

PATIENT Ishwar Chand Gurg REPORT DATE 29 August 2019 BOOKING ID 011908270432

## **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: Ishwar Chand Gurg

Sex: Male

Date of Birth/Age: 61 years

Disease: Moderately differentiated Rectal lesion

### Clinician

Clinician Name: Dr Archit Pandit Medical Facility: Max Hospital Pathologist: Not Provided

### **Specimen**

Site: Rectal Lesion

Sample Type: FFPE block H 5201/19 B Date of Collection: 27-08-2019 Date of Booking: 27-08-2019

# 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