

International Test Requisition Form - Page 1 of 7

COMPLETE ENTIRE FORM IN ENGLISH AND SUBMIT PEDIGREE/CLINIC NOTES TO AVOID DELAYS

COLLECTION DATE (REQUIRED)

If date of collection is not provided, three calendar days before specimen receipt will be used (for

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please provide the total annual gross household income: s and the number of family members in the household supported by the listed income: I authorize Ambry Genetics Corporation to	more about Ambry's privacy practices at https://www.ambrygen.com/legal/notice-of-privacy-practices.						
verify the above information for the sole purpose of assessing financial need, including the right to seek supporting documentation.	please provide the total annual gross household income: \$and t	he number of family	members in the household	supported by the listed income:			
Patient Signature (I agree to terms above): Date:	Patient Signature (I agree to terms above):					Date:	
Patient Consent To Testing I acknowledge and agree that my health care provider has ordered genetic testing for processing at Ambry Genetics, a laboratory based in the United States and subject to U.S. based privacy laws, including the Health Insurance Portability and Accountability Act (HIPAA). I have provided my health care provider with the appropriate consent acknowledgment and documentation as required under the applicable laws in my country of residence for my genetic information to be sent to and processed in the United States. I have reviewed Ambry's Notice of Privacy Practices and understand and agree to the use of my health information in accordance with such privacy practices. I understand that I can exercise my rights to privacy consistent with HIPAA and Ambry's privacy notices.	I acknowledge and agree that my health care provider has ordered genetic t Health Insurance Portability and Accountability Act (HIPAA). I have provid country of residence for my genetic information to be sent to and processed	ed my health care produced in the United States	ovider with the appropriat . I have reviewed Ambry's	e consent acknowledgment and do Notice of Privacy Practices and un	cumentation as	required under the a	applicable laws in my
Patient Signature (I agree to terms above): Date:	Patient Signature (I agree to terms above):					Date:	



Patient Name: _

DOB: _

Comprehensive Test Requisition Form - Page 2 of 7

PLEASE SUBMIT THE FOLLOWING WITH THE TRF:

