

## Summit™ 2.0 + Ascent™ + Vantage™ Report

Patient Information	Provisional Diagnosis	Specimen	Physician Information
Name: John Smith DOB: 01/01/1990 Sex Assigned at Birth: Male MRN: 11xx22xx33	Diagnosis: Glioblastoma ICD10: C71.9	Type: CSF Collected: 01/01/2026 Received: 01/02/2026 Specimen ID: Sum+AscPos-CNS	Institution: Belay Diagnostics Referring Physician: Provider Test

### RESULT SUMMARY

## POSITIVE

Comments
<p><i>TERT</i> promoter variants, chromosome 7 gain/chromosome 10 loss, and <i>EGFR</i> amplification are characteristic alterations in IDH-wildtype glioblastoma (GBM) (WHO, see Actionability Summary). <i>CDK4</i> and <i>MDM2</i> amplifications often co-occur in <i>TP53</i>-wildtype GBM, cooperating to activate cellular growth pathways and inhibit p53 tumor suppressor function (PMID: 8504413, 37509518). Clinical correlation is required.</p> <p><i>MGMT</i> promoter methylation in GBM is associated with increased sensitivity to alkylating chemotherapy, including temozolomide, and a more favorable treatment response (PMID: 15758010, 22877848, 22578793, 19805672). Clinical correlation is required.</p> <p>While most chromosomal arm-level alterations are considered variants of unknown significance (VUS) on their own, a high level of chromosomal loss and gain as observed in this specimen indicates chromosomal instability, a key driver of metastasis across cancer types (PMID: 38924459). Clinical correlation is required.</p>

### CLINICALLY SIGNIFICANT ALTERATION DETAILS (Tier 1 or 2 per AMP/ASCO/CAP)

SNV, MNV, Indel Variants				
Alteration	VAF	Actionability Summary		
		FDA/NCCN Therapies Associated	Prognostic/Diagnostic Guidelines	Clinical Trial Options
<i>TERT</i> c.-146C>T	34.4%	No	Yes	Yes

Copy Number Variants					
Alteration	Location	Fold Change	Actionability Summary		
			FDA/NCCN Therapies Associated	Prognostic/Diagnostic Guidelines	Clinical Trial Options
<i>CDK4</i> Amplification	chr12	38.69	No	No	Yes
<i>EGFR</i> Amplification	chr7	39.59	No	Yes	Yes
<i>MDM2</i> Amplification	chr12	61.13	No	Yes	Yes

<b>Fusion Variants:</b> None
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Biomarkers					
Tumor Mutation Burden (TMB)			Microsatellite Instability (MSI)		
Not Detected	Low	High	Stable	High	

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Ascent™ Chromosome Arm Level Loss or Gain				
Alteration	Type of Relevant Genes	Actionability Summary		
		FDA/NCCN Therapies Associated	Prognostic /Diagnostic Guidelines	Clinical Trial Options
Chr7 gain / Chr10 loss	Fully Contained: <i>GATA3, FGF8, FGFR2, PTEN, RET, SUFU, EGFR, BRAF, CDK6, MET, SMO</i>	No	Yes	Yes

Vantage™ MGMT Promoter Methylation				
Status	Guidelines	Actionability Summary		
		FDA/NCCN Therapies Associated	Prognostic/Diagnostic Guidelines	Clinical Trial Options
Methylated	NCCN	Yes	Yes	Yes

### VARIANTS OF UNKNOWN SIGNIFICANCE (Tier 3)

SNV/MNVs/Indels			
<i>BCOR</i> W534S <i>FLT4</i> R1354H	<i>H3-5</i> G13S <i>IGF1R</i> I197F	<i>INHA</i> G227R <i>KDM5A</i> V1238L	<i>SPEN</i> V1840I <i>SPTA1</i> D791E

Gene Level CNVs
None

Fusions
None

Ascent™ Aneuploidy Variants of Unknown Significance				
chr1q Gain chr2p Loss	chr2q Loss chr4q Loss	chr8q Loss chr9q Loss	chr11p Loss chr14q Loss	chr17q Gain

### ACTIONABILITY SUMMARY

FDA / NCCN Therapies for the Patient's Tumor Type (Tier 1A)			
Biomarker	Therapies	Setting	Source(s)
MGMT Promoter Methylation Positive	alkylating agent	Unspecified	NCCN

FDA / NCCN Therapies with Resistance / Decreased Response (Tier 1A): None


Prognostic Implications per NCCN			
Biomarker	Prognostic Association	Diseases	Note

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
MGMT Promoter Methylation Positive	Favorable	Glioma	MGMT promoter methylation confers a survival advantage in glioblastoma.
TERT c.-146C>T	Unfavorable	Glioma	In the absence of an IDH mutation, TERT mutations in diffusely infiltrative gliomas are associated with reduced overall survival compared to gliomas lacking TERT mutations.

