

PATIENT Zubaida Jan REPORT DATE 10 July 2019

BOOKING ID 011907070109

Test Description

MSI testing is used for Hereditary Cancer screening (Hereditary Non-Polyposis Colorectal Cancer -HNPCC or Lynch syndrome); As a biomarker (Prognostic and predictive biomarker for the response of Immunotherapy)

Patient Demographic

Name: Zubaida Jan Sex: Female

Date of Birth/Age: 50 years Disease: Carcinoma Urinary Bladder

Clinician

Clinician Name: Dr Manish Singhal Medical Facility: Apollo Hospital Pathologist: Not Provided

Specimen

Site: Bladder Mass

Sample Type: FFPE block S 10807/18 Date of Collection: 07-07-2019 **Date of Booking**: 07-07-2019

iMSI Rapid™ Assay

Result

Microsatellite status - Stable

BIOMARKER FINDINGS

ACVR2A	No mutation detected
BTBD7	No mutation detected
DIDO1	No mutation detected
MRE11	No mutation detected
RYR3	No mutation detected
SEC13A	No mutation detected
SULF2	No mutation detected

INTERPRETATION

Mutations are not detected in any of the 7 markers

*MSS <2 of the 7 markers demonstrate instability

#MSI-H ≥2 of the 7 markers demonstrate instability

*Microsatellite stable

Microsatellite Instability-High

For valid batch test results specific controls are being run with every batch.

METHODOLOGY

Multiplex detection of seven mononucleotide repeats using molecular beacon probe-based polymerase chain reaction followed by high resolution melt-curve analysis. The assay uses seven novel biomarkers ACVR2A, BTBD7, DIDO1, MRE11, RYR3, SEC31A and SULF2 as this set of biomarkers is stable over different cancer types and ethnicities and show high performance than other known assays like Bethesda Panel. This test is carried out on Idylla platform using the MSI/1.0 Cartridge based kit which is CE IVD approved.

REFERENCES

Zhao et al. (2014) eLife 3: e02725, 1-26. De Craene B. et al. (2018) ASCO Abstract #e15639.

Zhao et al. (2018) ASCO Abstract #e15654

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Dr Gulshan Yadav, MD, Consultant Pathology

Date