

PHONE 1.800.411.4363 FAX 1.800.434.9850

CONNECT







HEREDITARY CANCER REQUISITION

PATIENT INFORMATION (COMPLETE O	NE FORM FOR EACH PERSON TESTED)			
				1 1
Patient Last Name	Patient First Name		MI	Date of Birth (MM / DD / YYYY)
Address	City	State Patient discharged from	Zip Genetic Sex:	Phone
Accession # Ho	spital / Medical Record #	the hospital/facility: Yes No	Female (Gender identity (if different	Male Unknown ent from above):
REPORTING RECIPIENTS				
Ordering Physician		Institution Name		
Email (Required for International Clients)		Phone	Fax	
ADDITIONAL RECIPIENTS				
Name		Email	Fax	
Name		Email	Fax	
PAYMENT (FILL OUT ONE OF THE OPT	TONS BELOW)			
SELF PAYMENT	To Patient			
O INSTITUTIONAL BILLING				
Institution Name INSURANCE	Institution Code Insti		stitution Phone	Institution Contact Email
•	ent is Aware of Out-Of-Pocket Costs (exclude			
	of-pocket costs, the Baylor Genetics Billing Te ront/Back of Insurance Card(s) 2. ICD10 Diagr		atient for additional billi hysician 4. Insured Sig	
REQUIRED ITEMS 1. Copy of the Fi	ont/ back of insurance card(s) 2. ICD to blagi	. S. Name of Ordering F	nysician 4. msureu siç	Justine of Authorization
Name of Insured	Insured Date of Birth (MM / DD / YYYY)	Name of Insured	In	sured Date of Birth (MM / DD / YYYY)
Patient's Relationship to Insured	Phone of Insured	Patient's Relationship to	Insured Pl	none of Insured
Address of Insured		Address of Insured		
City	State Zip	City	St	ate Zip
Primary Insurance Co. Name	Primary Insurance Co. Phone	Secondary Insurance Co.	Name Se	econdary Insurance Co. Phone
Primary Member Policy #	Primary Member Group #	Secondary Member Polic	y # Se	econdary Member Group #
understand that I am responsible for any reasons including, but not limited to, nor	aylor Genetics to provide my insurance ca co-pay, co-insurance, and unmet deductibl n-covered and non-authorized services. I un payment for this test. Please note that Med	e that the insurance policy dictates, nderstand that I am responsible for	as well as any amount sending Baylor Geneti	ts not paid by my insurance carrier for
				11
Patient's Printed Name	Patient's S	ignature		Date (MM / DD / YYYY)
STATEMENT OF MEDICAL NECESSITY				
This test is medically necessary for the risk ass and treatment decisions. The person listed as the have consented to genetic testing.	essment, diagnosis, or detection of a disease, illnes le Ordering Physician is authorized by law to order	ss, impairment, symptom, syndrome, or dis the test(s) requested herein. I confirm that	sorder. The results will det t I have provided genetic te	ermine my patient's medical management sting information to the patient and they
Physician's Printed Name	Physician's	s Signature		//