Diagnostic Implications per WHO		
Biomarker	Diseases	Note
EGFR Amplification	Central Nervous System Neoplasm	EGFR amplification is a diagnostic criterion for glioblastoma, IDH-wildtype. Overall, about 60% of these tumors show evidence of EGFR amplification, mutation, rearrangement, or altered splicing (PMID: 24120142). The most frequent of these alterations is EGFR amplification (PMID: 1374522), which occurs in about 40% of all IDH-wildtype glioblastomas (PMID: 24120142; PMID: 30187121). Gain of EGFR is also present in ~50% of cases of the RTK2 subtype of diffuse pediatric-type high-grade glioma, H3-wildtype and IDH-wildtype. EGFR amplification has also been identified in hemangioblastomas, and a small subset of MN1-altered astroblastomas.
MDM2 Amplification	Central Nervous System Neoplasm	Among central nervous system neoplasms, amplification of MDM2 has been observed in glioblastoma, IDH-wildtype, where MDM2 or MDM4 (inhibitors of p53) are amplified in about 15% of cases (PMID: 24120142). Suppression of apoptosis via genetic dysregulation of the p53 pathway occurs in nearly 90% of glioblastomas (PMID: 24120142). MDM2 amplification is also observed in other CNS neoplasms, such as diffuse hemispheric glioma, H3 G34-mutant.
MET Amplification	Central Nervous System Neoplasm	MET amplifications or fusions occur in several central nervous system neoplasms. They are common in high-grade IDH-mutant astrocytomas (PMID: 30343896) and diffuse, pediatric-type high-grade glioma, H3-wildtype and IDH-wildtype (PMID: 27748748), but they may also occur in adult-type IDH-wildtype glioblastomas (PMID: 29718398; PMID: 30343896; PMID: 25135958). MET fusions are a diagnostic criterion for infant-type hemispheric glioma, where structural genomic variants, often driven by focal intragenic DNA copy-number changes, result in the acquisition of fusion genes involving numerous 5' partners and MET or other RTK genes.
MGMT Promoter Methylation Positive	Central Nervous System Neoplasm	Among central nervous system neoplasms, MGMT promoter methylation is detectable in the majority of oligodendrogliomas (PMID: 15455350). The MGMT gene encodes a DNA repair protein (PMID: 24071851) and is transcriptionally silenced by promoter methylation in approximately 40-50% of IDH-wildtype glioblastomas (PMID: 24120142; PMID: 10029064; PMID: 15758010; PMID: 22294349). MGMT promoter methylation in glioblastoma is a strong predictive marker of response to alkylating agents such as temozolomide and is associated with longer overall survival (PMID: 15758010; PMID: 22877848; PMID: 22578793; PMID: 19805672; PMID: 25655102; PMID: 24068788; PMID: 25035291; PMID: 24912512; PMID: 30782343), with more than 90% of long-term survivors harboring MGMT promoter methylation (PMID: 19269895). A higher frequency of methylation (>75%) is also associated with gliomas exhibiting the glioma CpG island methylator phenotype (G-CIMP), characteristic of IDH-mutant tumors (PMID: 22810491; PMID: 24120142; PMID: 23209033). Although H3 G34-mutant diffuse hemispheric gliomas display widespread DNA hypomethylation, MGMT is often methylated and may be associated with longer overall survival in the absence of oncogene amplifications (PMID: 23079654; PMID: 28966033; PMID: 30101054; PMID: 26482474). In high-grade astrocytoma with piloid features, a methylated MGMT promoter was reported in 46% of tumors, though no association with outcome was observed, and treatment data were unavailable (PMID: 29564591). In primary diffuse large B-cell lymphoma of the CNS, MGMT promoter methylation is observed in approximately 52% of cases and may have therapeutic implications, as a subset of elderly patients responded to temozolomide monotherapy (PMID: 16858686; PMID: 9546285; PMID: 15327516; PMID: 19494841; PMID: 19841864). In pituitary adenoma/pituitary neuroendocrine tumor (PitNET), MGMT protein expression appears inversely related to temozolomide response; however, promoter methylation status does not correlate with treatment outcomes (PMID: 29046323; PMID: 29330228; PMID: 20668043).
PTEN Loss	Central Nervous System Neoplasm	Among central nervous system neoplasms, PTEN alterations are common in glioblastoma, IDH-wildtype, in which alterations of RTK genes, PI3K pathway genes, and PTEN are found in about 90% of cases (PMID: 24120142; PMID: 18772890), and PTEN in particular shows mutation/deletion in about 40% of glioblastomas (PMID: 24120142). Germline PTEN mutations are a desirable diagnostic criterion in the diagnosis of dysplastic cerebellar gangliocytoma, a pathognomonic feature of Cowden syndrome. Germline PTEN mutations occur in 25%-85% of cases of Cowden syndrome (PMID:9140396; PMID:9467011; PMID:12844284). Genetic alterations of PTEN have also been reported in cases of papillary tumors of the pineal region (PMID:25003235).
TERT c.-146 C>T	Central Nervous System Neoplasm	Among central nervous system neoplasms, TERT promoter mutation is an essential diagnostic criteria for glioblastoma, IDH-wildtype. In oligodendroglioma, IDH-mutant and 1p/19q-codeleted, TERT promoter mutation is desirable for the diagnosis. Additionally, TERT promoter mutation is also observed in diffuse midline glioma, H3 K27-altered (PMID: 28966033; PMID: 29763623 ), diffuse pediatric-type high-grade glioma, H3-wildtype and IDH-wildtype (~64% of cases) (PMID: 2840133), high-grade astrocytoma with piloid features (2 [3%] of 74, both C228T mutations) (PMID:29564591), medulloblastoma, SHH-activated and TP53-wildtype, and meningioma. In pleomorphic xanthoastrocytoma, TERT promoter mutation and less frequently TERT amplifications have been identified (PMID: 24154961; PMID: 30240866; PMID: 30051528; PMID: 32619305).
Chr7 gain / Chr10 loss	Central Nervous System Neoplasm	Among central nervous system neoplasms, whole chromosome 7 gain (trisomy 7) and whole chromosome 10 loss (monosomy 10) is one of the essential diagnostic criteria for glioblastoma, IDH-wildtype (WHO grade 4). +7/-10 is the most frequent numerical chromosome alteration in glioblastoma (PMID: 30187121), and the sensitivity and specificity for the diagnosis of IDH-wildtype glioblastoma were reported as 59% and 98% for +7/-10 (PMID: 30187121).


