

HEREDITARY CANCER GENETIC REQUISITION FORM							
Practice information:							
1. Patient information:							
Last Name	First Name		S	Street Address			
City		State		Zip Code	Date of Birth (MM/DD/YY) / /		
Patient Phone #		Gender			Buccal Swab Sample Collection Date		
			☐ Male	☐ Female	(MM/DD/YY)'//		
Patient Ethnicity							
☐ White ☐ Mixed Race	□Hispanio □America	ic/Latino an Indian/Nat	itive Alaskan	☐ Asian ☐ Hawaiian/Paci	☐ Other/Unknown fic Islander ☐ African American		
2. Payment and Insurance Inf		III III Giding	IIVE AIGGRA	110000000000000000000000000000000000000	The Islander		
☐ Medicare ☐	Commercial:		☐ Othe	er D	☐ Patient Direct Pay		
☐ Bill Insurance							
Primary Insurance	ID N	Number	Group Numb	per			
Secondary Insurance	ID N	Number	Group Numb	per			
Name of Primary Insured	Reli	lationship to		red Date of Birth	red Date of Birth		
	· · · · · · · · · · · · · · · · · · ·	mary . ured	(MM/DD/YY)				
3. ICD-10 Codes (SEE SEPARATE DOC	UMENT and list all applica	able codes)					
4. Testing Options							
ALK (Lung, CNS)					Colorectal, Pancreatic, Gastric, Prostate, CNS, Renal)		
APC (Colorectal, Pancreatic, Gastric, CNS, Endo	ocrine)		!		Colorectal, Pancreatic, Gastric, Prostate, CNS, Renal)		
ATM (Breast, Pancreatic, Prostate) BARD1 (Breast, Ovarian)				I visito (ovarian, oterine, e	Colorectal, Pancreatic, Gastric, Prostate, CNS, Renal) tate, CNS, Sarcoma, Hematologic, Endocrine)		
BRCA1 (Breast, Ovarian, Pancreatic, Prostate)				NF1 (Breast, Gastric, Prost	ate, CNS, SafCuilla, Heinatologic, Endocrine,		
	BRCA2 (Breast, Ovarian, Skin, Pancreatic, Prostate)				PMS2 (Ovarian, Uterine, Colorectal, Pancreatic, Gastric, Prostate, CNS, Renal, Sarcoma, Hematologic)		
CDKN1B (CNS, Endocrine)				PTEN (Breast, Ovarian, Uterine, Colorectal, Skin, CNS, Renal)			
CDKN2A (Skin, Pancreatic, Prostate, CNS)				RET (Endocrine)			
EGFR (Lung)	·· C · · · Breedton Lung CN	Careen		SDHB (Gastric, Renal, Sarcoma, Endocrine) STK11 (Breast, Ovarian, Uterine, Colorectal, Pancreatic, Gastric)			
EPCAM (Ovarian, Uterine, Colorectal, Pancrea	tic, Gastric, Prostate, Lung, Civil	S, Renal, Sarcom	ia, Hematologic)	STK11 (Breast, Ovarian, Ot	erine, Colorectal, Pancreatic, Gastric)		
The following genes will also be tested for m	nutations:						
	DICER1, FH, FLCN, GPC3, HOXB13, M	лах, MEN1, MET, M	ITF, MUTYH, NBN, PALB?	2, PDGFRB, PHOX2B, POLD1, POLE, PRKAR1A,	, PTCH1, RAD50, RAD51C, RAD51D, RB1, RECQL4, RUNX1, SDHA, SDHAF2, SDHC, SDH		
5. Patient authorization and i	informed consen	nt					
limitations of this testing which have been explained to consistent with HIPAA, including for billing, audits, and responsible for payment of my account and any and all or denial by my insurance carrier, I hereby authorize Su underpaid claim. This authorization shall remain valid un circumstances, a patient's DNA may be used anonymou	my satisfaction by a qualified hother purposes. I hereby autho charges associated with its coll ureTox or their designee, to appuntil the charges for the orders of	I health profession porize SureTox or pollection. I hereby opeal my health pl s on this form are	onal. I hereby authorize their designee to bill y authorize my insura olan on my behalf* to e paid in full. (*SureTo	ize my physician to release personal h Il my insurance company and receive I ance company to pay SureTox or their o provide the actions and information Tox or their designee may perform this	s my acknowledgment that I have been informed of the benefits and nealth information to SureTox or their designee for any purposes, payment from them on my behalf. I acknowledge, however, that I am designee directly for services rendered. In the event of an underpaym necessary to overturn the denial or receive reimbursement for the s appeal on my behalf, but is not obligated to do so). In some be removed prior to re-testing and the DNA sample and results obtain		
will remain anonymous. Patient Name			Patient Signat	ture	Date/		
6. Physician informed conser	nt and medical no	ecessity !	statement	(Required rational ar	nd application options on back)		
Physician Certification: By their signature below, the head obtained the patient's informed consent in accordance with state at	ealthcare provider authorizes perform and local laws. I affirm each of the foll	mance of the test(s) a ollowing: I have provide	and indicates that he or ided genetic testing inform	r she has explained the purpose of the test, th rmation to the patient and the patient has con-	ne procedures, the benefits and the risks that are involved in testing to their patient sented to genetic testing. This test is medically necessary for the diagnosis or detectio Physician is authorized by law to order the test(s) requested herein.		