1. Clinic Notes 2. Pedigree 3. Insurance Card and Authorization Documents

PLEASE ATTACH PEDIGREE /CLINICAL CONSULTATION NOTES, IF AVAILABLE									
Birth and Neonatal History 🔲 Not Applicable					Developmental History Not Applicable				
Gestational age at bir	th		Birth we	aight.	Developmental delay: 🗌 yes 🗌 no 🗌 un	known			
Head circumference a					Delay prior to seizure onset: 🗌 yes 🗌 no	🗌 unknown 🔲 N/A			
					Type of delay (choose all that apply): 🗌 mo				
-	-				Intellectual disability: Uyes Ono Ounl				
Positive newborn s	creen, expla	in:			Regression or plateau: yes no ur – Does patient meet DSM-V diagnostic criteria				
Seizure History	lot Applicab	le			yes no unknown	in an autom speetram alsoraer			
Age at first unprovoke	ed seizure (f	rst se	izure without	fever or other acute	Cardiac History D Not Applicable				
metabolic or structura					Sudden cardiac arrest □Y □N (if yes): # E	pisodes:			
Seizure types (choose					Age first incident:	·			
☐ Infantile/epileptic s				Generalized tonic clonic	Syncope IY IN If yes, # Episodes:	Age first incident:			
			cal absence	Focal seizures	History of cardiomyopathy $\Box Y \Box N$ Age a	-			
Atonic									
Are seizures: Crefra					Cardiomyopathy type: History of Arrhythmia				
Has this patient been	diagnosed v	vith a	n epilepsy syr	ndrome?					
□ yes □ no □ unk	nown If ye	s, ple	ase specify: _		Arrhythmia type:				
Pulmonology History	🗌 Not Ap	plicat	ole		Congenital heart defect				
Positive newborn s	creen 🗌 C	BAVE	D 🗌 Mecon	ium ileus	-				
Infections:					Other History 🗌 Not Applicable				
Sweat chloride:	mmol/L	🗆 S	weat chloride	e: □<40 □ 40-60 □ >60	Hearing problems: I	□ Vision problems:			
Pancreatic insuffici	ency IRT le	vel: _			□ Migraine: □ F				
					Hematological:				
Respiratory assista	nce devices				□ Suspected genetic condition:				
Ultrasound finding					\Box Other clinical findings:				
	Cancer History 🗌 Not Applicable Metastatic: 🗌 Yes 🗌 No Tumor is 🗋 MSI-High or 🗋 IHC-Abnormal								
Cancer/Tumor	Age at Dx		thology and						
Brain									
Breast		Ту	ne:	FR](+) □(-) □unk PR□(+) □(-) □unk	HER2/neu□(+)□(-)□unk			
2nd primary breast		Ty			$ (+) \square (-) \square unk PR \square (+) \square (-) \square unk$	HER2/neu [(+) [(-)] unk			
Colorectal			Location:						
Ovarian		_	Fallopian tube Primary peritoneal						
Melanoma/skin									
Prostate Gleason Score:									
Uterine									
Hematologic* Type: Allogenic bone marrow or peripheral stem cell transplant*									
Other Cancer Type:									
GI polyps Adenomatous Polyp #: [] 1 [] 2-5 [] 6-9 [] 10-19 [] 20-99 [] 100+ Other type: Polyp #: [] 1 [] 2-5 [] 6-9 [] 10-19 [] 20-99 [] 100+									
*Blood or saliva from patients with active/recent hematological disease will undergo additional review and may not be accepted in some cases. For these, cultured fibroblasts or fresh/fresh frozen normal tis-									
sue are preferred. See ambrygen.com/specimen-requirements for details. PREVIOUS TEST HISTORY (Please include copy of test results if performed at another laboratory) Limited family history									
PREVIOUS TEST HI	STORY (Ple	ase inc	lude copy of test	results if performed at another labora	tory) 🗌 Limited family history				
Known Familial Variant:	Known Familial Variant: 🗌 Family 🗋 Self Gene: Variant (c. and/or p.): Testing Lab: Ambry ID:					Ambry ID:			
Patient previously tested at Ambry? Yes No Family previously tested at Ambry? Yes No									
Name: DOB:									
FAMILY MEMBER INFORMATION (Completion of this section is required for orders including parental samples)									
Mother - Name:				DOB: 🗆 un	affected 🗌 affected, list symptoms/dx:	Dx age:			
Father - Name:				DOB: 🗌 unat	ffected 🗌 affected, list symptoms/dx:	Dx age:			
Relationship to Patient	Mat	Pat	Age at Dx	Family Testing and Cancer Type D	Details	Reason relative has not been tested			
						Deceased Declines No Contact			
						Deceased Declines No Contact			
						Deceased Declines No Contact			



Patient Name: _

Comprehensive Test Requisition Form - Page 3 of 7

If this TRF is sent to Ambry without or ahead of the sample, it will be treated as a preverification. If test ordered is different than the test preverified, we will honor what is on the TRF order form with the sample. Preverification will only be performed for ExomeNext or SNP Array testing.