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			/ /	
Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex
ETHNICITY				
African American	Hispanic American		Pacific Islander (Philippines, Micrones	ia, Malaysia, Indonesia)
Ashkenazi Jewish	Mennonite		South Asian (India, Pakistan)	
East Asian (China, Japan, Korea)	Middle Eastern (Saudi Arabia, Qatar, Iraq, Tu	urkey)	Southeast Asian (Vietnam, Cambodia	a, Thailand)
Finnish	Native American		Southern European Caucasian (Spai	n, Italy, Greece)
French Canadian	Northern European Caucasian (Scandinavia	n, UK, Germany)	Other (Specify):	
SAMPLE		INDICATION FO	PR TESTING (REQUIRED)	
SAMPLE TYPE	ONA (Specify):	ICD10 Diagnosis	Code(s)	
Blood in EDTA (Purple-top)		Personal Hi	story	
Buccal Swab	Saliva	Type of Can	cor	
Cultured Skin Fibroblast	Skin Biopsy [†]	Type of Can		
		Cancer Loc	ation	
1	/	Age at Diag	nosis	
Date of Collection: ' DD	YYYY			
recent blood transfusion TESTING OPTIONS	who have had a bone marrow transplant or	HEREDITARY C	ANCER TESTS CANCER PANELS 24001	
		TEST NAME		SAMPLE *
Proband Last Name	Proband First Name	Comprehen	sive Hereditary Cancer (95 genes)	BE, DNA, CF, BUC, SA
	///	Common He	ereditary Cancer (43 genes)	BE, DNA, CF, BUC, SA
Relationship to Proband	Date of Birth (MM/DD/YYYY)	BRCA1 & BI	RCA2 Panel (2 genes)	BE, DNA, CF, BUC, SA
Proband testing location (Select one	e)	High-Risk H	lereditary Breast Cancer (9 genes)	BE, DNA, CF, BUC, SA
		Hereditary	Breast/Ovarian/Endometrial Cancer (27 genes)	BE, DNA, CF, BUC, SA
Baylor Genetics		High-Risk H	lereditary Colorectal Cancer (22 genes)	BE, DNA, CF, BUC, SA
Lab #	Family #	Hereditary	Colorectal/Gastrointestinal Cancer (37 genes)	BE, DNA, CF, BUC, SA
Lab #	raility #	Hereditary	Melanoma (10 genes)	BE, DNA, CF, BUC, SA
Another Laboratory		Hereditary	Prostate Cancer (12 genes)	BE, DNA, CF, BUC, SA
Attach a copy of the Probant A positive control sample of	d test results. the Proband is requested. Please provide, if available.	Hereditary	Paraganglioma/Pheochromocytoma (12 genes)	BE, DNA, CF, BUC, SA
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		Hereditary	Renal Cancer (19 genes)	BE, DNA, CF, BUC, SA
Full Gene Sequencing		Hereditary	Endocrine Cancer (22 genes)	BE, DNA, CF, BUC, SA
		Hereditary	Pancreatic Cancer (21 genes)	BE, DNA, CF, BUC, SA
Deletion/ Duplication Analysis			Brain/Central Nervous System/Peripheral stem Cancer (25 genes)	BE, DNA, CF, BUC, SA
			Leukemia/Lymphoma (18 genes)	BE, DNA, CF, BUC, SA

^{*} This sample type incurs an additional fee and typically adds 14 days to the turnaround time, depending on sample quality.
† Baylor Genetics will store this sample for up to 14 days after the report is issued, allowing for follow-up testing if needed.



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HEREDITARY CANCER REQUISITION

