



# Hereditary Cancer Test Request Form

- Please submit both pages of this form
- Make sure information is complete and legible

NOTE: Affix Patient Identifier Label to Specimen Tube

FOR LAB USE

SPECIMEN COLLECTION DATE (REQUIRED)

(MM/DD/YYYY)

At the time of specimen collection:  Hospital Inpatient (>24 hour stay) Discharge date: / / (MM/DD/YYYY)  Hospital Outpatient  Non-Hospital Patient

## 1. Patient Information (Complete information required)

Name (last)	Name (first)	(m.i.)	Gender <input type="checkbox"/> Male <input type="checkbox"/> Female	Birthdate (MM/DD/YYYY)	Patient ID #
Email		Cell phone		Daytime phone	
Address			City	State	Zip

## 2. Ordering Provider Information (Only name and HCP Account # required unless you're a new customer or HCP # is unknown)

Name (last)	Name (first)	Myriad HCP Account #	Degree	NPI #
Address		City	State	Zip
Office Contact Name	Phone	Fax	Email	

## 3. Send Results To (Optional - additional clinician can be listed to receive test status updates and the patient's copy of the test results)

Name (last)	Name (first)	Myriad HCP Account #	Degree	NPI #
Address		City	State	Zip
Office Contact Name	Phone	Fax	Email	

## 4. Test Requested (For test descriptions see reverse)

Tests ordered will be processed and billed based on payer criteria.

Myriad myRisk® Hereditary Cancer Panel Testing	Single-Syndrome Testing	Additional Tests*
<b>HEREDITARY BREAST AND OVARIAN CANCER SYNDROME CRITERIA:</b> <input type="checkbox"/> Integrated BRCAAnalysis® with Myriad myRisk® Hereditary Cancer Update Test <b>FOR PATIENTS OF ASHKENAZI JEWISH ANCESTRY:</b> <input type="checkbox"/> Multisite 3 BRCAAnalysis® <input type="checkbox"/> REFLEX to Integrated BRCAAnalysis® with Myriad myRisk® Hereditary Cancer Update Test if the Multisite 3 is negative <b>LYNCH SYNDROME OR MYH-ASSOCIATED POLYPOSIS (MAP) CRITERIA:</b> <input type="checkbox"/> COLARIS®PLUS with Myriad myRisk® Hereditary Cancer Update Test <b>FAMILIAL POLYPOSIS SYNDROME CRITERIA:</b> <input type="checkbox"/> COLARIS AP®PLUS with Myriad myRisk® Hereditary Cancer Update Test <b>FOR PATIENTS PREVIOUSLY TESTED AT MYRIAD:</b> <input type="checkbox"/> Myriad myRisk® Update Test (Available to patients previously tested negative with BRCAAnalysis®, COLARIS®, and/or COLARIS AP®. Full BRCA1/2 duplication and deletion analysis and/or PMS2 testing will be included in the test order unless previously performed or restricted by payer criteria.) <b>RISK ANALYSIS OPTIONS (to be excluded on report, see reverse for details):</b> <input type="checkbox"/> riskScore® is not appropriate for this patient <input type="checkbox"/> Tyrer-Cuzick and riskScore® are not appropriate for this patient	<b>HEREDITARY BREAST AND OVARIAN CANCER SYNDROME CRITERIA:</b> <input type="checkbox"/> Integrated BRCAAnalysis® <b>FOR PATIENTS OF ASHKENAZI JEWISH ANCESTRY:</b> <input type="checkbox"/> Multisite 3 BRCAAnalysis® <input type="checkbox"/> REFLEX to Integrated BRCAAnalysis® if the Multisite 3 is negative <b>LYNCH SYNDROME OR MYH-ASSOCIATED POLYPOSIS (MAP) CRITERIA:</b> <input type="checkbox"/> COLARIS®PLUS <b>FAMILIAL POLYPOSIS SYNDROME CRITERIA:</b> <input type="checkbox"/> COLARIS AP®PLUS	<input type="checkbox"/> <b>Single Site Testing:</b> Specify Gene: _____ Mutation: _____ <b>Relationship:</b> My patient is the _____ (e.g. maternal aunt) of the known mutation carrier. <b>Required: Include a copy of the known mutation carrier's report.</b> <input type="checkbox"/> <b>Other:</b> (e.g. single gene analysis)

## 5. Confirmation of Informed Consent & Statement of Medical Necessity

I affirm each of the following: I have provided genetic testing information to the patient and the patient has consented to genetic testing. This test is medically necessary for the diagnosis of a disease or syndrome. The results will be used in the patient's medical management and treatment decisions. The person listed as the Ordering Physician is authorized by law to order the test(s) requested herein.

