

Sample Receipt Details:

 POD : _____ Temp : _____
 Date & Time : _____ Sample Type : _____
 CS _____ Logistics _____
 Name & Sign: _____ Name & Sign : _____
 Prenatal Sample Yes No **Bill type**

TEST REQUISITION FORM

Disease Segment* _____

Each sample must be accompanied by this completed requisition. * Fields are mandatory

Test Details

EGFR gene analysis (Hot Spot) - 4 exons (18, 19, 20, 21)

Test Name:*	<input type="text"/>	Test Code:*	<input type="text" value="MGM190"/>
Sample type:	<input type="checkbox"/> Blood (in EDTA tube) <input type="checkbox"/> Blood (in Streck tube) <input type="checkbox"/> Amniotic Fluid <input type="checkbox"/> CVS <input type="checkbox"/> Fetal Blood (PUBS) <input type="checkbox"/> Maternal blood for MCC (please send for prenatal studies) <input type="checkbox"/> Fresh Frozen Tissue <input type="checkbox"/> Saliva	<input type="checkbox"/> DNA, Specify Source: _____ <input type="checkbox"/> Cultured CV <input type="checkbox"/> Products of Conception (POC), specify tissue: _____ <input type="checkbox"/> Other sample type (specify site) _____	<input type="checkbox"/> Buccal swab <input type="checkbox"/> Cultured amniocytes <input checked="" type="checkbox"/> FFPE tissue Block (Block no.) <input type="checkbox"/> DBS/FTA PR1584 One Wax Block

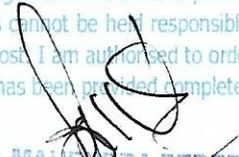
 Patient had a blood transfusion Yes No Date of last transfusion ____ / ____ / ____ (minimum 3 days of wait time is required for genetic testing)
 Has he/she undergone allogenic bone marrow transplant: Yes No.

Patient Details
Name:* Mrs. Chitra Pathirana (In Capital Letters) D.O.B. DD MM YY Age: 60Y/F Gender: M / F
Address:
Phone: **E-mail I.D:**

Clinician Details
Clinician's Name:* Prof. Jayantha Balawardana **Hospital Affiliation:** Aegle Omics Pvt Ltd
Address:
Phone :
Email id :

Date of sample collection * 4/11/2024YY

I understand that the current analysis is limited to variants which co-relate with disease phenotype/symptoms/terms as mentioned in the clinical details provided by me. Incidental findings which may or may not be actionable are not routinely reported. They can however be provided on request after informed consent from the patient/guardian. As disease phenotype may evolve over time, the appearance of new symptoms/signs may alter test results or their significance: MedGenome laboratories cannot be held responsible for this. A re-analysis or a re-test may be required due to the former; this will be performed (if deemed necessary) at an additional cost. I am authorised to order the above tests as I am the treating physician/consulting physician in this case. I confirm that the patient/guardian (in case of minors) has been provided complete information regarding the test, including its limitations in a language of their understanding.


Dr. MAHEERA PERERA
 MBBS (Cey), MD (Coy), Dip RT
 Consultant in Clinical Oncology & Radiotherapy

Medical Professional Signature* _____ **Date:** _____ **Place:** _____

Clinical notes/diagnosis:

Disease affection status **Parental consanguinity present** **Age of manifestation:** _____
Affected Siblings **Details:** _____

GOVERNING LAW, JURISDICTION AND DISPUTE RESOLUTION

These Terms and Conditions and this Test Requisition Form shall be governed by and construed in accordance with Indian law and the courts in Bangalore shall have exclusive injunctive jurisdiction. In the event of any dispute, controversy or claim whatsoever arising from these Terms and Conditions and/or this Test Requisition Form, the parties shall undertake to make every effort to reach an amicable settlement within fifteen (15) days upon reference of the dispute by any party through discussions among the concerned representatives of parties, failing which the dispute, controversy or claim shall be settled by Arbitration by a Sole Arbitrator appointed by the 'President-Arbitration Centre-Karnataka', Bangalore as per Indian Arbitration and Conciliation Act, 1996 as amended from time to time. The venue of arbitration shall be Bangalore and it shall be conducted in English language. The award passed by the Sole Arbitrator shall be final and binding upon the parties.