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Patient Last Na	ame	Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex	
HEREDITARY	CANCER TESTS 24001	l				
SINGLE GENE	ANALYSIS					
	sive test codes are listed		•	e analysis, which includes both sequencing and dele ations codes, please obtain the test code from our w		
Test Code	Gene	Test Code	Gene	Test Code	Gene	
Test Name		Test Name		Test Name		
TEST CODE	TEST NAME		DISORDER		SAMPLE TYPE *	
6720	APC Comprehensive (Se	equence & Deletion/Duplication Analysis)	APC-Associated	Polyposis Conditions	BE	
6520	RUNX1 Sequence Analy	sis	Familial Thromb	ocytopenia with Propensity to AML	BE	
3740	FH Sequence Analysis		Hereditary Leiom	Hereditary Leiomyomatosis and Renal Cell Cancer (FH-Related Disorders)		
6705	MLH1 Comprehensive (Sequence & Deletion/Duplication Analysis)		Hereditary Non-I	Hereditary Non-Polyposis Colon Cancer (HNPCC) - Blood Analysis		
6710 & 6888		Sequence & Deletion/Duplication Analysis) uplication Analysis (by MLPA)	Hereditary Non-l	Hereditary Non-Polyposis Colon Cancer (HNPCC) - Blood Analysis		
6715	MSH6 Comprehensive (Sequence & Deletion/Duplication Analysis)		Hereditary Non-I	Polyposis Colon Cancer (HNPCC) - Blood Analysis	BE	
6795	PMS2 Deletion/Duplication Analysis		Hereditary Non-I	Polyposis Colon Cancer (HNPCC)	BE	
6890	PMS2 Comprehensive (Sequence & Deletion/Duplication Analysis)		Hereditary Non-I	Hereditary Non-Polyposis Colon Cancer (HNPCC) - Blood Analysis		
6888	EPCAM Deletion/Duplication Analysis (by MLPA)		Hereditary Non-I	Hereditary Non-Polyposis Colon Cancer (HNPCC) - Blood Analysis		
6821	TP53 Comprehensive (S	Sequence & Deletion/Duplication Analysis)	Li-Fraumeni Syn	Li-Fraumeni Syndrome (LFS)		
3665	MEN1 Sequence Analys	is	Multiple Endocri	Multiple Endocrine Neoplasia, Type 1		
3660	RET Sequence Analysis		Multiple Endocri	Multiple Endocrine Neoplasia, Type 2 (RET-Related Disorders)		
6120	MUTYH (MYH) Sequence	Analysis	MUTYH (MYH) - A	MUTYH (MYH) - Associated Polyposis		
3600	SDHB, SDHC, & SDHD Sequence Panel		PHEO and PGL S	PHEO and PGL Syndromes		
6790	PTEN Comprehensive (Sequence & Deletion/Duplication Analysis)		PTEN-Related Di	PTEN-Related Disorders		
6121	RECQL4 Sequence Anal	ysis	Rothmund-Thom	nson Syndrome (RECQL4 -Related Disorders)	BE	
6770	VHL Comprehensive (Se	equence & Deletion/Duplication Analysis)	Von Hippel-Linda	au Syndrome	BE	
SAMPLE SPE	CIFICATIONS TABLE					
ABBREVIATION	SAMPLE NAME -	RECOMMENDED AMOUNT	SHIPPING INSTRU	CTIONS SPECIAL NO	OTES	

		RECOMMENDED AMOUNT				
ABBREVIATION	SAMPLE NAME	(2 YRS - ADULT) (NEWBORN - 2YRS)		SHIPPING INSTRUCTIONS	SPECIAL NOTES	
BE	Blood in EDTA tube (purple-top)	3-5 cc	2 -3 cc	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze.		
BUC	Buccal Swab	See Special Notes	See Special Notes	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze. Sample must arrive within 72 hours.	Collect with ORAcollect•Dx (OCD-100) self- collection kit (provided by Baylor Genetics with instructions). It is highly recommend that the sample be collected by a healthcare professional.	
CF	Cultured Skin Fibroblast	See Special Notes	See Special Notes	Ship at ambient temperature (18-25°C/64-77°F) in an insulated container by overnight courier. Specimens should arrive in the laboratory within 48 hrs of collection. Do not heat or freeze.	Send two T25 flasks at 80-100% confluence	
DNA	DNA, Extracted	10 ug	10 ug	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze.		
SA	Saliva	See Special Notes	See Special Notes	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze.	Collect with Oragene DNA Self-Collection Kit.	



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INFORMED CONSENT FOR HEREDITARY CANCER TESTING

			/ /	
Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex
Your physician has advised you	to undergo genetic testing for herec	litary cancer and is red	questing testing for:	
Name of Test				
TEST INFORMATION				

This consent form will provide you with information regarding genetic testing, which you should discuss with your healthcare provider or a genetic counselor. To assist you in understanding the reason for this testing, we have provided information about the testing process and potential results below.

The purpose of genetic testing is to determine if a genetic disease may be present or if there is an increased risk for a genetic disease to occur in a patient or their family. DNA is the genetic material that we receive from our parents. Genes are made of DNA and are the instructions for maintaining the health of our body. Each person has a unique set of DNA and most of the differences in our DNA do not impact our health. Genetic testing analyzes DNA to find any abnormal changes (mutations also called variants) that might cause disease, make it more likely to develop disease, and/or increase the chance of having a child affected by disease.