	Concurrent Testing: There is no action needed on your part if this is your desired strategy.								
Reflex Testing: Please select this option if you wish to have testing performed in a reflex manner, and indicate the order of testing below:									
Test 1: Test 2:									
See Refle	ex or Concu	rrent Testing sectio	on of the S	Supplemental Information page for more information	ition.				
CANCE	R TEST O	RDERS							
Primary	/ Test Ord	er							
l re	QUIRED: S	elect a Primary Te	st Order						
For Pati	ents Meet	ing BRCA1/2 Te	sting Cri	teria	For Patie	nts Mee	ting Colorectal Ca	ncer Syr	ndrome Testing Criteria (polyposis)
BRCA	1/2 test				Polyposis	s test: 🗌	APC/MUTYH		
For Pati	ents Meet	ing Colorectal C	ancer Sy	ndrome Testing Criteria (Lynch)	Other:				
Lynch Sy	ndrome te	est: 🗌 MLH1, MSH	2, MSH6,	PMS2, EPCAM	None o	of the abo	ve (patient does no	t meet a	ny genetic testing criteria)
Select a	n Optiona	al Supplemental	Test (Pe	r payer policy, all tests in this section will b	e process	ed and b	illed separately; te	ests may	be performed as a reflex.)
Order	Test Code	Test Name		Description	Order	Test Code	Test Name		Description
	8857	BRCANext®		19 gene breast & gynecologic cancer test		8821	ColoNext®		21 gene colorectal cancer & polyposis test
		Add on: 🗌 Limite	d Evidenc	e (Additional 7 genes)		1	Add on: 🗌 Limited	Evidence	(Additional 5 genes)
	8836	BRCAPlus®		13 gene STAT breast management test	-		CustomNext-Can	cer®	
	8824	CancerNext®		40 gene pan-cancer test			Notes:		up to 90 gene custom test
	8875	CancerNext-Exp		77 gene pan-cancer test		9511			Gene content is required. Use CustomNext- <i>Cancer</i> supplemental form for guidance.
				e (Additional 8 genes)	-				cuncer supplemental torm for guidance.
Others		Add on: Pancre		0					
		ntal Test Options	-	us, or STAT orders; PAXgene® tube required f					
	linsignt° (i		вксари	as, or STAT orders, FAXgene® tube required in					
Order	Test Nam	e	Test Code	Description	Order	Test Na	me	Test Code	Description
Breast and/or Ovarian Cancer					Gastroi	ntestinal	Cancer (Cont.)		
	ATM		9014	Ataxia-telangiectasia		MLH1		8508	Lynch syndrome
	BRCA1/2		8838			MSH2+	EPCAM del/dup	8510	Includes MSH2 inversion
		/2 Ashkenazi Jew- ite mutation panel 5892		Hereditary breast and ovarian cancer		MSH2 i	nversion	2226	Lynch syndrome
	CHEK2		9016			MSH6		8512	Lynch syndrome
	DICER1		5260			MUTYH	1	4661	MUTYH-associated polyposis
						PMS2		4646	Lynch syndrome
	PALB2	2366				STK11		2766	Peutz-Jeghers syndrome
	PTEN	N 2106		PTEN-related disorders (including Cowden syndrome)	Genito	burinary Cancer			
□ TP53 2866		2866	Li-Fraumeni syndrome		BAP1		9044		
Endocrine Tumors				FH		6301	Hereditary leiomyomatosis and renal cell cancer		
	MEN1 2646		2646	Multiple endocrine neoplasia type 1		FLCN		5921	Birt-Hogg-Dubé syndrome
	RET gene	sequence	2680	Multiple endocrine neoplasia type 2				2606	Von-Hippel Lindau disease
Gastroi	ntestinal (Cancer				TSC1 ar	nd TSC2	5904	Tuberous sclerosis complex
	APC		3040	Familial adenomatous polyposis	Skin Ca	Skin Cancer/Melanoma			
	concurrent		8726	Adenomatous polyposis		- CDKN2A and CDKA		4708	Familial atypical multiple mole melanoma (FAMMM)
	BMPR1A and SMAD4 8604		8604	Juvenile polyposis syndrome		PTCH1		5684	Gorlin syndrome
		Hereditary diffuse gastric cancer	Other H	Other Hereditary Cancer Testing					
	EPCAM d	el/dup	8519	Lynch syndrome		NF1		5704	Neurofibromatosis type 1
	Lynch syr	ndrome	8517	MLH1, MSH2, MSH6, PMS2 + EPCAM del/		NF2		9024	Neurofibromatosis type 2
	(concurre	ent)	0.517	dup		RB1		5426	Hereditary retinoblastoma
						SMARC	B1	7180	Schwannomatosis
					Other S	Single Syr	ndrome Orders		· · · · · · · · · · · · · · · · · · ·
						<u> </u>		/heredit	ary-cancer-single-gene-tests for details.