### CLINICAL TRIALS / INVESTIGATIONAL THERAPIES

CDK4 Amplification + CDK6 Amplification			
Therapy	Clinical Trial	Location/Sponsor	
abemaciclib	<a href="#">NCT03310879</a> (Phase 2) Study of the CDK4/6 Inhibitor Abemaciclib in Solid Tumors Harboring Genetic Alterations in Genes Encoding D-Type Cyclins or Amplification of CDK4 or CDK6	Boston, Massachusetts Dana-Farber Cancer Institute <a href="mailto:geoffrey_shapiro@dfci.harvard.edu">geoffrey_shapiro@dfci.harvard.edu</a>	 <a href="https://genomoncology.wetrials.com/v1/NCT03310879">https://genomoncology.wetrials.com/v1/NCT03310879</a>




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palbociclib	<a href="#">NCT02896335</a> (Phase 2) Palbociclib and Pembrolizumab In Central Nervous System Metastases	Boston, Massachusetts Massachusetts General Hospital PBRASTIANOS@mgh.harvard.edu	 <a href="https://genomoncology.wetrial.com/v1/NCT02896335">https://genomoncology.wetrial.com/v1/NCT02896335</a>
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### CDK4 Amplification + CDK6 Amplification + PTEN Loss

Therapy	Clinical Trial	Location/Sponsor	
everolimus + ribociclib	<a href="#">NCT05843253</a> (Phase 2) Study of Ribociclib and Everolimus in HGG and DIPG or Ribociclib and Temozolomide in DHG, H3G34-mutant	Washington D.C., District of Columbia Nationwide Children's Hospital kelsey.troyer@nationwidechildrens.org	 <a href="https://genomoncology.wetrial.com/v1/NCT05843253">https://genomoncology.wetrial.com/v1/NCT05843253</a>

### EGFR Amplification

Therapy	Clinical Trial	Location/Sponsor	
CARv3-TEAM-E T-cells	<a href="#">NCT05660369</a> (Phase 1) CARv3-TEAM-E T Cells in Glioblastoma	Boston, Massachusetts Marcela V. Maus, M.D., Ph.D. carteamingbm@mgb.org	 <a href="https://genomoncology.wetrial.com/v1/NCT05660369">https://genomoncology.wetrial.com/v1/NCT05660369</a>
ERAS-801 + surgery	<a href="#">NCT07089641</a> (Phase 1) ERAS-801 for the Treatment of Resectable and Progressive or Recurrent IDH Wildtype Grade IV Glioblastoma or Astrocytoma With an EGFR Amplification or Mutation, ERAS801-SARG Trial	Los Angeles, California Jonsson Comprehensive Cancer Center QuanLi@mednet.ucla.edu	 <a href="https://genomoncology.wetrial.com/v1/NCT07089641">https://genomoncology.wetrial.com/v1/NCT07089641</a>
verteporfin	<a href="#">NCT04590664</a> (Phase 1/Phase 2) Verteporfin for the Treatment of Recurrent High Grade EGFR-Mutated Glioblastoma	Atlanta, Georgia Emory University william.l.read@emory.edu	 <a href="https://genomoncology.wetrial.com/v1/NCT04590664">https://genomoncology.wetrial.com/v1/NCT04590664</a>



### EGFR Amplification + MET Amplification



Therapy	Clinical Trial	Location/Sponsor	
MK-0472; MK-0472 + pembrolizumab; MK-0472 + MK-1084	<a href="#">NCT05853367</a> (Phase 1) Study of MK-0472 in Participants With Advanced /Metastatic Solid Tumors (MK-0472-001)	Chicago, Illinois Merck Sharp & Dohme LLC Trialsites@msd.com	 <a href="https://genomoncology.wetrial.com/v1/NCT05853367">https://genomoncology.wetrial.com/v1/NCT05853367</a>


### EGFR Amplification + Monosomy 10 + TERT c.-146C>T + Trisomy 7



Therapy	Clinical Trial	Location/Sponsor
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
bevacizumab + temozolomide + radiation therapy; temozolomide + radiation therapy	<a href="#">NCT05271240</a> (Phase 3) Repeated Superselective Intraarterial Cerebral Infusion (SIACI) of Bevacizumab With Temozolomide and Radiation Compared to Temozolomide and Radiation Alone in Newly Diagnosed GBM	New York, New York Northwell Health <a href="mailto:jboockvar@northwell.edu">jboockvar@northwell.edu</a>	 <a href="https://genomoncology.wetrials.com/v1/NCT05271240">https://genomoncology.wetrials.com/v1/NCT05271240</a>
pembrolizumab + olaparib + temozolomide	<a href="#">NCT05463848</a> (Phase 2) Surgical Pembro +/- Olaparib w TMZ for rGBM	Boston, Massachusetts L. Nicolas Gonzalez Castro, MD, PhD <a href="mailto:lgonzalez-castro@partners.org">lgonzalez-castro@partners.org</a>	 <a href="https://genomoncology.wetrials.com/v1/NCT05463848">https://genomoncology.wetrials.com/v1/NCT05463848</a>




MDM2 Amplification			
Therapy	Clinical Trial	Location/Sponsor	
BTP-114	<a href="#">NCT02950064</a> (Phase 1) A Study to Determine the Safety of BTP-114 for Treatment in Patients With Advanced Solid Tumors With BRCA Mutations	Sarasota, Florida Placon Therapeutics	 <a href="https://genomoncology.wetrials.com/v1/NCT02950064">https://genomoncology.wetrials.com/v1/NCT02950064</a>
LP-184	<a href="#">NCT05933265</a> (Phase 1/Phase 2) Study of LP-184 in Patients with Advanced Solid Tumors	Springdale, Arkansas Lantern Pharma Inc. <a href="mailto:lyza@lanternpharma.com">lyza@lanternpharma.com</a>	 <a href="https://genomoncology.wetrials.com/v1/NCT05933265">https://genomoncology.wetrials.com/v1/NCT05933265</a>




MDM2 Amplification + MET Amplification			
Therapy	Clinical Trial	Location/Sponsor	
crizotinib + talazoparib; axitinib + talazoparib; palbociclib + talazoparib	<a href="#">NCT04693468</a> (Phase 1) Talazoparib and Palbociclib, Axitinib, or Crizotinib for the Treatment of Advanced or Metastatic Solid Tumors, TalaCom Trial	Houston, Texas M.D. Anderson Cancer Center <a href="mailto:tyap@mdanderson.org">tyap@mdanderson.org</a>	 <a href="https://genomoncology.wetrials.com/v1/NCT04693468">https://genomoncology.wetrials.com/v1/NCT04693468</a>

