

Comprehensive Test Requisition Form - Page 1 of 7

Ambry Genetic	S		COMPLETE ENTIRE FORM AND SUBMIT PEDIGREE/CLINIC NOTES TO AVOID DELAYS					
COLLECTION DATE (REQUI	RED)							
If date of collection is not provided, three cale specimens stored longer than 30 days, the da)					
2. PATIENT INFORMATION								
Legal Name (Last, First, MI)				Date of Birth (MM/DD/Y	 Sex Assigned at Birth □ F □ M 		der (optional) 1an □Woman elf-described	□ Nonbinary
Genetic Ancestry: □Ashkenazi Jewi □Middle Eastern □Native Americ					Mediterranean		MRN	
Address			City		St	tate	1	Zip
Mobile #		Email						1
SPECIMEN INFORMATION*	(Please see ambrygen.com/sp	ecimen-requireme	ents for details)					
Personal history of allogenic bone	marrow or peripheral stem ce	ell transplant						
Specimen ID			Medical Record #					
* Fetal specimens, cord blood and POC sample submission test codes.	will have maternal cell contami	nation studies add	led for a charge. Maternal a	nd fetal specimen require	d. Please see botton	n of page	e 5 for Maternal C	ell Contamination
Collection Assistance: Phlebotomy	_				,			<i>(</i> ,)
** As the patient's clinician, I am unawa patient if the safety of the phlebotomist	t and/or patient(s) are in questi		arawing blood for the liste	u patient(s). I understand	unat trie phleboton	nisť nas j	un authority to re	juse to araw any
INDICATION(S) FOR TESTIN	NG							
Will the medical management chang	re depending on the results of	the test? □Vec						
Was genetic counseling completed?			etic Counseling was Perfor	rmed:				
PRENATAL SAMPLES ONLY		Dute com		<u></u>				
Sample type: 🗌 Direct CVS 🔲	Cultured CVS 🔲 Cultured	amnio 🗌 POC	C Cultured POC	Gestational	age at sample coll	ection		
ORDERING LICENSED PROV	IDER/SENDING FACILI	TY (Each listed	person will receive a copy	of the report)				
Facility Name (Facility Code)	Address		City		ate /Country	Zip	Pho	ne
Ordering Licensed Provider Name (La	ast, First)(Code)	NPI#	Phone	Fax/Em	ail			
ADDITIONAL RESULTS RECI	PIENTS							
Genetic Counselor or Other Medical	Provider Name (Last, First) (Code)	Phone/Fax/Ema	ail				
Genetic Counselor or Other Medical	Provider Name (Last, First) (Code)	Phone/Fax/Ema	ail				
CONFIRMATION OF INFORMED The undersigned person (or represen consent. I confirm that testing is med genetic counseling services by a thirc applies to the attached letter of medi	ntative thereof) ensures he/sh lically necessary and that tesi d-party service, as required by	ie is a licensed me t results may imp	edical professional authori act medical management	zed to order genetic tes for the patient. I agree t	ting and confirms o allow Ambry Ge	that the netics to	facilitate the pr	ovision of pre-test
Signature Required for Processing	Medical Professional Sig	gnature:				0	Date:	
INSURANCE BILLING (Inclu		rance card)			ISTITUTIONA			
Patient Relation to Policy Holder? □Self □Spouse □Child	Name and DOB of Policy Holder (if not self)			Facili	ty Name	∐ Ser	nd invoice to facilit	ty address above
Insurance Company	Policy #		HMO Auth #	Addr	ess			
Special Billing Notes:				Cont	act Name			
				Phon	e Number		Email/Fax	
				□ P/	ATIENT PAYM	ENT	I	e to Ambry Genetics) all 949-900-5795)
Patient Acknowledgement: I acknowledg (Ambry), authorize <u>Ambry</u> to release mer medical records for this purpose. I unders I agree to be contacted regarding futur more about Ambry's privacy practices at For patient payment by credit card: I here	dical information concerning my stand that I am financially respon re research studies for which I ma https://www.ambrygen.com/leg	testing to my insur- sible for any amour ay be a candidate. A cal/notice-of-privac	er, to be my designated repre nts not covered by my insurer Any future research projects v y-practices.	sentative for purposes of a and responsible for sendii vill be subject to a separate	ppealing any denial ng Ambry money rec informed consent p	of benefi ceived fro process a	ts as needed and t om my health insur nd participation is	o request additional ance company. voluntary. Learn

please provide the total annual gross household income: verify the above information for the sole purpose of assessing financial need, including the right to seek supporting documentation.

For NY Residents:

By checking this box, I agree that Ambry Genetics will retain my sample for 6 months after the testing above has been completed. By not checking this box, I understand that under New York State law, Ambry Genetics must discard my sample after the longer of (a) testing completion and (b) 60 days after the Date of Collection above.



