

Phone 866.776.590
Fax 239.690.4237

FOLR1 IHC CDx Sponsored Testing Program Request Form

immun•gen

Program Description:

Eligible patients may receive one (1) FOLR1 FDA (ELAHERETM) for Ovarian Carcinoma test regardless of test results or treatment decision. Patients must meet all of the following criteria to be eligible:

· Patient has ovarian cancer (including epithelial ovarian cancer, primary peritoneal cancer or primary fallopian tube cancer)

Date:

- Patient lives and receives treatment in the United States or a US Territory
- Patient does not have a known FR α expression from a previous test
- · Patient has not previously been tested under this Program

No patient, health care program, or beneficiary shall be billed for this test. This test shall not be included in a bundled payment to any health care facility including, but not limited to, a hospital. The ordering physician shall not be compensated any fees in connection with this testing, such as for specimen collection, handling, or data reporting. Program is not valid where prohibited by law. NeoGenomics and Immunogen reserve the right to rescind, revoke, or amend the program for any reason without notice.

Client Information	
Required Information Account #: Account Name:	
Street Address:	
City, ST, ZIP:	
Phone: Fax:	
Additional Reporting Fax:	
Requisition Completed by:	Date:
Ordering Physician (please print: Last, First):	NPI #:
Treating Physician (please print: Last, First):	NPI #:
The undersigned certifies that he/she is (1) licensed to order agrees to the terms in Program Description; (2) that such test treatment of this patient; and (3) patient meets eligibility requi	(s) are medically necessary for the care/

Specimen Information	ONCOLOGY OFFICE & PATHOLOGY TO COMPLETE
Oncology office to complete Specimen ID and	Collection Date when possible.
Specimen ID:	Block ID:
Fixative/Preservative:	_ Retrieved Date: mm / dd / yyyy
Collection Date: mm / dd / yyy	Collection Time: 🗖 AM 🗖 PM
□ Slides # Unstained	Stained 🗖 H&E
🗆 Primary 🛛 Metastasis – If Metastasis, I	ist Primary:
Paraffin Block(s) #:	
Predictive Marker Fixation (CAP/ASCO Req *Indicated markers/panels/profiles require fixatio.	
Cold ischemic duration (mins):	Unknown
Fixative: 10% NBF Other:	
Fixation duration (hours):	Unknown

3rd Party Specimen Location ONCOLOGY OFFICE TO COMPLETE

Clinical Information

Authorized Signature:

perform the services described.

Required: Please attach patient's pathology report (required), clinical history, and other applicable report(s). ICD 10 (Diagnosis) Code/Narrative (Required):					
Reason for Referral:					
□ New Diagnosis		Relapse	🗖 In Rem	iission	Monitoring
Staging: 🗖 0				D IV	Note:

Patient Information

Last Name:		🗆 Male	🗆 Female	
First Name:	M.I.	Other Pt ID/Acct #:		
Date of Birth: mm / dd	_ / уууу	Medical Record #:		
By completing this section, Client represents it has obtained informed consent from patient to perform the services described herein.				

Client will arrange separate tissue shipment to NeoGenomics OR
Complete the following so that NeoGenomics Client Services will request specimen from
Pathology site. Please fax this completed requisition and pathology report to 239.690.4237.

Location of Specimen:		
Street Address:		
City:	ST:	ZIP:
Phone:	Fax:	
Body Site:		

Select Testing

□ FOLR1 IHC CDx, Sponsored Testing Program⁺

Specimen Requirements

A formalin-fixed, paraffin-embedded (FFPE) tissue block is the preferred specimen type

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One (1) unbaked, unstained slide for H&E staining (required) and two to three (2-3) positively charged unstained slides (all cut at 4-5 microns)

Block and slide identifiers should be clearly written and match exactly with the specimen ID and specimen labeling as noted on the requisition.

Test Notations

Specimen Usage

NeoGenomics makes every effort to preserve and not exhaust tissue, but in small and thin specimens, there is a possibility of exhausting the specimen in order to ensure adequate material and reliable results.