SIGN HERE: Medical Professional (required to process form)

X

Date: / / (MM/DD/YYYY)

(Signature date is the specimen collection date if a different date is not provided above)

## 6. Billing/Payment Information

**OPTION 1: BILL INSURANCE** (Please attach copy of authorization/referral)

Name of Policy Holder: / / (MM/DD/YYYY) DOB: / / (MM/DD/YYYY)

Insurance ID#: Patient Relation to Policy Holder:  Self  Spouse  Child  Other Authorization/Referral: / / (MM/DD/YYYY)

SIGN HERE: Patient/Responsible Party I AGREE TO THE BILLING TERMS ON REVERSE

X

DATE: / / (MM/DD/YYYY)

Reminder: Include a copy of BOTH SIDES of your insurance card(s).

If you submit more than one card, indicate which is primary.

I understand that Myriad will contact me if I will be financially responsible for any non-covered service. To be considered for the Myriad Financial Assistance Program, please provide the following information: Annual household income \$ . Number of family members in household .

**OPTION 2: PATIENT PAYMENT** (Please call Customer Service for questions regarding test prices or for credit card payment)

**OPTION 3: OTHER BILLING** (To establish an account, submit billing information with this form)

Bill our institutional account #: or established research project code #: or Authorization/Voucher #:



# Testing for Myriad myRisk® Hereditary Cancer

## IMPORTANT INFORMATION FOR PATIENT\*

**BILLING TERMS:** I represent that I am covered by insurance and authorize Myriad Genetic Laboratories, Inc. (MGL) to give my designated insurance carrier, health plan, or third party administrator (collectively "Plan") the relevant health information necessary for reimbursement. I authorize Plan benefits to be payable to MGL. I understand MGL will contact me if I will be financially responsible for any non-covered service. By agreeing to testing I also authorize Myriad to obtain a consumer credit report on me from a consumer reporting agency selected by Myriad. I understand and agree that Myriad may use my consumer credit report to confirm whether my income qualifies me for financial assistance. I further understand that this is not a credit application and will not impact my credit score. I agree to assist MGL in resolving insurance claim issues and if I don't assist, I may be responsible for the full test cost. I permit a copy of this authorization to be used in place of the original.

**NON-DISCRIMINATION:** Federal law (GINA) and laws in most states prohibit discrimination regarding employment eligibility, health benefits, or health insurance premiums based solely on genetic information. Myriad Genetic Laboratories, Inc. (Myriad) complies with applicable Federal civil rights laws and does not discriminate on the basis of race, color, national origin, age, disability, or sex.

## AFFORDABILITY: Myriad Promise™

- The majority of appropriate patients pay \$0
- Myriad will work with your insurance provider to help you get the appropriate coverage
- If you encounter ANY financial hardship associated with your bill, Myriad will work with you toward your complete satisfaction
- For more information please refer to the billing information at MyriadPromise.com

\*Translation of Billing Terms are available in Mandarin and Spanish at MyriadPromise.com. Myriad also provides free language services to people whose primary language is not English through qualified interpreters. If you need these services, contact Customer Service at 800-469-7423.

## TEST DESCRIPTIONS (For a full list of tests offered, visit [www.myriadpro.com](http://www.myriadpro.com))

<b>Integrated BRACAnalysis®:</b> Analysis of <i>BRCA1</i> and <i>BRCA2</i> for susceptibility to Hereditary Breast and Ovarian Cancer syndrome.
<b>Integrated BRACAnalysis® with Myriad myRisk® Hereditary Cancer Update Test:</b> Analysis of <i>BRCA1</i> and <i>BRCA2</i> for susceptibility to Hereditary Breast and Ovarian Cancer syndrome with additional genes associated with hereditary cancer risk (see table below). May also include riskScore® breast cancer analysis. See below details for inclusion criteria.
<b>Multisite 3 BRACAnalysis®:</b> Three-mutation <i>BRCA1</i> and <i>BRCA2</i> analysis for individuals of Ashkenazi Jewish ancestry: <i>BRCA1</i> c.68_69del (p.Glu23Valfs*17) (aka <i>BRCA1</i> 185delAG, 187delAG); <i>BRCA1</i> c.5266dupC (p.Gln1756Profs*74) (aka <i>BRCA1</i> 5382insC, 5385insC); <i>BRCA2</i> c.5946del (p.Ser1982Argfs*22) (aka <i>BRCA2</i> 6174delT).
<b>COLARIS®PLUS:</b> Analysis of <i>MLH1</i> , <i>MSH2</i> , <i>MSH6</i> , <i>PMS2</i> , <i>MUTYH</i> , and <i>EPCAM</i> for susceptibility to Lynch syndrome (HNPCC) and <i>MYH</i> -Associated Polyposis (MAP).
<b>COLARIS®PLUS with Myriad myRisk® Hereditary Cancer Update Test:</b> Analysis of <i>MLH1</i> , <i>MSH2</i> , <i>MSH6</i> , <i>PMS2</i> , <i>MUTYH</i> , and <i>EPCAM</i> for susceptibility to Lynch syndrome (HNPCC) and <i>MYH</i> -Associated Polyposis (MAP) with additional genes associated with hereditary cancer risk (see table below). May also include riskScore® breast cancer analysis. See below details for inclusion criteria.
<b>COLARIS AP®PLUS:</b> Analysis of <i>APC</i> for susceptibility to FAP/AFAP.
<b>COLARIS AP®PLUS with Myriad myRisk® Hereditary Cancer Update Test:</b> Analysis of <i>APC</i> for susceptibility to FAP/AFAP with additional genes associated with hereditary cancer risk. May also include riskScore® breast cancer analysis. See below details for inclusion criteria.
<b>Single Site Testing:</b> Analysis of single, familial mutation.
<b>Myriad myRisk® Update Test:</b> Analysis of 35 hereditary cancer genes for patients who previously tested negative for <i>BRACAnalysis®</i> , <i>COLARIS®</i> , and/or <i>COLARIS AP®</i> . Full <i>BRCA1/2</i> duplication and deletion analysis and/or <i>PMS2</i> testing will be included in the test order unless previously performed or restricted by payor criteria.