The testing ordered by your healthcare provider can determine if you or your child have a variant associated with a genetic disease. "Your child" can also mean your unborn child, for the purposes of this consent.

Depending on why genetic testing is needed, you might be tested for:

- · A known variant that has already been found in your family
- A single gene or variant that causes a specific, suspected disease.
- Multiple genes at the same time. These genes might cause similar diseases or might cause diseases that are unrelated to each other.
- · Multiple types of testing that each test for different variants.

RESULTS

There are several types of test results that may be reported including:

- Positive: Positive or "abnormal" results mean there is a change in the DNA found that is related to your/your child's medical issues or that you/your child are at an increased risk of developing a disease in the future. It is possible to test positive for more than one variant. Positive results might include pathogenic variants (variants known to be associated with disease) and likely pathogenic variants (variants that are likely to be associated with disease).
- Negative: Negative or "normal" results mean no relevant variants related to your/your child's medical issues were detected or that you/your child are not expected to be at an increased risk for developing a disease in the future. This might indicate that there are no variants associated with disease in the gene(s) tested. Genetic testing, while highly accurate, might not detect a variant present in the gene(s) tested. This can be due to limitations of the information available about the gene(s) being tested, or limitations of the testing technology.
- · Variant of Uncertain Significance: Testing can detect variant(s) in DNA which we do not yet fully understand. These are also referred to as variants of uncertain significance (VUS). Additional testing may be recommended for you or your family if a VUS is identified in a gene that may be associated with your/your child's medical condition.
- Secondary / Incidental Findings: Testing can sometimes detect a variant in a person's DNA unrelated to the reason for testing. If this variant is expected to have medical or reproductive significance, it is called a secondary or incidental finding.

CONSIDERATIONS AND LIMITATIONS

- This consent form cannot be used for whole exome sequencing (WES), whole genome sequencing (WGS), or Huntington's disease testing. These tests have specific consents that are located at https://www.baylorgenetics.com/consent/.
- Results may indicate you have a genetic disease, are at increased risk to develop a genetic disease, and/or be at an increased risk to have a child with a genetic disease. It is important to understand that genetic tests, even if negative, cannot rule out every variant. It is not possible to exclude risks for all genetic diseases for you and your family members.
- Depending on the type of genetic testing performed and the results, additional genetic testing or other testing may be needed to fully understand the likelihood of your developing the disease or the severity of the disease. This additional testing might be needed for you/your child or other members of your family.
- It is recommended that you discuss genetic testing with your healthcare provider or genetic counselor before signing this consent and again after results are made available.
- · It may not always be possible to complete testing, as sometimes the sample does not have enough DNA to perform testing or other reasons. In these cases, another sample may need to be sent to the laboratory to perform testing.



will be made via secure email if possible):

