Please submit both pages of this form	page <b>1</b> o	f 2								
Make sure information is complete ar	nd legible	1 2								
FOR LAB USE	SPECIMEN COLLECTION DATE (RE	QUIRED)								
i i										
- <u> </u>	(MM/DD/YYYY)				NOTE: A	ffin Daklan	t Idautifian Iah	al da Carad	iman Tuba	
1. Patient Information (Complete	e information required)				NUIE: AI	mix Patien	t Identifier Lab	ei to speci	imen lube	
Name (last)	Name (first)	(m.i.)	Gender Male	Birthdate	e (MM/DD/	/YYYY)		Pat	ient ID #	
Email	Call	phone	Female			avtime r	phono			
Enlan	Cell	priorie				аушпе р	onone _			
Address			City					State	Zip	
2. Ordering Provider Informat	CION (Only name and HCP Ad	ccount # re	equired unless	you're a	new cu	stomer	or HCP #	is unk	nown)	
Name (last)	Name (first)		Myriad HCP Acco		Degree		NPI#			
			City					State	Zip	
Address			City					State	ΣΙΡ	
Office Contact Name	Phone	Fax			Email					
3. Send Results To (Optional - add	litional clinician can be listed to	o receive te	est status upda	ates and	I the pati	ient's c	opv of the	e test re	esults)	
Name (last)	Name (first)		Myriad HCP Acco		Degree		NPI#			
			City		1 1	1		State	Zip	
Address			City					State	ΖΙΡ	
Office Contact Name	Phone	Fax			Email					
4. Test Requested (For test des		,	Tosts ord	orod wil	l bo proc	ensend :	and billad	hasad (	on payer cr	itoria
□ BRACAnalysis CDx* - BRCA1 and BRCA Results of the test are used as an aid in i or Talzenna* (talazoparib). In addition, re for treatment/maintenance with Lynparz associated with enhanced progression-fi □ Myriad myRisk* Update Test - Analysis required by payer medical policy, myRisk	dentifying breast cancer patients as all the test are used as an eas (olaparib) or Rubraca* (ruca ree survival (PFS) from Zejula* of additional hereditary cancer.	nts who ar n aid in ide aparib). A (niraparib er genes fo	e or may becontifying ovariate positive BRAC or with Rubror patients who	ome elig an cance CAnalysi aca® (ru o have k	lible for the patients CDx responding to the patients of the p	treatments who esult in maint	ent with L are or ma ovarian c enance th h BRACA	ynparz ay beco ancer p nerapy. nalysis	za® (olapariome eligib patients is CDx®. Who	le also en
test has not been reviewed, cleared or a Risk Analysis Options (to be excluded or	•		kScore® is not rer-Cuzick and					or this	natient	
			Ter Guziek arie				лорпасе т		patient	
5. Confirmation of Informed	d Consent & Staten	nent of	f Medical	Nec	essity	/				
I affirm each of the following: I have provi is medically necessary for the diagnosis of decisions. Assay result may have implicat listed as the Ordering Physician is authori	f a disease or syndrome. The rions concerning the patient's s	esults will susceptibili	be used in the ty to Heredita	patient	t's medic	cal mar	nagement	and tr	eatment	n
(required to process form)		(Signatu	re date is the spe	cimen col	lection dat	te if a dif	ferent date	is not pr	ovided above	e)
6. Billing/Payment Informa	tion									
OPTION 1: BILL INSURANCE (Please attack										
Name of Policy Holder:			00B:	/				CC	<b>eminder:</b> Inc opy of <u>BOTH</u>	I SIDES
,				abian /Defe	-1.	(M	IM/DD/YYYY)		f your insura ard(s).	nce
Insurance ID#: Patie    SIGN HERE: Patient/Responsible Party   I AGREE TO THE BILLING TERMS ON REVERSE   X	nt Relation to Policy Holder: ☐ Self ☐ Spous		ATE: / .	ation/Referr	al:		4/55 0000		ou submit more than licate which is prima	
I understand that Myriad will contact me if	I will be financially responsible t			co To b	o concida	(Mi	The Mari	d Eine	ncial Assist	ance
Program, please provide the following info		-	-covered servi				-			ance
OPTION 2: PATIENT PAYMENT (Please call	Customer Service for questions	regarding	test prices or fo	or credit	card pay	ment)				
☐ OPTION 3: OTHER BILLING (To establish a	n account, submit billing informa	ation with t	his form)							
☐ Bill our institutional account #:	or established research project of	rode #·			or Authori	ization/Vou	ıcher #			

## Testing for BRACAnalysis CDx®

#### 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**

BRACAnalysis CDx\* - BRCA1 and BRCA2 gene sequence and large rearrangement analysis to identify the presence of BRCA1/2 mutation(s). Results of the test are used as an aid in identifying breast cancer patients who are or may become eligible for treatment with Lynparza\* (olaparib) or Talzenna\* (talazoparib). In addition, results of the test are used as an aid in identifying ovarian cancer patients who are or may become eligible for treatment/maintenance with Lynparza\* (olaparib) or Rubracar\* (rucaparib). A positive BRACAnalysis CDx result in ovarian cancer patients is also associated with enhanced progression-free survival (PFS) from Zejula\* (niraparib) or with Rubraca\* (rucaparib) maintenance therapy.