Test Code(s): _

_Gene/Test Name(s):



Comprehensive Test Requisition Form - Page 4 of 7

Order	Test Name	Test Code	Description	Order	Test Name	Test Code	Description		
CARDI	OLOGY								
Compre	hensive Cardiovascular Pa	inels		Familia	Hypercholesterolemia				
	CardioNext®	8911	92 genes for hereditary cardiomyopathies and arrhythmias		FHNext®	8680	4 genes (APOB, LDLR, LDLRAP1, PCSK9)		
	Custom Naut Cardia®	0520	Up to 167 genes related to hereditary cardiomyopathies, arrhythmias, TAAD,	Check this box if you would like to have the <i>SLC01B1</i> c.521T>C polymorphism reported with FHNext, which has been associated in medical literature with statin-induced myopathies					
	CustomNext-Cardio®	9520	HHT, Noonan, and lipidemias. Required: completed CustomNext-Cardio supplemental form. ambrygen.com/forms		FCSNext (Familial Chylo- micronemia Syndrome)	8920	APOA5, APOC2, GPIHBP1, LMF1, LPL		
Arrhyth	imia Panels				Sitosterolemia	8930	ABCG5, ABCG8		
	LongQTNext™	8890	17 genes for long QT, Brugada and short	Aneury	sms and Related Disorders	5			
	RhythmNext®	8900	QT syndromes 42 genes for long QT syndrome, Brugada and short QT syndromes, CPVT and ARVC		TAADNext®	8789	35 genes for thoracic aortic aneurysms/ dissections, Marfan syndrome, Ehlers-Danlos and related disorders		
	CPVTNext®	8902	4 genes for catecholaminergic polymorphic ventricular tachycardia		Marfan reflex to TAADNext	8783	FBN1 reflex to TAADNext		
Cardior	nyopathy Panels	1		Heredit	ary Hemorrhagic Telangied	ctasia (H	IHT)		
	HCMNext®	8936	30 genes for hypertrophic cardiomyopathy		HHTNext®	8672	ACVRL1, ENG, EPHB4, GDF2, RASA1, SMAD4		
	HCMNext Reflex	8883	MYBPC3, MYH7 reflex to HCMNext	Noonar	n Syndrome				
	DCMNext®	8884	37 genes for dilated cardiomyopathy		NoonanNext™	8402	18 genes for RASopathies		
	CMNext [®]	8887	56 genes for hereditary cardiomyopathy	Other	Hoondinitext	0402			
	ARVCNext™	8904	11 genes for arrhythmogenic right ventricular cardiomyopathy		Transthyretin amyloidosis	1560	TTR		
CLINIC	AL GENOMICS								
For Refl	ex or Concurrent Testing:								
		Poflov to	Test 2: [] Reflex to	Test 3:				
iest i		Concurr							
See Refl	ex or Concurrent Testing sec	ction of t	he Supplemental Information page.						
Chromo	osomal Microarray								
	SNP Array	5490	Chromosomal microarray (>2.6 million copy number probes and 750,000 SNP probes)		Familial targeted microarray	5495	Paid option. Only available following SNP Array (5490) completed at Ambry. Incidental findings unrelated to the variant(s) detected in the proband will NOT be reported. Name of proband tested at Ambry:		
Exome	Exome								
REQUIRED: Select a Primary Test Order									
	ExomeNext®-Proband	9993	Proband only exome sequencing		ExomeNext-Trio	9995	Trio exome sequencing		
	ExomeNext-Proband plus mtDNA	9994	Proband only exome sequencing plus mtDNA sequencing		ExomeNext- <i>Trio</i> plus mtDNA	9996	Trio exome sequencing plus mtDNA sequencing		
	ExomeNext-Duo	9991	Duo exome sequencing				Rapid Trio exome sequencing plus mtDNA		
	ExomeNext- <i>Duo</i> plus mtDNA	9992	Duo exome sequencing plus mtDNA sequencing		ExomeNext- <i>Rapid</i> ®	9999R	sequencing (Institutional billing or patient payment only)		
If ordering ExomeNext/ExomeNext-Rapid, please complete: Secondary Findings Report: Check below to opt-out of the ACMG Recommended List of secondary findings. If left unchecked, secondary findings will be reported. Opt-out: I choose to decline the ACMG Recommended List of secondary findings									
Exome	Next Supplemental Test Op	otions							
	ExomeReveal™	9990	RNA analysis available with all ExomeNext orders except for ExomeNext- <i>Rapid</i> , EDTA and PAXgene RNA tubes required						
ENDOC	RINOLOGY								
	Hereditary leiomyomatosis renal cell carcinoma	6301	FH		Multiple endocrine neoplasia type 2 and familial medullary thyroid	2680	<i>RET</i> gene sequence		
	Multiple endocrine neoplasia type l	2646	MEN1		cancer (FMTC) Neurofibromatosis type 1	5704	NF1		
	1				von-Hippel Lindau disease	2606	VHL		



