomal ROH greater than5Mb are considered for this estimate) consistent with AOH at a level of greater than 10 percent will generally be reported; AOH less than 10 percent may be reported, dependent uponon the concern for masked CN-LOH and/or a recessive disorder

Limitations

This analysis cannot provide structural (positional) information associated with genomic imbalance. Therefore, additional cytogenetic testing by chromosome analysis or fluorescence in situ hybridization (FISH) may be recommended.

Certain genomic alterations may not or cannot be detected by this technology. Thesealterations may include, but are not limited to:

-CNVs below the limit of resolution of this platform -Sequence-level variants (mutations) including point mutations andindels

-Low-level mosaicism (generally, less than3Opercent)
-Balanced chromosomal rearrangements (translocations, inversions

and insertions) -Genomic imbalance in repetitive DNA regions (centromeres, telomeres, segmental duplications, and acrocentric chromosome short arms)

This result has been reviewed and approved by

A portion of this analysis was performed at the following

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INTERPRETIVE INFORMATION: Cytogenomic Molecular Inversion
Probe Array, FFPE Tissue
Oncology
For detection of copy number alterations and loss of
heterozygosity in FFPE specimens.

This test was developed and its performance characteristics
determined by ARUP Laboratories. It has not been cleared or
approved by the US Food and Drug Administration. This test was
performed in a CLIA certified laboratory and is intended for
clinical purposes.

EER Cytogenomic MIP Microarray, FFPE

EERUnavailable

Block ID

22SU-1007 A9

VERIFIED/REPORTED DATES				
Procedure	Accession	Collected	Received	Verified/Reported
Cytogenomic MIP Array, FFPE	22-119-147877	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00
EER Cytogenomic MIP Microarray, FFPE	22-119-147877	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00
Block ID	22-119-147877	00/00/0000 00:00	00/00/0000 00:00	00/00/0000 00:00

END OF CHART

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