

Sample Collection – Min 6 mLs of CSF
Date*: (MM/DD/YY)
Time*: XX:XX am/pm
Kit # _____
Lumbar Puncture Tube Collection # :
#1 #2 #3 #4

Belay Lab Use Only

*Required fields; otherwise, testing may be delayed

PATIENT INFORMATION				ORDERING CLINICIAN INFORMATION			
Last Name*	Middle Name	First Name*		Clinician Name*	NPI #*		
Date of Birth* (MM/DD/YY)	Sex at birth: Male Female	MRN		Account#/Clinic/Institution*	Phone #		
Phone	Email Address*			Street Address			
Street Address*				City	State	Zip	
City*		State*	Zip*	Additional Recipient		Email or Fax	
Ancestry (select all that apply): Ashkenazi Jewish Native American Asian Pacific Islander Black/African American White/Non-Hispanic Hispanic/Latino Unknown Middle Eastern Other: _____				BILLING INFORMATION			
PROVISIONAL DIAGNOSIS				SUMMIT™ 2.0 COMPREHENSIVE GENOMIC PROFILE			
Example: Glioma				SNVs, MNVs, Indels – 520 GENES CNVs – 62 GENES Fusions – 27 GENES Chromosome Arm LOSS/GAIN Key Biomarkers – TMB, MSI			
ORDER CHOICE* & SPECIMEN REQUIREMENTS				For specifications, see Summit 2.0 Comprehensive Genomic Profile at bit.ly/3ISmeX or scan this code:			
Summit™ 2.0 and/or Vantage™							
Minimum 6 mLs of CSF collected in a sterile tube via: Lumbar Puncture Ommaya reservoir Ventricular Catheter Aspiration of CSF Space: Ventricle Cistern Other				VANTAGE™ TEST CONTENT			
				MGMT promoter methylation status			
PATIENT HISTORY							
Is the patient either (check all that apply): Recurrent Relapsed Refractory Metastatic Advanced stage/grade Other:							
Cancer History Questions				Yes	No	Unk	Primary Cancer Data (complete if answered "yes" to history of cancer):
Is test ordered to guide clinical decision-making for current stage/grade of cancer treatment?							Brain CA Stage:
Is test ordered to assess treatment plans due to suspected resistance or progression?							Breast
Is patient seeking systemic treatment and/or clinical trial participation based on test results?							Colon Date of DX: (MM/DD/YY)
Does patient have a history of cancer? (If "Yes", complete Primary Cancer Data section at right)							Kidney
Has patient undergone tissue biopsy/resection? (If "Yes", attach pathology/cytology report)							Lung
Is tissue biopsy/resection infeasible or contraindicated for this patient?							Lymphoma
Has genomic profiling on tumor been completed for patient? (If "Yes", attach report)							Melanoma
Has patient received/currently receiving chemotherapy?							Prostate
Has patient received/currently receiving radiotherapy?							Other (please specify):
Was CSF cytology testing performed? If "Yes", the results are: Negative Positive							
ICD10/Diagnostic Codes:				https://www.icd10data.com/ICD10CM/Codes/R00-R99/R90-R94/R94-/R94.02			
C 70.0	Malignant neoplasm of cerebral meninges			C 71.7	Malignant neoplasm of brain stem		
C 70.1	Malignant neoplasm of spinal meninges			C 71.9	Malignant neoplasm of brain, unspecified		
C 70.9	Malignant neoplasm of meninges, unspecified			C 72.0	Malignant neoplasm of spinal cord, cranial nerves and other parts of CNS		
C 71.0	Malignant neoplasm of cerebrum, except lobes and ventricles			C 79.31	Secondary malignant neoplasm of the brain		
C 71.1	Malignant neoplasm of frontal lobe			C 79.32	Secondary malignant neoplasm of cerebral meninges		
C 71.2	Malignant neoplasm of temporal lobe			C 79.40	Secondary malignant neoplasm of unspecified part of nervous system		
C 71.3	Malignant neoplasm of parietal lobe			C 79.49	Secondary malignant neoplasm of other parts of nervous system		
C 71.4	Malignant neoplasm of occipital lobe			G 96.198	Other disorders of meninges, not elsewhere classified		
C 71.5	Malignant neoplasm of cerebral ventricles			Other:			
C 71.6	Malignant neoplasm of cerebellum						
MEDICAL NECESSITY AND ORDERING PROVIDER AUTHORIZATION							
I am the patient's treating physician and my signature certifies: (1) I am authorized under applicable law to order the tests on this test requisition form; (2) the clinical information entered on this form is accurate and this test is medically necessary; (3) I have prepared documentation demonstrating the medical necessity of the test, included it in the patient's medical record, and will make it available to Belay upon request; (4) I will use the test results to inform my treatment decisions and medical management of the patient; (5) I have explained the nature, purpose, potential benefits, risks, and alternatives to the patient, and the patient has had the opportunity to ask questions regarding the test including the collection, use, and disclosure of their sample and health information; and (6) I have obtained informed consent from the patient to have the test performed using the consent form enclosed with this test requisition form, notified the patient they may receive a copy of this informed consent for their records, and will provide a copy of this informed consent to Belay upon request.							
Ordering Clinician/Authorized Signature*:						Date*:	