Patient Name: ____

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GASTR	GASTROENTEROLOGY						
	CFTR gene sequence and deletion/duplication	1007	Report poly T/TG status		Juvenile polyposis syndrome	8604	BMPR1A, SMAD4
	analysis Hirschsprung disease				Pancreatitis	8022	CASR, CFTR, CPA1, CTRC, PRSS1, SPINK1
	(RET-related)	2680	<i>RET</i> gene sequence		Peutz-Jeghers syndrome	2766	STK11
HEMA	HEMATOLOGY/ONCOLOGY						
	Shwachman-Diamond syndrome	1440	SBDS				
NEURC	NEUROLOGY						
For patie you do n	ot check this box, VUS will Not ntal samples provided for co	r neurode OT be rep osegreg a	evelopmental disorder panel, checking this box in ported.				
For Refl	ex or Concurrent Testing:						
Test 1: _		Reflex to		Reflex to			
See Ref			he Supplemental Information page.] Concurr	ent with		
Order	Test Name	Test Code	Description	Order	Test Name	Test Code	Description
Epileps	y	1		Neurod	evelopmental Disorders		-
	EpilepsyNext®	6864	124 genes for epilepsy		AutismNext®	6863	72 genes for non-syndromic autism spectrum disorders and/or intellectual disability
	EpilepsyNext- <i>Expanded</i> ™	6865	>890 genes associated with seizures, primarily with neonatal to childhood onset		Autism, macrocephaly	2106	PTEN
Heredit	ary Neuropathy				Fragile X syndrome	4544	FMR1 repeat expansion analysis and
	Familial transthyretin amyloidosis	1560	TTR		NeurodevelopmentNext™	6861	methylation studies 202 genes known to cause developmental delay, intellectual disability and/or autism
							spectrum disorders
-	utaneous/Neuro-Oncolog	-					Ι
	Ataxia-telangiectasia	9014	ATM		Neurofibromatosis 2 Nevoid basal cell	9024	NF2
	HHTNext®	8672 5724	ACVRL1, ENG, EPHB4, GDF2, RASA1, SMAD4 SPRED1		carcinoma syndrome/ Gorlin syndrome	5684	РТСН1
	Legius syndrome	2866	TP53	Image: Commission of the synthesis of the synthesyntemes of the synthesis of the synthesis of the synt		TSC1, TSC2	
	Neurofibromatosis 1	5704	NF1	von Hippel-Lindau disease 2606 VHL			
PULMONOLOGY							
Congenital Central Hypoventilation Syndrome Primary Ciliary Dyskinesia							
	Congenital central hypoventilation syndrome	1580	PHOX2B gene sequence		PCDNext®	8122	21 genes for primary ciliary dyskinesia Report poly T/TG status
Cystic Fibrosis Pulmonary Fibrosis							
	CFTR gene sequence and deletion/duplication	1007	Report poly T/TG status		Telomere-related pulmonary fibrosis	8140	TERT, TERC
	analysis			Respira	tory Distress Syndrome	1	
					Surfactant dysfunction (respiratory distress syndrome)	8100	ABCA3, SFTPB, SFTPC gene sequence
VASCULAR							
	HHTNext®	8672	ACVRL1, ENG, EPHB4, GDF2, RASA1, SMAD4		TAADNext®	8789	35 genes for thoracic aortic aneurysms
	Marfan syndrome reflex to TAADNext	8783	FBN1 reflex to TAADNext				
SPECIFIC SITE ANALYSIS (Please include a copy of relative's report)							
Gene(s): Mutation(s): Relative Name:							
Relationship to Relative: Accession # (if tested at Ambry):							
Positive control sample: 🗌 will be provided 🔲 already at Ambry 🔲 not available							