Authorizing Physician Name			Authorizing P	Physician Signature	Date/		
			7.				
Supre.	- -	I					
a division of Suretox haboratory		Date	D.O.B				
241 Molnar Drive, Elmwood Park, NJ 07407 P: (201) 791-7293 F: (866) 425-4630 , CLIA# 31D2063148			_	<u> </u>			

BAR CODE//////////



To the second second								
8. Genetic Counseling PRE-GENETIC COUNSELING: If genetic counseling is required by the patient's insurance company for the test ordered, the ordering provider agrees to								
•	_		e contacted so they can refer patient locally					
POST-GENETIC COUNSELING: SureTox laboratory will facilitate genetic counseing for any patient with abnormal test results (ie. Pathogenic or Likely								
Pathogenic variant) through Informed I								
Yes, please refer my patient to ID	ONA for genetic co	ounseling if test resu	Its are abnormal.					
☐ No. please do not refer my patier	nt to IDNA for gen	etic counseling if tes	t results are abnormal. I will recommend another genetic counseling resource					
for my patient.		-						
Physician Signature:	refer to IDNA for post-tes	t counseling a letter confirming	this will be sent with all reports for patients with pathogenic/likely pathogenic results. Date:					
9. Patient Personal History of Cancer & Other Clinical Information (Select all that apply.) No Personal History of Cancer								
Patient has been diagnosed with:	Diagnosis Age	Currently Being	Pathology/ Other Info					
ratione has been alagnosed with	Diagnosis Age	currently being	Ductal Invasive Lobular Invasive Premenopausal Metastatic					
☐ Breast cancer ☐ L ☐ R		☐ Yes ☐ No	☐ Bilateral ☐ DCIS					
☐ Endometrial / Uterine		☐ Yes ☐ No	☐ Tumor MSI-High or IHC Abnormal - Result: ☐ Tumor not available for MSI-High or IHC Abnormal Testing					
Ovarian Cancer		Yes No	☐ Non-epithelial					
☐ Prostate Cancer		Yes No	☐ Gleason Score : ☐ Metastatic					
			Type: ☐ Mucinous ☐ Tumor Infiltrating Lymphocytes ☐ Medullary Growth Pattern ☐ Signet ring ☐ Crohn's-like Lymphocytic Reaction					
Colon / Rectal Cancer		Yes No	☐ Tumor is MSI-High or IHC Abnormal- Result:					
			☐ Tumor not available for MSI-High or IHC Abnormal Testing					
			Cumulative Adenomatous Polyp #:					
Colon / Rectal Adenomas		☐ Yes ☐ No	☐ 1 ☐ 2-5 ☐ 6-9 ☐ 10-19 ☐ 20-99 ☐ 100+					
Hematologic Cancer		Yes No						
Other Cancer		Yes No	TYPE:					
Other Cancer		Yes No	TYPE:					
10. Family History of Cancer (Provide complete and specific information to ensure proper insurance reimbursement, determine cancer risk estimates, and optimize medical management recommendations.)								