 \square Email \square Phone \square Mail

BAYLOR GENETICS 2450 HOLCOMBE BLVD. GRAND BLVD. RECEIVING DOCK HOUSTON, TX 77021-2024

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I	NFORMED CONSENT FOR	R HEREDITARY CANCER TES	TING			
				/ /		
Р	atient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex	
P	ATIENT CONFIDENTIALITY AND SI	PECIMEN RETENTION				
•	family members. In rare cases,	ested, the correct interpretation of th genetic testing can reveal that the tru to share this information with the hea	ue biological relation	ships in a family are not as they wer		
•		te, however in rare cases, inaccurate of clinical/medical information, or rare		easons for this include, but are not l	mited to, mislabeled	
•	If you sign this consent form, but you no longer wish to have your sample(s) tested, you can contact the healthcare provider who ordered the test to cancel the test. If you wish to cancel testing, the laboratory must be notified of the cancellation request before 5 PM CST the business day after the sample has begun testing. If the laboratory is not notified of your cancellation request until after this time, you will be charged for the full cost of the test.					
•	will only be released to the follo representative, and (iv) those al Genetics by providing a written	r Genetics contracted partners will ha owing person(s): (i) a licensed healthca lowed access to test results by law. I request. I also understand that labora n request or HIPAA Authorization For	are provider, (ii) thos understand that I ha atory raw data, while	e authorized in writing, (iii) the patie ve the right to access any test result	nt or their personal s directly from Baylor	
•	enacted several laws that prohi	etic diagnoses have experienced prot bit discrimination based on genetic te e of this information. For more inform	est results by health	insurance companies and employers		
•	Samples will be retained in the	laboratory in accordance with the lab	oratory retention po	licy.		
•		e-identified submitted specimen may less. DNA specimens are not returned to.				
•		York State will not be included in resple. No tests other than those authori			e retained for more than	
•	By signing this consent form, I understand and agree that variants identified may also be submitted to public databases, such as ClinVar. Such submission serves to contribute knowledge to the medical community. I understand that limited clinical information is also required for the submission of information to ClinVar's database and further that the contents of this limited clinical information may, although unlikely, include information that may identify me personally.					
•	It is possible that even if the test identifies the underlying genetic cause for the disease in your family, this information may not help in predicting the progression of disease or change management or treatment of disease.					
F	INANCIAL AGREEMENT AND GUAF	RANTEE				
b o d p h to	illing, I hereby authorize Baylor Gomy insurance carrier which is refappealing any denial of benefits irectly to Baylor Genetics. I undeleart of a verification of benefits in ealth insurance plan. If my insurate on endorse the insurance check as	ept full and complete financial resporsenetics to bill my health insurance pleasonably required for billing. I addit by my insurance carrier. I irrevocablestand that my out-of-pocket costs may estigation. I agree to be financially reance provider sends a payment direct appropriate and forward such checks rendered. If I do not have health insume by Baylor Genetics.	an on my behalf, and ionally designate Ba ly assign associated ay be different than tesponsible for all am ly to me for unpaid sto Baylor Genetics v	I further authorize Baylor Genetics to ylor Genetics as my designated repr payment to Baylor Genetics, and dir- he estimated amount indicated to mo ounts as indicated on the explanatio ervices performed by Baylor Genetivithin thirty (30) days of receipt there	o release any information esentative for purposes ect that payment be made e by Baylor Genetics as n of benefits issued by my cs on my behalf, I agree of, as payment towards	
I	understand that a completed Adv	rance Beneficiary Notice (ABN) is requ	uired for Medicare pa	itients if the service is deemed not m	edically necessary.	
R	ECONTACT FOR RESEARCH CONS	ENT				
c r	ontact patients or their provider(esearch involving the sample(s) a	search relating to health, disease prevs) directly as part of this research. I and/or information associated with this e information on research at Baylor G	gree to allow Baylor is testing. I understa	Genetics to contact me or my provid nd that patients generally receive no	er(s) about possible	
lf	I wish to opt out of being reconta	acted for research purposes by Baylor	r Genetics, I understa	and that I may check the box below:		
	Please do not contact me regard	ling any research that uses information	on obtained from thi	s testing.		

For any research I may be contacted about, I prefer contact through the following methods (please check all that apply – if no choices are selected, contact



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INFORMED CONSENT FOR HE	REDITARY CANCER	TESTING		
Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYY	Y) Genetic Sex
PATIENT AUTHORIZATION				
By signing this statement of consent, I appropriate explanations from my heal provider about the availability and impor medical geneticist who can provide sinformed decision about the genetic test. I hereby give permission to Baylor General constants and the second s	thcare provider about the ortance of genetic counsel such counseling services. st(s).	planned genetic test(s) ar ing and have been provide All my questions have bee	nd possible results. I have beer ed with written information ide en answered and I have had the	n informed by my healthcare entifying a genetic counselor
Patient's Printed Name	Pa	tient's Signature		Date (MM / DD / YYYY)
Patient's Parent / Personal Representative* No	ame Pa	tient's Parent / Personal Repr	esentative Signature	//
				///
Relationship of Personal Representative to the	e Patient Or	dering Provider's Signature		Date (MM / DD / YYYY)

^{*}If you are signing as a person with legal authority to act on behalf of the patient, you may be required to provide evidence of your authority.