		lest Requisition			01.07	,		by Ambry	ncomplete or mis Genetics	sing, testi	ng may i	e detaye	u.			
A.PATIENT INFORMATIO		.893.0276   e: sup	pport@temp	us.com			1 owerea	Бу Липоту	Genetics							
Last Name		MI		First Name												
DOB (MM/DD/YYYY)	Medical Record	#	_	Biological Sex F M			Email						Phone			
Address (Street, Unit)	(Street, Unit)				City			State			Postal (	Postal Code Co		Country		
Ancestry Ashkenazi Jewish Black/A	rican American	East Asian	Hispanic	Middl	e Easter	rn Na	ative Ame	rican	South Asian	White/Ca	ucasian	Othe	er:			
B.ORDERING PROVIDER	INFORMATIO	ON														
Ordering Provider (full legal name)														NPI #		
Facility Name				ccount #	#			Email (required for report d		t delivery)	delivery)			Fax		
Facility Address (Street, Unit)				City						State	State		Postal Code Cou		,	
Additional person to be copied								mpleted	by							
Name	Email/Fax				Facility Name					Email/Fax			Facility Name			
C.TESTING OPTIONS																
xG+ (CancerNext-Expanded®	) Add	1		_	-			-	•	equires Bl	ood (EDT	A), Saliva,	, or Culture	ed Fibrob	last (Cultured Fibr	oblast
xG (CancerNext®)  **RNAinsight® specimen requires the completion of the 'Test Requisition for Tissue Culturing' form).  +RNAinsight®: Supplemental germline RNA sequencing, powered by Ambry Genetics. Requires Blood (PAXgene® tube required for RNA).																
Familial Variant Testing (i.e (CancerNext®) or xG+ CancerN												genic or li	kely patho	ogenic va	riant on the Temp	us xG
D.SPECIMEN RETRIEVAL		specimen guide	lines for co	llection	ı instru	ctions a	nd furthe	er details	5.							
Blood / Saliva / Cultured	l Fibroblast								T							
Mobile phlebotomy Send saliva kit to patient								Patient status at time of specimen collection: Office/Non-Hospital Hospital Outpatient Hospital Inpatient Hospital Inpatient  Not yet discharged OR Discharge								
Date of Collection:	Speci	men Collection Fac	cility:						Hospital Inpa	atient _	→ No	t yet disch	narged <b>OF</b>	<b>?</b> Dischai	rge date:	
E.CLINICAL HISTORY																
Breast Colorectal Enc *Blood or saliva samples may not be appro		Polyps Hemato th active hematologic n	-	Ovarian	Pa	increatic	Prost	ate 1	No personal histo	ry of cance	er O	ther:				
Stage: I II III IV Other: Age at diagnosis:				Primary ICD-10 Codes (C, I				D, & Z codes): Additional detail				ls (pathology, number of polyps, etc.):				
Other patient history:							_		marrow or periph				Yes bone marro	No w or periph	eral stem cell transplo	ant.
F.BILLING INFORMATIO	N															
Primary insurance plan name	Policy #	cy #				Group#			Policy Holder Name				Policy Holder DOB			
Patient relationship to policy holde	Other:		Bill Type:			Insurance Hospital/Institution				Self pay/International						
G.FAMILY HISTORY  None/No known family history	Unknown	Adopted														
Relationship to patient Matern		·	Details of re	levant h	istory											
H.PRIOR PERSONAL OR	FAMILY HIST	ORY OF GEN														
No personal or family history of	Relations Self		<b>atient:</b> ly memb	oer:		Microsatellite instability Stable (MSS) Uns				anatysis: table/High (MSI-High) Unstable/Low (MSI-Low)						
Germline testing Test performed:	1	Somatic/tumor testing (inclu Test performed:				al germlir	ne findings)		unohisto present		cal staining Proteins absent:					
I.FAMILIAL VARIANT T	ESTING INF	ORMATION	Section is	require	ed if ord	dering F\	/T testin	g.								
Proband Name			Proband D					iship to Pi	roband			Probano	d Accessio	on #		
Variant Information Attaching th	e family member's	test report is recon	nmended.												No. of Variants	i:
Gene Coding DNA (c.)								Amino Acid (p.)				Transcript (NM#)				
Gene	Coding DNA (c.)	(بن				Amino /	cid (p.)				Transcr	Franscript (NM#)				
Gene					Amino Acid (p.)				Transcript (NM#)							
J.ORDERING PROVIDER	GENETIC CO	IINSELOB'S S	SIGNATII	DE AA	ID CO	NCENT										

I certify that the patient has received an explanation of the purpose, risks, and benefits of the ordered test(s). My signature below certifies medical necessity of the test(s) (including that the test results will inform the treatment plan) and that the patient has provided informed consent that meets the requirements of applicable law for Tempus or its reference lab to: (a) collect and use the patient's samples (including genetic material) and health information and perform the ordered test(s); (b) obtain, receive, and release health information (including test results) as necessary for reimbursement or the processing of increase and release and information for an indefinite period of time in accordance with applicable law; and (d) de-identify such samples and information and use and share the resulting de-identified samples and information in accordance with applicable law.

Ordering Provider's Signature: Printed Name (full legal name):

Today's Date (MM/DD/YYYY):