Supplemental Information - Page 6 of 7

Hereditary Cancer Multi-Gene Tests

TEST NAME	TEST CODE	GENES					
Pan-cancer							
CancerNext® (40 genes)	8824	APC, ATM, AXIN2, BAP1, BARD1, BMPR1A, BRCA1, BRCA2, BRIP1, CDH1, CDKN2A, CHEK2, EPCAM, FH, FLCN, GREM1, HOXB13, MBD4, MET, MLH1, MSH2, MSH3, MSH6, MUTYH, NF1, NTHL1, PALB2, PMS2, POLD1, POLE, PTEN, RAD51C, RAD51D, RPS20, SMAD4, STK11, TP53, TSC1, TSC2, VHL					
CancerNext- <i>Expanded</i> ® (77 genes or up to 90 genes w/ add-ons)	8875	AIP, ALK, APC, ATM, AXIN2, BAP1, BARD1, BMPR1A, BRCA1, BRCA2, BRIP1, CDC73, CDH1, CDK4, CDKN1B, CDKN2A, CEBPA, CHEK2, CTNNA1, DDX41, DICER1, EGFR, EPCAM, ETV6, FH, FLCN, GATA2, GREM1, HOXB13, KIT, LZTR1, MAX, MBD4, MEN1, MET, MITF, MLH1, MSH2, MSH3, MSH6, MUTYH, NF1, NF2, NTHL1, PALB2, PDGFRA, PHOX2B, PMS2, POLD1, POLE, POT1, PRKAR1A, PTCH1, PTEN, RAD51C, RAD51D, RB1, RET, RPS20, RUNX1, SDHA, SDHAF2, SDHB, SDHC, SDHD, SMAD4, SMARCA4, SMARCB1, SMARCE1, STK11, SUFU, TMEM127, TP53, TSC1, TSC2, VHL, WT1					
		Optional Add-on 1 - Limited Evidence Genes (8 genes): ATRIP, EGLN1, KIF1B, MLH3, PALLD, RAD51B, RNF43, TERT					
		Optional Add-on 2 - Pancreatitis Genes (5 genes): CFTR, CPA1, CTRC, PRSS1, SPINK1					
STAT Breast Management							
BRCAPlus [®] (13 genes)	8836	ATM, BARD1, BRCA1, BRCA2, CDH1, CHEK2, NF1, PALB2, PTEN, RAD51C, RAD51D, STK11, TP53					
Breast & gynecologic	·						
BRCANext [®] (19 genes or up to 26	8857	ATM, BARD1, BRCA1, BRCA2, BRIP1, CDH1, CHEK2, EPCAM, MLH1, MSH2, MSH6, NF1, PALB2, PMS2, PTEN, RAD51C, RAD51D, STK11, TP53					
genes w/ add-on)		Optional Add-on - Limited Evidence Genes (7 genes): ATRIP, CDC73, FH, NTHL1, POLD1, POLE, RAD51B					
Colorectal & polyposis							
ColoNext® (21 genes or up to 26	8821	APC, AXIN2, BMPR1A, CDH1, EPCAM, GREM1, MBD4, MLH1, MSH2, MSH3, MSH6, MUTYH, NTHL1, PMS2, POLD1, POLE, PTEN, RPS20, SMAD4, STK11, TP53					
genes w/ add-on)		Optional Add-on - Limited Evidence Genes (5 genes): ATM, CHEK2, CTNNA1, MLH3, RNF43					
Customizable							
		To order all genes on Ambry's oncology menu, please order CancerNext-Expanded.					
CustomNext- <i>Cancer®</i> (up to 90 genes) Required: complete CustomNext- <i>Cancer</i> supplemental form. <u>ambrygen.com/forms</u>	9511	AIP, ALK, APC, ATM, ATRIP, AXIN2, BAP1, BARD1, BMPR1A, BRCA1, BRCA2, BRIP1, CDC73, CDH1, CDK4, CDKN1B, CDKN2A, CEBPA, CFTR, CHEK2, CPA1, CTNNA1, CTRC, DICER1, DDX41, EGFR, ELGN1, EPCAM, ETV6, FH, FLCN, GATA2, GREM1, HOXB13, KIF1B, KIT, LZTR1, MAX, MBD4, MEN1, MET, MITF, MLH1, MLH3, MSH2, MSH3, MSH6, MUTYH, NF1, NF2, NTHL1, PALB2, PALLD, PDGFRA, PHOX2B, PMS2, POLD1, POLE, POT1, PRKAR1A, PRSS1, PTCH1, PTEN, RAD51B, RAD51C, RAD51D, RB1, RET, RNF43, RPS20, RUNX1, SDHA, SDHAF2, SDHB, SDHC, SDHD, SMAD4, SMARCA4, SMARCB1, SMARCE1, SPINK1, STK11, SUFU, TERT, TMEM127, TPS3, TSC1, TSC2, VHL, WT1					
		For Medicare Patients: At a minimum, the following core genes must be included in the panel to ensure Medicare coverage: APC, ATM, BRCA1, BRCA2, CHEK2, EPCAM, MLH1, MSH2, MSH6, PALB2, PMS2, PTEN, TP53.					
Syndrome specific							
Adenomatous polyposis	8726	APC, MUTYH					
<i>BRCA1/2</i> -associated hereditary breast and ovarian cancer (HBOC)	8838	BRCA1, BRCA2					
Lynch syndrome	8517	MLH1, MSH2, MSH6, PMS2 + EPCAM del/dup					