Certain payers do not cover genetic testing when Single Nucleotide Polymorphisms (SNPs) are a component of the test. For payers who do not reimburse for a hereditary cancer test due to SNP analysis inclusion, Myriad will report the myRisk Hereditary Cancer Test without SNPs and these patients will not receive a SNP based riskScore®. Please visit [www.myriadpro.com/payeroptout](http://www.myriadpro.com/payeroptout) to determine if your patient's payer does not reimburse for hereditary cancer genetic testing with SNP analysis.

Genes & Associated Cancers*	Br	Ov	Co	En	Me	Pa	Ga	Pr
<i>BRCA1</i>	•	•				•		•
<i>BRCA2</i>	•	•				•		•
<i>MLH1</i> , <i>MSH2</i> , <i>MSH6</i> , <i>PMS2</i> , <i>EPCAM**</i>		•	•	•			•	•
<i>APC</i>			•				•	
<i>MUTYH</i>			•					
<i>CDK4</i> , <i>CDKN2A (p16INK4a, p14ARF)</i>					•	•		
<i>TP53</i>	•	•	•	•	•	•	•	•
<i>PTEN</i>	•		•	•	•			
<i>STK11</i>	•	•	•	•			•	•
<i>CDH1</i>	•		•				•	
<i>BMPR1A</i> , <i>SMAD4</i>			•			•	•	
<i>PALB2</i> , <i>ATM</i>	•					•		
<i>CHEK2</i>	•		•					
<i>NBN</i>	•							•
<i>BARD1</i>	•							
<i>BRIP1</i>		•						
<i>RAD51C</i> , <i>RAD51D</i>		•						
<i>POLD1</i> , <i>POLE</i> , <i>GREM1</i>			•					
<i>AXIN2</i> , <i>GALNT12</i> , <i>MSH3</i> , <i>NTHL1</i> , <i>RPS20</i> , <i>RNF43</i>			•					
<i>HOXB13</i>								•

Br: Breast / Ov: Ovarian / Co: Colorectal / En: Endometrial / Me: Melanoma / Pa: Pancreatic / Ga: Gastric / Pr: Prostate

\*Additional risks may be associated with each gene/syndrome. \*\*Large rearrangement only.

## Turnaround Time:

- The majority of Myriad myRisk® results are completed within 14 days
- We will notify you in the unusual event results take longer than 21 days

## Myriad myRisk® Report includes:

- myRisk Genetic Result
- riskScore® Result
  - Personalized breast cancer risk assessment based on an analysis of biomarkers combined with patient clinical and family history data
- myRisk Management Tool
  - Guideline based (NCCN, CAPS, Amsterdam, and others) cancer management for both positive and negative results
  - Includes a Tyrer-Cuzick breast cancer risk estimate

## Completing the Test Request Form:

- Please include:
  - Age, cancer diagnosis, ancestry, gender, and cancer family history

The myRisk Management Tool and riskScore may not be reported without an accurate and specific personal and family history included on the Patient Cancer Family History in Sections 7 - 11.

riskScore® is calculated for women under age 85, of solely White/Non-Hispanic and/or Ashkenazi Jewish ancestry, without a personal history of breast cancer, LCIS, hyperplasia, atypical hyperplasia, or a breast biopsy with unknown results. riskScore® is not calculated if a woman or blood relative is known to carry a mutation in a breast cancer risk gene. riskScore® and Tyrer-Cuzick model will not be calculated if provider indicates that they are not appropriate for the patient by selecting the check box in Section 4. Not all data collected on the TRF is incorporated into Tyrer-Cuzick or riskScore® calculations. Some fields may be used for anonymized, internal validation studies only.

- The genes associated with Myriad myRisk® Hereditary Cancer Panel are subject to change. To ensure you have a current version of the TRF and the genes included with the Myriad myRisk panel please visit [www.myriadpro.com/documents-and-forms/test-request-forms](http://www.myriadpro.com/documents-and-forms/test-request-forms) and [www.myriadpro.com/myrisk/why-myriad-myrisk/gene-selection](http://www.myriadpro.com/myrisk/why-myriad-myrisk/gene-selection).
- For additional information visit [MySupport360.com](http://MySupport360.com) and [MyriadPro.com](http://MyriadPro.com)