MET Amplification			
Therapy	Clinical Trial	Location/Sponsor	
ANS014004	<a href="#">NCT06307795</a> (Phase 1) A Study to Investigate ANS014004 in Participants With Locally Advanced or Metastatic Solid Tumors	San Diego, California Avistone Biotechnology Co., Ltd. <a href="mailto:information.center@avistonebio.com">information.center@avistonebio.com</a>	 <a href="https://genomoncology.wetrials.com/v1/NCT06307795">https://genomoncology.wetrials.com/v1/NCT06307795</a>
VERT-002	<a href="#">NCT06669117</a> (Phase 1/Phase 2) FIH Trial of VERT-002 in Patients With Locally Advanced or Metastatic Solid Tumors With MET Alterations	Washington, District of Columbia Pierre Fabre Medicament <a href="mailto:yuhua.wang@pierre-fabre.com">yuhua.wang@pierre-fabre.com</a>	 <a href="https://genomoncology.wetrials.com/v1/NCT06669117">https://genomoncology.wetrials.com/v1/NCT06669117</a>


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vebreltinib	<a href="#">NCT03175224</a> (Phase 2) APL-101 Study of Subjects With NSCLC With c-Met EXON 14 Skip Mutations and c-Met Dysregulation Advanced Solid Tumors	Los Angeles, California Apollomics Inc. <a href="mailto:clinops@apollocinc.com">clinops@apollocinc.com</a>	 <a href="https://genomoncology.wetrial.com/v1/NCT03175224">https://genomoncology.wetrial.com/v1/NCT03175224</a>
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MGMT Promoter Methylation Positive			
Therapy	Clinical Trial	Location/Sponsor	
lomustine + temozolomide + radiation therapy	<a href="#">NCT05095376</a> (Phase 3) Testing the Addition of the Chemotherapy Drug Lomustine (Gleostine) to the Usual Treatment (Temozolomide and Radiation Therapy) for Newly Diagnosed MGMT Methylated Glioblastoma	Crystal Lake, Illinois NRG Oncology	 <a href="https://genomoncology.wetrial.com/v1/NCT05095376">https://genomoncology.wetrial.com/v1/NCT05095376</a>
temozolomide; dasatinib + quercetin; fisetin; fisetin + temozolomide; dasatinib + quercetin + temozolomide	<a href="#">NCT07025226</a> (Early Phase 1) Sequential Treatments or Combinations Including Dasatinib, Quercetin, Fisetin and/or Temozolomide for the Treatment of Previously Treated Glioma With Residual Disease	Rochester, Minnesota Mayo Clinic <a href="mailto:mayocliniccancerstudies@mayo.edu">mayocliniccancerstudies@mayo.edu</a>	 <a href="https://genomoncology.wetrial.com/v1/NCT07025226">https://genomoncology.wetrial.com/v1/NCT07025226</a>
tuvusertib + temozolomide	<a href="#">NCT05691491</a> (Phase 1/Phase 2) Testing the Combination of the Anti-Cancer Drugs Temozolomide and M1774 to Evaluate Their Safety and Effectiveness	Irvine, California National Cancer Institute (NCI)	 <a href="https://genomoncology.wetrial.com/v1/NCT05691491">https://genomoncology.wetrial.com/v1/NCT05691491</a>

PTEN Loss			
Therapy	Clinical Trial	Location/Sponsor	
TER-2013	<a href="#">NCT07109726</a> (Phase 1/Phase 2) A Phase 1/2 Trial of TER-2013 in Patients With Solid Tumors Harboring AKT/PI3K/PTEN Pathway Alterations	Orlando, Florida Terremoto Biosciences Inc. <a href="mailto:clinicaltrials@terremotobio.com">clinicaltrials@terremotobio.com</a>	 <a href="https://genomoncology.wetrial.com/v1/NCT07109726">https://genomoncology.wetrial.com/v1/NCT07109726</a>
berzosertib + sacituzumab govitecan	<a href="#">NCT04826341</a> (Phase 1/Phase 2) A Phase I/II Study of Sacituzumab Govitecan Plus Berzosertib in Small Cell Lung Cancer, Extra-Pulmonary Small Cell Neuroendocrine Cancer and Homologous Recombination-Deficient Cancers Resistant to PARP Inhibitors	Bethesda, Maryland National Cancer Institute (NCI) <a href="mailto:linda.sciuto@nih.gov">linda.sciuto@nih.gov</a>	 <a href="https://genomoncology.wetrial.com/v1/NCT04826341">https://genomoncology.wetrial.com/v1/NCT04826341</a>
dostarlimab + niraparib	<a href="#">NCT05700721</a> (Phase 2) Phase II Trial of the PARP Inhibitor Niraparib and PD-1 Inhibitor Dostarlimab in Patients With Advanced Cancers With Active Progressing Brain Metastases (STARLET)	Houston, Texas M.D. Anderson Cancer Center <a href="mailto:tyap@mdanderson.org">tyap@mdanderson.org</a>	 <a href="https://genomoncology.wetrial.com/v1/NCT05700721">https://genomoncology.wetrial.com/v1/NCT05700721</a>

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ipatasertib + paclitaxel	<b>NCT05554380</b> (Phase 2) Study of Chemotherapy Plus Ipatasertib for People With Solid Tumors With PTEN/AKT Mutations, A ComboMATCH Treatment Trial	Lancaster, Ohio National Cancer Institute (NCI)	 <a href="https://genomoncology.wetrials.com/v1/NCT05554380">https://genomoncology.wetrials.com/v1/NCT05554380</a>
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## TIER 1A THERAPY DETAILS

MGMT Promoter Methylation Positive		
Therapy	Approval / Guideline Summary	Underlying Evidence
alkylating agent	Per NCCN, methylation of the MGMT promoter in glioma silences MGMT, making the tumor more sensitive to treatment with alkylating agents (Category 2A).	The NCCN guideline for alkylating agents was supported by data from a retrospective analysis of the MGMT promoter in tumor DNA by a methylation-specific polymerase chain reaction at the University Hospital of Navarre (PMID: 11070098). Clinical data demonstrated methylation of the promoter was positively correlated with the clinical response and with overall and disease-free survival; 63% (n = 12/19) of the patients with methylated tumors had a partial or complete response to carmustine, as compared with 4%; P < 0.001 (n = 1/ 28) patients with unmethylated tumors. Additionally, the median time to the progression of disease was 21 mo. for methylated gliomas vs. 8 mo. for unmethylated glioma; P < 0.001.