By checking this box, I confirm that I have discussed Belay's billing options, including if applicable the Assay Evolution Program Protocol, with the patient, and the patient requests that Belay provide the patient with additional information regarding such options.

INFORMED CONSENT FOR SAMPLE RELEASE, TESTING, AND FINANCIAL RESPONSIBILITY

Patient Full Name	Date of Birth* (MM/DD/YY)	MRN
Address	Email	Phone

Your healthcare provider intends to order Summit™ and/or Vantage™ (Test(s)) offered by Belay Diagnostics (Belay). By signing below, you acknowledge receipt of information regarding the financial responsibility, potential risks, benefits, and limitations of testing and provide your authorization as to the matters listed in this consent. If you have any questions or need additional information about Test(s), please consult your healthcare provider before signing this consent.

Testing Overview

Summit analyzes cerebrospinal fluid samples on a molecular level to detect variants, chromosome arm loss/gain, TMB and MSI including those that are known to be associated with brain and spinal cord cancers. Vantage analyzes cerebrospinal fluid samples to assess *MGMT* promoter methylation status. The results of Summit and/or Vantage may assist your healthcare provider in choosing a treatment plan that is best for your medical condition. However, undergoing Test(s) does not guarantee that a successful treatment will be identified or that all relevant aneuploidy, mutations, and methylation signatures will be found. Test(s) may not provide information about susceptibility to developing disease in the future, and a negative result does not rule out the presence of aneuploidy, mutations, and methylation.

Financial Responsibility

Coverage for Test(s) may be available through private third-party health insurance payers. For patients who elect to use their insurance, the cost of the test will be determined by their insurance plan (i.e. the allowable amount). For patients who opt to self-pay, Belay will directly bill the patient for the cost of the Test(s), which is Twelve Hundred Dollars (\$1200) for Summit and Two Hundred and Seventy-Five Dollars (\$275) for Vantage. The patient is solely responsible for this cost. Belay will generally invoice the patient at the time the Test(s) are released to the healthcare provider. However, Belay offers a patient assistance program (BelayAccess Program) through which financial assistance may be available on the basis of need. Belay cannot determine if the patient would qualify for financial assistance unless the patient submits a complete program application. If the patient submits an application, Belay will not charge the patient until it makes a determination on the application (except the patient may be charged sooner if the patient fails to submit a complete program application within 15 days of receipt). The patient may contact Belay at contact@belaydiagnostics.com or by calling +1 (331) 320-0155 at any time with questions related to financial responsibility for Test(s). Pricing policies are available for eligible patients and are subject to change by Belay Diagnostics.

Sample Collection and Release

Belay will perform Test(s) using genomic material extracted from the patient's cerebrospinal fluid sample. By signing below you authorize Belay to work with your healthcare provider to obtain your sample and any information related to you or your medical condition that is relevant for Test(s). Performing the requested Test(s) may exhaust the sample that is sent to Belay, and additional Test(s) may not be possible unless you provide additional samples.