NOTICE

All notices, statements or other communication required or permitted to be given or made shall be in writing and in English language. Such notices will deliver by hand or sent by prepaid post with recorded delivery, or facsimile transmission addressed to the intended recipient at the address mentioned in this Test Requisition Form.

INDEPENDENT PARTIES

All parties effected hereunder are independent entities and neither of the parties are an agent, employee or joint venture of the other and they shall not represent themselves as such to any third parties.

REFUND

Refund of fees for any reason has to be claimed by the Patient or the guardians of the Patients within 90 days from the date of delivery of report.

Patient/Guardian Authorization

By my signature below I attest to the following:

I have read and I understand the information provided on this form.

Patient Consent (sign here or on the consent document)

I have read the Informed Consent document and I give permission to MedGenome to perform genetic testing as described. I also give permission for my specimen / genetic data to be used in (de-identified) studies at MedGenome to improve genetic testing for other patients.

By agreeing to this informed consent below, I am confirming that I understand the benefits, risks and limitations associated with genetic testing. Furthermore, I am affirming that I recognize the seriousness of conditions for which {I am/my child} being tested, and that disease descriptions, prognoses, and treatment options have been made available to me by {my/my child's} health care provider. Finally, if I have the legal authorization to provide this informed consent on behalf of another person, I am attesting that the sample provided belongs to that person.

Patient/Guardian Name Mrs. Chitra Pathirana

First Name	Middle Name	Last Name	Date of Birth: mm/dd/yyyy

Patient/Guardian Signature*	Date:	Place:

Father Name	Mother Name

Signature*	Date and time
Signature*	Date and time

Relationship with the proband 

Note :

Signature of both parents is requested for prenatal testing.

For trio testing, each parent should provide separate informed consent for the sequencing of his or her sample.

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Prof. Jayantha Balawardane
MBBS, MD(Col), FRCP (Edin), FRCP (Lond), FSLCO
Professor in Oncology
Head of the Department of Oncology
Faculty of Medicine
University Hospital
General Sir John Kotelawala Defence University

CONFIDENTIAL LABORATORY REPORT

PATIENT NAME : MRS. I. CHITRA PATHIRANA
AGE : 69 YEARS
BILL NO :
OPD / IP : OPD
REFERRED BY : DR DHAMMIKE RASNAYAKE
COLLECTED TIME : 2024-11-04 17:11
GENDER : FEMALE
LAB REF NO : EH_MC_266984
WARD :
REPORTED TIME : 2024-11-11 07:52

TEST : HISTOPATHOLOGY

Clinical history :- Left lung tumour, involving both upper and lower lobes.

Specimen :- Left lung bronchial biopsy for histology.

Macroscopy :- Received four brownish tissue pieces together measuring 7 x 4 x 2 mm. All passed in 1 block.

Microscopy :- Sections reveal possible fragments of bronchial mucosa, focally covered by respiratory epithelium. Sub epithelium shows an infiltrating carcinoma, composed of very few cohesive sheets and many irregular small glands lined by atypical epithelial cells with moderate nuclear atypia in a background of marked desmoplasia. These cells are rounded with vesicular nuclei, prominent eosinophilic nucleoli and moderate thin cytoplasm. Necrosis, vascular emboli or perineural invasion are not seen.

Diagnosis :- **Left lung, bronchial, biopsy:-**
- Adenocarcinoma, acinar growth pattern-see comment.

Comment :- Suggest TTF1 for confirmation as lung in origin.

RP - 1584/24

G.P.H Ramani

Dr. Ramani Punchihewa
MBBS, D.Path, MD (Histopathology)
Consultant Histopathologist

**** Blocks and slides of this specimen will be retained as follows;
Slides - Five years, Blocks - Ten years & Specimen - Three months from
the date of the report issued ****.