## TEST DETAILS

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PANEL CONTENT AND REPORTING TRANSCRIPTS				
ABL1 NM_005157.4 ^ ABL2 NM_007314.3 ACVR1 NM_001105.4 ACVR1B NM_020328.3 AKT1 NM_001014432.1 + AKT2 NM_001626.4 + AKT3 NM_005465.4 ALK NM_004304.4 ^+ ANKRD11 NM_001256182.1 ANKRD26 NM_014915.2 APC NM_000038.5 + AR NM_000044.3 + ARAF NM_001654.4 ARFRP1 NM_003224.4 ARID1A NM_006015.4 ARID1B NM_020732.3 ARID2 NM_152641.2 ARID5B NM_032199.2 ASXL1 NM_015338.5 ASXL2 NM_018263.4 ATM NM_000051.3 + ATR NM_001184.3 ATRX NM_000489.3 + AURKA NM_198433.1 AURKB NM_004217.3 AXIN1 NM_003502.3 AXIN2 NM_004655.3 AXL NM_021913.4 B2M NM_004048.2 BAP1 NM_004656.3 BARD1 NM_000465.2 BBC3 NM_001127240.2	DNAJB1 NM_006145.1 DNMT1 NM_001130823.1 DNMT3A NM_022552.4 DNMT3B NM_006892.3 DOT1L NM_032482.2 E2F3 NM_001949.4 EED NM_003797.3 EGFL7 NM_016215.4 EGFR NM_005228.3 ^+ EIF1AX NM_001412.3 EIF4A2 NM_001967.3 EIF4E NM_001130679.1 EML4 NM_019063.3 EP300 NM_001429.3 EPCAM NM_002354.2 EPHA3 NM_005233.5 EPHA5 NM_004439.5 EPHA7 NM_004440.3 EPHB1 NM_004441.4 ERBB2 NM_004448.2 + ERBB3 NM_001982.3 + ERBB4 NM_005235.2 ERCC1 NM_001983.3 + ERCC2 NM_000400.3 + ERCC3 NM_000122.1 ERCC4 NM_005236.2 ERCC5 NM_000123.3 ERG NM_001136154.1 ERRF1 NM_018948.3 ESR1 NM_001122742.1 + ETS1 NM_001143820.1 ETV1 NM_004956.4 ^	H2BC5 NM_021063.3 H3C1 NM_003529.2 H3C2 NM_003537.3 H3C3 NM_003531.2 + H3C4 NM_003530.4 H3C6 NM_003532.2 H3C7 NM_021018.2 H3C8 NM_003534.2 H3C10 NM_003536.2 H3C11 NM_003533.2 H3C12 NM_003535.2 H3C15 NM_001005464.2 H3C14 NM_021059.2 H3C13 NM_001123375.2 H3-4 NM_003493.2 HLA-A NM_002116.7 HLA-B NM_005514.6 HLA-C NM_002117.5 HNF1A NM_000545.5 HNRNPK NM_002140.3 HOXB13 NM_006361.5 HRAS NM_005343.2 + HSD3B1 NM_000862.2 HSP90AA1 NM_001017963.2 ICOSLG NM_015259.4 ID3 NM_002167.4 IDH1 NM_005896.2 + IDH2 NM_002168.2 + IGF1 NM_001111283.1 IGF1R NM_000875.3 IGF2 NM_001127598.1 IKBKE NM_014002.3 IKZF1 NM_006060.4	MYC NM_002467.4 + MYCL NM_001033082.2 + MYCN NM_005378.4 + MYD88 NM_002468.4 + MYOD1 NM_002478.4 NAB2 NM_005967.3 ^ NBN NM_002485.4 NCOA3 NM_181659.2 NCOR1 NM_006311.3 NEGR1 NM_173808.2 NF1 NM_001042492.2 + NF2 NM_000268.3 + NFE2L2 NM_006164.4 + NFKBIA NM_020529.2 NKX2-1 NM_001079668.2 NKX3-1 NM_006167.3 NOTCH1 NM_017617.3 NOTCH2 NM_024408.3 NOTCH3 NM_000435.2 NOTCH4 NM_004557.3 NPM1 NM_002520.6 NRAS NM_002524.4 + NRG1 NM_013964.3 + NSD1 NM_022455.4 NTRK1 NM_002529.3 ^ NTRK2 NM_006180.3 ^ NTRK3 NM_001012338.2 NUP93 NM_014669.4 NUTM1 NM_175741.1 ^ PAK1 NM_001128620.1	COP1 NM_022457.5 RHEB NM_005614.3 RHOA NM_001664.2 RICTOR NM_152756.3 + RIT1 NM_006912.5 RNF43 NM_017763.4 ROS1 NM_002944.2 ^ RPS6KA4 NM_003942.2 RPS6KB1 NM_003161.3 + RPS6KB2 NM_003952.2 RPTOR NM_020761.2 RUNX1 NM_001754.4 RUNX1T1 NM_175635.2 RYBP NM_012234.5 SDHA NM_004168.2 SDHAF2 NM_017841.2 SDHB NM_003000.2 SDHC NM_003001.3 SDHD NM_003002.3 SETBP1 NM_015559.2 SETD2 NM_014159.6 + SF3B1 NM_012433.2 SH2B3 NM_005475.2 SH2D1A NM_002351.4 SHQ1 NM_018130.2 SLIT2 NM_004787.1 SLX4 NM_032444.2 SMAD2 NM_005901.5 SMAD3 NM_005902.3 SMAD4 NM_005359.5 + SMARCA4 NM_001128849.1 + SMARCB1 NM_003073.3 +