Myriad myRisk\* Update Test: Analysis of additional hereditary cancer genes for patients who have been tested with BRACAnalysis CDx\*. When required by medical policy, myRisk Update may be performed as a reflex with genes from the original testing excluded.

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 patents will not receive a SNP based riskScore\*. Please visit 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 <sup>*</sup>	Br	Ov	Со	En	Ме	Pa	Ga	Pr
BRCA1	•	•				•		•
BRCA2	•	•			•	•		•
MLH1, MSH2, MSH6, PMS2, EPCAM**		•	•	•		•	•	•
APC			•			•	•	
MUTYH			•					
CDK4, CDKN2A (p16INK4a, p14ARF)					•	•		
TP53	•	•	•	•	•	•	•	•
PTEN	•		•	•	•			
STK11	•	•	•	•		•	•	
CDH1	•		•				•	
BMPR1A, SMAD4			•			•	•	
PALB2, ATM	•					•		
CHEK2	•		•					
NBN	•							•
BARD1	•							
BRIP1		•						
RAD51C, RAD51D		•						
POLD1, POLE, GREM1			•					
AXIN2, GALNT12, MSH3, NTHL1, RPS20, RNF43			•					
HOXB13								•

 $\label{eq:Br.Breast} \textbf{Po:} \ \text{Ov: Ovarian / Co: Colorectal / En: Endometrial / Me: Melanoma / Pa: Pancreatic / Ga: Gastric / Pr. Prostate *Additional risks may be associated with each gene/syndrome. **Large rearrangement only.$ 

#### 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 and www.myriadpro.com/myrisk/whymyriad-myrisk/gene-selection.
- For additional information visit MySupport360.com, MyriadPro.com, and Myriad myChoice.com



# **BRACAnalysis CDx® Test Request Form**

Name (last)	ormation		re information is to		page 1) Birthdate (MM/I	DD/YYYY)					
8. Ancestry (ris	skScore® is curre	ently only	validated and pro	vided for patients of so	olely White/N	Ion-Hispan	ic and/or Ashkenazi Jewis	h ancestry)			
Select all that apply:			Black / African Hispanic / Latino	☐ Midd	le Eastern re American	ander on-Hispanic					
9. Patient Per	sonal Hi	story	of Cancer	& Other Clini	cal Info	rmatio	(Select all that apply)				
☐ No personal histor	y of cancer										
Patient has been dia	Age at Diagnosis	Patient is Currently Being Treated	Pathology / Other Info								
☐ Breast Cancer	☐ Left ☐ Right			☐ Ductal Invasive ☐ DCIS ☐ Triple-Negative (ER-, PR-, HER2-)	☐ Lobular II ☐ Bilateral ☐ Metastati		Yes □ No crine Therapy: or inappropriate				
☐ Ovarian Cancer				☐ Non-epithelial							
☐ Other Cancer				Туре							
☐ Other Cancer				Туре							
	☐ Bone marro	w transplar	nt recipient Typ	e: 🗆 Autologous 🗆 A	llogeneic (If	allogeneic p	blease call 800-469-7423 x3	(850)			
Check if applicable to patient:	☐ Blood transf	isfusion recipient Type: Whole blood Packed red blood cells Date:									
to patient.	☐ Diagnosis of	a hematol	ogic cancer Ty <sub>l</sub>	pe:	1 1 1						
10. Family His	story of C	Cancer	· 💝				nsure proper insurance rein nedical management recom				
☐ No Known Family	History of Can	cer	-				ited family history available such				
Relationship to Patio	ent	Maternal (mother's side		ancer Site or Polyp T	Age at Each						
			e) (lather's side)			·	·	Diagnosis			
								1 1 1 1			
11. Breast Car	ncer Risk	Mode	l Informat	ion							
Patient information:				INFORMATION abo	ut PATIENT'S	OTHER INFORMATION:					
Height - ft:	in:	Weight (	bs):	FEMALE RELATIVE	S:	•	graphic Density:				
Patient's age at time of first menstrual period:				Number of daught	OKC!	Has the pa □ No	atient had her breast densit	y assessed?			
Is patient ☐ Pre-menor currently: ☐ Post-menor				Number of daught	ers.		nplete one of the following	for the most			
Has this patient No had a live birth? Yes: patient's age at first child's birth:				recent assessment:  Volpara* Volumetric Density:  VAS Percentage Density:  WAS Percentage Density:  WAS Percentage Density:							
Has patient ever used Hormone Replacement Therapy? ☐ No ☐ Yes				es			☐ BI-RADS® ATLAS Density (Select one of the				
If Yes, Treatment Type: ☐ Combined ☐ Estrogen only ☐ Progesterone only				Number of mater aunts (mother's siste		following):  ☐ Almost entirely fatty ☐ Heterogeneously dense					
If Yes, is patient a: Current User: Started years ago				aunts (mother's siste	51 <b>3 )</b> .	□Sc	attered	emely dense			
_	nore years _ years ago	Number of pater aunts (father's siste	ers):	fibroglandular density Unknown  NOTE: Risk associated with mammographic density is not incorporated into riskScore (v.1), nor Tyrer-Cuzick (v.7) calculations provided on the clinical report.							
1		-		r more of the following		/A (No biop	sy or none of the listed resu	ılts)			

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