Disclosure of Results

Belay will report the results of Test(s) to the patient's healthcare provider, who will discuss results and next steps with the patient. Based on results, the healthcare provider will determine if any follow-up testing is appropriate. The results and other data and information generated during the performance of Test(s) may be used and disclosed in a manner consistent with our Notice of Privacy Practices, which can be found at belaydiagnostics.com/privacy-practices. Belay is under no ongoing obligation to update, revisit, or later re-evaluate Test(s) results after those results have been made available to your healthcare provider through Test(s) reports described above.

Retention of Samples and Secondary Data

New York Residents Only: I authorize Belay to retain my specimen for potential future testing, for research ordered by my healthcare provider and/or for quality control purposes. If this box is not checked, unused specimen will be destroyed 60 days after testing is completed. Opting in or out will not impact the quality of care or testing you receive.

By signing this consent below, you nevertheless authorize the use of your sample, results, and other data and information generated during the performance of Test(s) for the purposes described in this consent. At the end of the testing process, Belay may choose to destroy or return your sample, maintain it for a future Test(s) ordered by your healthcare provider, or convert it into "Residual Information and Materials" (as defined below) and retain it indefinitely.

Belay may redact information that directly identifies you from your sample, the results of Test(s), other data and information generated during performance of Test(s), and other health or demographic information that Belay receives about you to create "Residual Information and Materials." Belay may maintain and use the Residual Information and Materials for any purpose permitted by federal and state law, including but not limited to:

- Ongoing development of testing methodologies to aid in improved diagnosis of primary and metastatic brain cancers.
- Performing quality assurance, test validation, and other operations purposes.
- Conducting commercial development and research, including performing additional analyses using the Residual Information and Materials for scientific and/or research purposes.
- Aggregating the Residual Information and Materials with similar residual information from other individuals, which may be used to create, or be disclosed to, databases or datasets that are solely or jointly owned by Belay or may be submitted to public databases to advance medical research.

The Residual Information and Materials may also be shared with third-parties, including, but not limited to, pharmaceutical and medical device companies, hospitals and universities, and other entities. You are not entitled to compensation for the use of the Residual Information and Materials or rights to any products or discoveries resulting from use of the Residual Information and Materials. Notwithstanding the foregoing, Belay will retain any of your identifiable or de-identified data as required by applicable federal or state laws or regulations.

By signing below, I confirm that I have read this consent in its entirety, understand it, and have had the opportunity to speak with my healthcare provider about Test(s) including the cost and financial responsibility, purpose, risks, benefits, and testing alternatives. I understand that I may raise any future questions or concerns related to Test(s) or this consent with my healthcare provider at any time. I request that my Test(s) proceed and authorize my sample to be taken and released for the performance of Test(s). I further authorize Belay to maintain, use, and disclose my sample and any information related to my Test(s) as described in this consent.

I agree that I will be solely responsible for the full cost of Test(s). I agree and acknowledge that I am requesting this testing for the purpose of informing my diagnosis and/or further treatment if indicated by the results. I understand that Belay may charge me in this amount at the time my Test(s) results are released to my healthcare provider unless I request an application for the BelayAccess™ program. I understand that if I complete an application, Belay will not charge me until it makes a determination on my application (except that Belay may charge me sooner if I fail to submit a complete program application within 15 days of receipt). If my application is not accepted, I agree that Belay may charge me for the Test(s).

I agree that Belay may contact me at the email address or telephone number listed above for any additional information relating to my medical history that may be required for Test(s). I also understand that this consent is voluntary, treatment from my healthcare provider is not conditioned upon it, and I may opt out of it at any time by contacting Belay at contact@belaydiagnostics.com or by calling +1 (331) 320-0155.

Patient or Representative/Guardian Signature*	Date*
Representative/Guardian Printed Name and Relationship to Patient (If Applicable)*	

By checking this box, I am requesting an application for the BelayAccess™ program through which financial assistance may be available to certain patients on the basis of need.