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BCL10 NM_003921.4	ETV4 NM_001079675.2 <sup>^</sup>	IL10 NM_000572.2	PAK3 NM_002578.3	SMARCD1 NM_003076.4
BCL2 NM_000633.2	ETV5 NM_004454.2	IL7R NM_002185.3	PAK5 NM_020341.3	SMC1A NM_006306.3
BCL2L1 NM_138578.1	ETV6 NM_001987.4 <sup>^</sup>	INH A NM_002191.3	PALB2 NM_024675.3	SMC3 NM_005445.3
BCL2L11 NM_001204108.1	EWSR1 NM_013986.3 <sup>^</sup>	INHBA NM_002192.2	PRKN NM_004562.2	SMO NM_005631.4 <sup>+</sup>
BCL2L2 NM_001199839.1	EZH2 NM_004456.4	INPP4A NM_001134224.1	PARP1 NM_001618.3	SNCAIP NM_005460.2
BCL6 NM_001706.4	AMER1 NM_152424.3	INPP4B NM_003866.2	PAX3 NM_181457.3 <sup>^</sup>	SOC S1 NM_003745.1
BCOR NM_001123385.1	ABRAXAS1 NM_139076.2	INSR NM_000208.2	PAX5 NM_016734.2	SOX10 NM_006941.3
BCORL1 NM_021946.4	TENT5C NM_017709.3	IRF2 NM_002199.3	PAX7 NM_001135254.1	SOX17 NM_022454.3
BCR NM_004327.3 <sup>^</sup>	FANCA NM_000135.2	IRF4 NM_002460.3	PAX8 NM_013953.3 <sup>^</sup>	SOX2 NM_003106.3
BIRC3 NM_001165.4	FANCC NM_000136.2	IRS1 NM_005544.2	PBRM1 NM_018313.4	SOX9 NM_000346.3
BLM NM_000057.2	FANCD2 NM_033084.3	IRS2 NM_003749.2	PDCD1 NM_005018.2	SPEN NM_015001.2
BMPR1A NM_004329.2	FANCE NM_021922.2	JAK1 NM_002227.2	PDCD1LG2 NM_025239.3	SPOP NM_001007228.1
BRAF NM_004333.4 <sup>+</sup>	FANCF NM_022725.3	JAK2 NM_004972.3 <sup>+</sup>	PDGFRA NM_006206.4 <sup>+</sup>	SPTA1 NM_003126.2
BRCA1 NM_007294.3 <sup>+</sup>	FANCG NM_004629.1	JAK3 NM_000215.3	PDGFRB NM_002609.3 <sup>+</sup>	SRC NM_198291.2
BRCA2 NM_000059.3 <sup>+</sup>	FANCI NM_001113378.1	JUN NM_002228.3	PDK1 NM_001278549.1	SRSF2 NM_003016.4
BRD4 NM_058243.2	FANCL NM_001114636.1	KAT6A NM_006766.3	PDPK1 NM_002613.4	STAG1 NM_005862.2
BRIP1 NM_032043.2	FAS NM_000043.4	KDM5A NM_001042603.1	PGR NM_000926.4	STAG2 NM_001042749.1
BTG1 NM_001731.2	FAT1 NM_005245.3	KDM5C NM_004187.3	PHF6 NM_032458.2	STAT3 NM_139276.2
BTK NM_000061.2	FBXW7 NM_033632.3 <sup>+</sup>	KDM6A NM_021140.2	PHOX2B NM_003924.3	STAT4 NM_003151.3
EMSY NM_020193.3	FGF1 NM_001144934.1 <sup>+</sup>	KDR NM_002253.2	PIK3C2B NM_002646.3	STAT5A NM_003152.3
CALR NM_004343.3	FGF10 NM_004465.1 <sup>+</sup>	KEAP1 NM_012289.3	PIK3C2G NM_004570.4	STAT5B NM_012448.3
CARD11 NM_032415.4	FGF14 NM_175929.2 <sup>+</sup>	KEL NM_000420.2	PIK3C3 NM_002647.2	STK11 NM_000455.4
CASP8 NM_001228.4	FGF19 NM_005117.2 <sup>+</sup>	KIF5B NM_004521.2	PIK3CA NM_006218.2 <sup>+</sup>	STK40 NM_032017.1
CBFB NM_001755.2	FGF2 NM_002006.4 <sup>+</sup>	KIT NM_000222.2 <sup>+</sup>	PIK3CB NM_006219.2 <sup>+</sup>	SUFU NM_016169.3 <sup>+</sup>
CBL NM_005188.3	FGF23 NM_020638.2 <sup>+</sup>	KLF4 NM_004235.4	PIK3CD NM_005026.3	SUZ12 NM_015355.2
CCND1 NM_053056.2 <sup>+</sup>	FGF3 NM_005247.2 <sup>+</sup>	KLHL6 NM_130446.2	PIK3CG NM_002649.2	SYK NM_003177.5
CCND2 NM_001759.3	FGF4 NM_002007.2 <sup>+</sup>	KMT2B NM_014727.1	PIK3R1 NM_181523.2	TBX3 NM_016569.3
CCND3 NM_001760.3 <sup>+</sup>	FGF5 NM_004464.3 <sup>+</sup>	KMT2C NM_170606.2	PIK3R2 NM_005027.3	ELOC NM_005648.3
CCNE1 NM_001238.2 <sup>+</sup>	FGF6 NM_020996.1 <sup>+</sup>	KMT2D NM_003482.3	PIK3R3 NM_003629.3	TCF3 NM_003200.3
CD274 NM_014143.3	FGF7 NM_002009.3 <sup>+</sup>	KRAS NM_004985.3 <sup>+</sup>	PLG2 NM_002661.3	TCF7L2 NM_030756.4
CD276 NM_001024736.1	FGF8 NM_033163.3 <sup>+</sup>	LAMP1 NM_005561.3 <sup>+</sup>	PLK2 NM_006622.3	TERC
CD74 NM_001025159.2 <sup>^</sup>	FGF9 NM_002010.2 <sup>+</sup>	LATS1 NM_004690.3	PLK3 NM_006622.3	TERT NM_198253.2 <sup>+</sup>
CD79A NM_001783.3	FGFR1 NM_023110.2 <sup>+</sup>	LATS2 NM_014572.2	PMAIP1 NM_021127.2	TET1 NM_030625.2
CD79B NM_000626.2 <sup>+</sup>	FGFR2 NM_000141.4 <sup>+</sup>	LMO1 NM_002315.2	PMS1 NM_000534.4	TET2 NM_001127208.2
CDC73 NM_024529.4	FGFR3 NM_000142.4 <sup>+</sup>	LRP1B NM_018557.2	PMS2 NM_000535.5	TFE3 NM_006521.4 <sup>^</sup>
CDH1 NM_004360.3 <sup>+</sup>	FGFR4 NM_213647.1 <sup>+</sup>	LYN NM_002350.3	PNRC1 NM_006813.2	TFRC NM_003234.2 <sup>+</sup>
CDK12 NM_016507.2	FGFR5 NM_000143.3	LZTR1 NM_006767.3	POLD1 NM_001256849.1	TGFBR1 NM_004612.2
CDK4 NM_000075.3 <sup>+</sup>	FLC N NM_144997.5	MAGI2 NM_012301.3	POLE NM_006231.2	TGFBR2 NM_001024847.2
CDK6 NM_001259.6 <sup>+</sup>	FLI1 NM_002017.4	MALT1 NM_006785.3	PPARG NM_138712.3 <sup>^</sup>	TMEM127 NM_017849.3
CDK8 NM_001260.1	FLT1 NM_002019.4	MAP2K1 NM_002755.3	PPM1D NM_003620.3	TMPRSS2 NM_001135099.1 <sup>^</sup>
CDKN1A NM_000389.4	FLT3 NM_004119.2	MAP2K2 NM_030662.3	PPP2R1A NM_014225.5	TNFAIP3 NM_006290.3
CDKN1B NM_004064.3	FLT4 NM_182925.4	MAP2K4 NM_003010.3	PPP2R2A NM_001177591.1	TNFRSF14 NM_003820.2
CDKN2A NM_000077.4 <sup>+</sup>	FOXA1 NM_004496.3	MAP3K13 NM_004721.4	PPP6C NM_001123355.1	TOP1 NM_003286.2
CDKN2B NM_004936.3	FOXL2 NM_023067.3	MAP3K14 NM_003954.3	PRDM1 NM_001198.3	TOP2A NM_001067.3
CDKN2C NM_001262.2	FOXP1 NM_032682.5	MAP3K4 NM_005922.2	PREX2 NM_024870.2	TP53 NM_000546.5 <sup>+</sup>
CEBPA NM_004364.3	FRS2 NM_001278351.1	MAPK1 NM_002745.4	PRKAR1A NM_212472.2	TP63 NM_003722.4
GENPA NM_001809.3	FUBP1 NM_003902.3 <sup>+</sup>	MAPK3 NM_002746.2	PRKCI NM_002740.5	TRAF2 NM_021138.3
CHD2 NM_001271.3	FYN NM_002037.5	MAX NM_002382.4	PRKDC NM_006904.6	TRAF7 NM_032271.2 <sup>+</sup>
CHD4 NM_001273.2	GABRA6 NM_000811.2	MCL1 NM_021960.4	PRSS8 NM_002773.3	TSC1 NM_000368.4
CHEK1 NM_001114122.2 <sup>+</sup>	GATA1 NM_002049.3	MDC1 NM_014641.2	PTCH1 NM_000264.3 <sup>+</sup>	TSC2 NM_000548.3
CHEK2 NM_007194.3 <sup>+</sup>	GATA2 NM_0032638.4	MDM2 NM_002392.5 <sup>+</sup>	PTEN NM_000314.4 <sup>+</sup>	TSHR NM_000369.2
CIC NM_015125.3 <sup>+</sup>	GATA3 NM_001002295.1 <sup>+</sup>	MDM4 NM_002393.4 <sup>+</sup>	PTPN11 NM_002834.3	U2AF1 NM_006758.2
CREBBP NM_004380.2	GATA4 NM_002052.3	MED12 NM_005120.2	PTPRD NM_002839.3	VEGFA NM_001025366.2
CRKL NM_005207.3	GATA6 NM_005257.4	MEF2B NM_001145785.1	PTPRS NM_002850.3	VHL NM_000551.3 <sup>+</sup>
CRLF2 NM_022148.2	GEN1 NM_182625.3	MEN1 NM_130799.2	PTPRT NM_133170.3	VTCN1 NM_024626.3
CSF1R NM_005211.3	GID4 NM_024052.4	MET NM_000245.2 <sup>+</sup>	QKI NM_006775.2	CCN6 NM_003880.3
CSF3R NM_156039.3	GLI1 NM_005269.2	MGA NM_001164273.1	RAB35 NM_006861.6	WT1 NM_024426.4
CSNK1A1 NM_001025105.2		MITF NM_000248.3	RAC1 NM_018890.3	XIAP NM_001167.3
CTCF NM_006565.3		MLH1 NM_000249.3	RAD21 NM_006265.2	XPO1 NM_003400.3
CTLA4 NM_005214.4		KMT2A NM_001197104.1	RAD50 NM_005732.3	XRCC2 NM_005431.1
		MLL T3 NM_004529.2	RAD51 NM_002875.4	YAP1 NM_001130145.2
		MPL NM_005373.2	RAD51B NM_133509.3	YES1 NM_005433.3
		MRE11 NM_005591.3	RAD51C NM_058216.2	ZBTB2 NM_020861.1
			RAD51D NM_002878.3	ZBTB7A NM_015898.2