Supplemental Information - Page 7 of 7

Specimen Requirements

Blood/saliva from patients with a history of allogenic bone marrow or stem cell transplant cannot be used for genetic testing. Blood/saliva from patients with active hematological disease is not recommended. An alternative specimen may be needed. Please see <u>ambrygen.com/specimen-requirements</u> for details.

Buccal swab sample available for chromosomal microarray (SNP array, familial targeted microarray), CustomNext-*Neuro®*, epilepsy, ExomeNext, fragile X syndrome, hereditary neuropathy (familial transthyretin amyloidosis), HHTNext, and neurodevelopmental disorder tests. Buccal swab samples from patients from patients with a history of allogenic bone marrow or stem cell transplant should not be used for genetic testing. For these patients, an alternative specimen (e.g. cultured fibroblasts) is required. Testing on buccal swab samples from patients with active hematological disease is not recommended. An alternative specimen (e.g. cultured fibroblasts) is recommended. Please see ambrygen.com/specimen-requirements for details.

Please note that Ambry cannot guarantee the viability of your specimen for testing at our laboratory, given the logistics of international specimen transfer. Testing may not be completed on specimens of inadequate quality due to specimen transfer issues not under Ambry's control, including, but not limited to, delays at customs, or other transfer-related delays. Ambry or your health care provider will reach out to you in such a case to rearrange for a specimen collection and transfer for completion of the ordered test.

Specific site analysis for variants identified at an external laboratory must be accompanied by a copy of the original testing report. A positive control from a known positive family member is recommended (required for prenatal testing).

Reflex or Concurrent Testing

Concurrent testing is when multiple tests are initiated at the same time. When multiple tests are ordered on the same test requisition form, testing will be run concurrently unless otherwise specified.

Reflex testing is when a subsequent test is initiated pending the outcome of the initial test. Reflex testing may result in delayed reporting of results.

For reflex test orders:

- Any diagnostic finding at any step will result in cancellation of any subsequent reflex tests.
- Non-diagnostic findings (including VUS or Uncertain results) will automatically reflex to the subsequent test.
- · Secondary findings results do not impact whether a subsequent test is initiated or canceled.

When ordering STAT panels (such as BRCAplus^{*}), the results of the STAT panel will be prioritized and reported with a shorter turnaround time, even if the tests were run concurrently.

Known Familial Variants

Variant-specific report comments about the presence or absence of known familial variant(s) require the "Known Familial Variant" section of this form to be completed accurately, including an internal Ambry reference ID and/or a copy of the positive family member's lab report. Acceptable types of Ambry identifiers include:

- Accession number
- Order number
- Name and date of birth

Variant requests without an internal Ambry reference ID or positive family member's lab report will not receive a variant-specific report comment.

Variant-specific report comments are not included in ExomeNext or Neurology panel reports.