Relationship to Patient Mater	nal Paternal	Cancer Site or Polyn	Type (add # for colon/rectal adenomas) Diagnosis Age					
Relationship to Fatient Water	nai Taternai	cancer site of 1 diyp	Type (add # for cololly rectal adellorinas)					
11. PATIENT CONSENT FOR NGS (Next Generation	n Sequencing) CAN	ICER TESTING					
What is NGS Testing: The purpose of this molecular genetics test is to ascertain if you carry any mutation(s) causing increased cancer susceptibility. This test will include analysis of relevant genes included on the cancer panel indicated above.								
General Purpose and Clinical Information. NGS refers to a test that uses massively parallel platforms, allowing sequencing of large stretches of DNA. All genes on our NGS panel have been implicated in cancer predisposition and are associated with increased lifetime cancer risk(s). If mutations are identified in more than one gene on this panel, there may not be sufficient information available to determine your precise cancer risk. Therefore, the results of this genetic test may or may not have implications for your medical management and optio including preventive screening/intervention or therapeutics based on your genetic testing result may change over time. If you are found to carry a mutation/variant in any of the genes analyzed, this may also have implications for your family members. This should be discussed with your healthcare provider. There are several types of results that can be generated as a result of genetic testing, including:								
Pathogenic/Likely Pathogenic- a mutation was identified in a gene(s) associated with increased cancer susceptibility that is clinically significant from the evidence that has been found in research papers and case-studies. This means that you may be at an increased risk of developing certain cancer(s). This risk is associated with specific cancers based upon the type of variant and which gene it is present in. Your healthcare provider will make cancer screening and medical management recommendations based on what is known about the								
mutation that was found. Variant of Unknown Significance- An alteration was identified in one or more genes; however, there is not enough information to determine whether this change is associated with an increased risk for cancer. A thorough review of the variant and the associated literature may suggest that a variant is more likely to be disease-ransing or begin. However, in some cases the significance remains unclear. Your healthcare provider will make cancer screening and medical management recommendations based on your personal and/or family history.								
may suggest that a variant is more likely to be disease-causing or benign. However, in some cases the significance remains unclear. Your healthcare provider will make cancer screening and medical management recommendations based on your personal and/or family history. Description and principle of the test: This test uses targeted next-generation sequencing (NGS) to analyze coding regions of the genes listed in this requisition. This panel represents genes with known implications in hereditary cancer risk and is intended to provide information for physicians that assist them with cancer screening decisions.								
Technical Limitations of this test: While this test is designed to identify most detectable mutations in the genes analyzed, it is still possible that there are mutations that this testing technology is unable to detect. In addition, there may be other genes associated with cancer								
susceptibility that are not included on this panel or that are not known at this time. What is required to perform this test? You will be asked to provide 2 buccal swabs containing brushings from the inside of your cheeks. DNA will be extracted from these samples and tested according to our validated SOPM and compliance policies. As a CLIA-certified								
laboratory, we strictly adhere to all the rules regarding compliance with regulations related to patient confidentiality, diagnosis coding, professional courtesy, proficiency testing and other similar regulatory requirements. Your sample and DNA will be discarded at the end of testing process and stored for no more than 60 days. In some circumstances, a patient's DNA may be used anonymously as a negative or positive control sample in future testing, but in this circumstance, all identifiers will be removed prior to re-testing and the DNA sample and results obtained will remain anonymous.								
and results obtained will remain anonymous. I consent that SureTox laboratory may use my DNA for the above	duration and purpose.	Patient Name:	Patient Signature:					
□ I grant consent to SureTox laboratory to use my sample for further research if deemed useful: Patient Name: Patient Signature: How will I obtain results from this test? Due to the complexity of DNA-based testing in general and NGS-testing in particular, as well as the important implications of the test results, these results will be reported through your designated physician or genetic counselor. To								
the extent permitted by law, all of your laboratory records and results								

Patient Attestation of Informed Consent: My signature indicates that I have received information about this test, and I have read and understood the material in this document. I have been given a full opportunity to ask questions that I may have about the testing procedure and related issues. I agree to undergo this testing. The decision to consent to, or to refuse, the above testing is entirely mine. No test(s) will be performed and reported on my sample other than the one(s) authorized by my doctor, and any unused portion of my original sample will be destroyed within 60 days of receipt of the sample by the laboratory.