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CTNNA1 NM_001903.2	GNA11 NM_002067.2	MSH2 NM_000251.2	RAD52 NM_134424.2	ZFX3 NM_006885.3
CTNNB1 NM_001904.3 <sup>+</sup>	GNA13 NM_006572.4	MSH3 NM_002439.4	RAD54L NM_001142548.1	ZNF217 NM_006526.2
CUL3 NM_003590.4	GNAQ NM_002072.3	MSH6 NM_000179.2	RAF1 NM_002880.3 <sup>+</sup>	ZNF703 NM_025069.1
CUX1 NM_181552.3	GNAS NM_000516.4 <sup>+</sup>	MST1 NM_020998.3	RANBP2 NM_006267.4	ZRSR2 NM_005089.3
CXCR4 NM_003467.2	ADGRA2 NM_032777.9	MST1R NM_002447.2	RARA NM_000964.3	MTAP NM_002451.3 <sup>++</sup>
CYLD NM_015247.2	GPS2 NM_004489.4	MTOR NM_004958.3	RASA1 NM_002890.2	
DAXX NM_001141970.1	GREM1 NM_013372.6	MUTYH NM_001128425.1	RB1 NM_000321.2 <sup>+</sup>	
DCUN1D1 NM_020640.2	GRIN2A NM_000833.3	MYB NM_001130173.1	RBM10 NM_005676.4	
DDR2 NM_001014796.1	GRM3 NM_000840.2		RECQL4 NM_004260.3	
DDX41 NM_016222.2	GSK3B NM_002093.3		REL NM_002908.2	
DHX15 NM_001358.2	H3-3A NM_002107.4 <sup>+</sup>		RET NM_020975.4 <sup>^+</sup>	
DICER1 NM_177438.2	H3-3B NM_005324.3			
DIS3 NM_014953.3	H3-5 NM_001013699.2			
	HGF NM_000601.4			
	H1-2 NM_005319.3			

