

A. PATIENT INFORMATION															
Last Name				MI	First Name										
DOB (MM/DD/YYYY)		Medical Record #		Biological Sex F M		Email			Phone						
Address (Street, Unit)				City		State	Postal Code	Country							
Ancestry Ashkenazi Jewish Black/African American East Asian Hispanic Middle Eastern Native American South Asian White/Caucasian Other:															
B. ORDERING PROVIDER INFORMATION															
Ordering Provider (full legal name)								NPI #							
Facility Name				Tempus Account #		Email (required for report delivery)			Fax						
Facility Address (Street, Unit)				City		State	Postal Code	Country							
Additional person to be copied					Form completed by										
Name		Email/Fax		Facility Name		Name		Email/Fax		Facility Name					
C. TESTING OPTIONS															
xG+ (CancerNext-Expanded*)		Add +RNAinsight*		xG+: 76-gene or xG: 39-gene hereditary cancer test, powered by Ambry Genetics. Requires Blood (EDTA), Saliva, or Cultured Fibroblast (Cultured Fibroblast specimen requires the completion of the 'Test Requisition for Tissue Culturing' form).											
xG (CancerNext*)		+RNAinsight*: Supplemental germline RNA sequencing, powered by Ambry Genetics. Requires Blood (PAXgene® tube required for RNA).													
Familial Variant Testing (i.e. Cascade Testing) is offered at no additional cost for blood relatives (out to 3rd degree) of patients who are found to have a pathogenic or likely pathogenic variant on the Tempus xG (CancerNext*) or xG+ CancerNext-Expanded*) test. No-cost testing is offered for 90 days from the original xG report date. Requires Blood (EDTA) or Saliva.															
D. SPECIMEN RETRIEVAL See Tempus' specimen guidelines for collection instructions and further details.															
Blood / Saliva / Cultured Fibroblast															
Mobile phlebotomy Send saliva kit to patient				Patient status at time of specimen collection: Office/Non-Hospital Hospital Outpatient Hospital Inpatient <input type="checkbox"/> → Not yet discharged OR Discharge date:											
Date of Collection:		Specimen Collection Facility:													
E. CLINICAL HISTORY															
Breast		Colorectal		Endometrial		GI Polyps		Hematologic*		Ovarian	Pancreatic	Prostate	No personal history of cancer		Other:
*Blood or saliva samples may not be appropriate for patients with active hematologic malignancies.															
Stage: I II III IV Other:		Age at diagnosis:		Primary ICD-10 Codes (C, D, & Z codes):			Additional details (pathology, number of polyps, etc.):								
Other patient history:				Personal history of allogeneic bone marrow or peripheral stem cell transplant:** Yes No											
**Using a blood or saliva sample is not appropriate for patients who have undergone an allogeneic bone marrow or peripheral stem cell transplant.															
F. BILLING INFORMATION															
Primary insurance plan name			Policy #			Group#		Policy Holder Name		Policy Holder DOB					
Patient relationship to policy holder: Self Spouse Child Other:						Bill Type: Insurance Hospital/Institution Self pay/International									
G. FAMILY HISTORY															
None/No known family history			Unknown			Adopted									
Relationship to patient		Maternal	Paternal	Age at diagnosis	Details of relevant history										
H. PRIOR PERSONAL OR FAMILY HISTORY OF GENETIC TESTING															
No personal or family history of molecular and/or genetic testing.				Relationship to patient: Self Family member:			Microsatellite instability analysis: Stable (MSS) Unstable/High (MSI-High) Unstable/Low (MSI-Low)								
Germline testing Test performed:		Results:		Somatic/tumor testing (including potential germline findings) Test performed:			Immunohistochemical staining Proteins present: Proteins absent:								
I. FAMILIAL VARIANT TESTING INFORMATION Section is required if ordering FVT testing.															
Proband Name			Proband DOB (MM/DD/YYYY)		Relationship to Proband			Proband Accession #							
Variant Information Attaching the family member's test report is recommended.									No. of Variants:						
Gene		Coding DNA (c.)			Amino Acid (p.)			Transcript (NM#)							
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J. ORDERING PROVIDER/GENETIC COUNSELOR'S SIGNATURE AND CONSENT															
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 insurance claims; (c) retain and use samples and health 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):								