<sup>^</sup>Summit™ also reports fusion events for this gene  
<sup>+</sup>Summit™ also reports copy number alterations for this gene  
<sup>\*</sup>Summit™ only reports copy number alterations for this gene

Aneuploidy (chromosome arm level loss and gain)									
chr1p	chr3p	chr5p	chr7p	chr9p	chr11p	chr13q	chr16q	chr18q	chr20q
chr1q	chr3q	chr5q	chr7q	chr9q	chr11q	chr14q	chr17p	chr19p	chr21q
chr2p	chr4p	chr6p	chr8p	chr10p	chr12p	chr15q	chr17q	chr19q	chr22q
chr2q	chr4q	chr6q	chr8q	chr10q	chr12q	chr16p	chr18p	chr20p	

### Methods and Limitations

The Summit™ 2.0 comprehensive genomic profiling (CGP) next-generation sequencing (NGS) test investigates tumor derived nucleic acid extracted from cerebrospinal fluid (CSF) for clinically relevant single/multi nucleotide variants (SNVs, MNVs), insertions and deletions (indels), gene level copy number variants (CNVs), and other biomarkers such as tumor mutation burden (TMB) and microsatellite instability (MSI). Methodology involves evaluation of 520 genes for SNVs, MNVs, Indels, 62 genes for CNVs, 28 genes for fusions, as well as TMB and MSI (PMID: 41595175). The LOD (limit of detection) for SNVs, MNVs and Indels was determined to be 0.3% variant allelic frequency (VAF), for CNVs was determined to be  $\geq 2$ -fold change for amplifications and  $\leq 0.5$ -fold change for deletions, and for fusions was determined to be  $\geq 2$  supporting reads. Reporting thresholds for TMB and MSI are:  $< 10$  Mut/Mb (TMB low),  $\geq 10$  Mut/Mb (TMB high), and when total unstable sites are  $< 30\%$  (MSS) and  $\geq 30\%$  (MSI-High). Ascent™ evaluates chromosomal arm level loss/gain (aneuploidy), and focal alterations (gene level amplification/deletion) using  $> 0.1x$  low pass whole genome sequencing (LP-WGS) (PMID: 37014860). The LOD (limit of detection) for aneuploidy was determined to be  $\log_2(r)$  of abs (0.09), and for focal alteration was determined to be seq.mean cutoff of 0.1 for amplification and -0.2 for deletions. Variants are called against the human genome build reference hg19 using Summit™Omics pipeline version 1.3.0, developed at Belay Diagnostics.

The Vantage™ MGMT Promoter Methylation Assay utilizes a quantitative PCR (qPCR) followed by high-resolution melt analysis (HRM) using the EpiMelt MGMT kit (MethylDetect) after enzymatic conversion (NEBNext Enzymatic Methyl-seq, New England Biolabs) on a portion of the library generated in the Summit™ workflow. Methylated and unmethylated melting temperature peaks are evaluated using the LightCycler® 480 Software v. 1.5.1 (Roche LifeScience). Qualitative results are reported as "Negative - Unmethylated", "Positive - Methylated", or "Indeterminate Results were equivocal". Specimens with results above the validated 25% methylated control are interpreted as "Positive". Specimens with results between unmethylated and methylated control are interpreted as "Indeterminate".

Tertiary analysis is performed using the precision oncology workbench (GenomOncology) based on the joint AMP/ASCO/CAP consensus guidelines for interpretation of sequence variants in cancer (PMID: 27993330). Please reach out to contact@belaydiagnostics.com for additional information or queries.

### Disclaimers

This test was developed, and its performance characteristics determined by Belay Diagnostics Laboratory (CLIA# 14D2302605), which is certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA) as qualified to perform high complexity testing. This test has not been cleared or approved by the U.S. Food and Drug Administration (FDA). This test may be used for clinical purposes. However, the results of this test do not establish a diagnosis and should not be used alone for diagnosis or patient care decisions or otherwise replace the judgment of a treating physician and must always be interpreted in the context of all relevant clinical and pathological data.

This test is performed only to evaluate for somatic (i.e., tumor-specific) variants within the genes listed and cannot distinguish between germline and somatic alterations with absolute certainty. This test therefore does not report on incidental findings as defined by the American College for Medical Genetics and Genomics (ACMG) (PMID: 37347242). If a germline variant is suspected, follow-up germline testing using non-neoplastic (normal) tissue should be performed by a laboratory permitted to perform germline genetic testing along with genetic counseling. It is possible for a genomic variant to be present yet go undetected by our assay either due to the heterogeneous nature of the specimen or the limits of detection of our assay. Therefore, to

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the extent a particular genomic variant is not reported, Belay Diagnostics LLC does not guarantee that the variant does not exist in the specimen provided. Likely benign, and benign variants are not reported. For any reported variant of uncertain significance (VUS), if the classification changes, there is no obligation to send out a new report updating this information.

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### ACTIONABILITY REFERENCES

**FDA:** U.S. Food & Drug Administration ([fda.gov](http://fda.gov))

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**WHO:** World Health Organization Classification of Tumours online ([tumourclassification.iarc.who.int](http://tumourclassification.iarc.who.int))

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