Patient Name: _

DOB: _

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PLEASE SUBMIT THE FOLLOWING WITH THE TRF:

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

CLINICAL HISTORY									
PLEASE ATTACH PEDIGREE /CLINICAL CONSULTATION NOTES, IF AVAILABLE									
Birth and Neonatal Hi	story 🗌 N	ot App	olicable		Developmental History 🗌 Not Applicable				
Gestational age at hir	th		Birth w	eight:	Developmental delay: 🗌 yes 🔲 no 🗌 un	iknown			
Head circumference a					Delay prior to seizure onset: 🗌 yes 🗌 no	🗌 unknown 🔲 N/A			
	-				Type of delay (choose all that apply): 🗌 mo				
_					Intellectual disability: 🗌 yes 🗌 no 🗋 un				
Positive newborn s	creen, explai	n:			Regression or plateau: yes no ur Does patient meet DSM-V diagnostic criteria				
Seizure History	lot Applicabl	le			yes no unknown				
Age at first unprovoke	ed seizure (fi	rst sei	zure without	fever or other acute	Cardiac History 🗌 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:			
Tonic			al absence	Focal seizures	History of cardiomyopathy \Box Y \Box N Age at dx:				
Atonic			cal absence						
Are seizures: 🗌 refra	,				Cardiomyopathy type:				
Has this patient been	diagnosed w	ith an	epilepsy syr	ndrome?	History of Arrhythmia $\Box Y \Box N$ Age at d				
□ yes □ no □ unki	nown If ye	s, plea	se specify: _		Arrhythmia type:				
Pulmonology History	🗌 Not App	plicabl	е		Congenital heart defect				
Positive newborn s	creen 🗌 Cl	BAVD	☐ Mecon	ium ileus	-				
Infections:					Other History 🗌 Not Applicable				
Sweat chloride:	mmol/L	🗆 Sv	weat chloride	e: □<40 □ 40-60 □ >60	Hearing problems:	□ Vision problems:			
Pancreatic insuffici	ency IRT lev	vel:			☐ Migraine: ☐ Psychiatric:				
Respiratory distres	s, explain:				□ Hematological:				
Respiratory assista	nce devices:				□ Suspected genetic condition:				
Ultrasound finding	s:				□ Other clinical findings:				
Cancer History 🗌 Not Applicable Metastatic: 🗌 Yes 🗌 No Tumor is 🗌 MSI-High or 🗌 IHC-Abnormal									
Cancer/Tumor	Age at Dx	Pat	hology and	Other Info					
Brain									
Breast		Тур	e:	ER](+) □(-) □unk PR□(+) □(-) □unk	HER2/neu□(+) □(-) □unk			
2nd primary breast		Тур	e:	ER](+) □(-) □unk PR□(+) □(-) □unk	HER2/neu□(+)□(-)□unk			
Colorectal		Loc	ation:						
Ovarian		I I F	Fallopian tub	e 🔲 Primary peritoneal					
Melanoma/skin									
Prostate		Gle	ason Score:						
Uterine									
Hematologic*		Тур	e:	All	ogenic bone marrow or peripheral stem cell tra	ansplant*			
Other Cancer		Тур	e:						
GI polyps			Adenomatou Other type:	S	Polyp #: □1 □2-5 □6-9 □10-19 □2 Polyp #: □1 □2-5 □6-9 □10-19 □20	20-99 □100+ 0-99 □100+			
		/recent	hematological		and may not be accepted in some cases. For these, culture				
sue are preferred. See amb									
			12	t results if performed at another labora					
Known Familial Variar	nt: 🗆 Family	🗆 Sel	lf Gene:	Variant (c. and/or p.):	Testing Lab:	Ambry ID:			
				Family previously tested at Amb					
					DOB: Relation:				
				f this section is required for orders incl					
				· · · · · · · · · · · · · · · · · · ·		Πν ασει			
Father - Name:	Mother - Name: DOB: unaffected [] affected, list symptoms/dx: Dx age: Father - Name: DOB: [] unaffected [] affected, list symptoms/dx: Dx age:								
Relationship to Patient Mat Pat Age at Dx Family Testing and Cancer Type Details Reason relative has not been tested									
isolationship to ratient			Age at DA	ranny resting and cancer type L					
						Deceased Declines No Contact			
						Deceased Declines No Contact			
						Deceased Declines No Contact			



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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 Reflex or Concurrent Testing section of the Supplemental Information page for more information.										
CANCE	CANCER TEST ORDERS									
Primary	Primary Test Order									
R	EQUIRED: S	Select a Primary Te	st Order							
For Pati	ents Mee	ting BRCA1/2 Te	sting Cri	iteria	For Patients Meeting Colorectal Cancer Syndrome Testing Criteria (polyposis)					
□ BRCA1/2 test						s test: 🗌	APC/MUTYH			
For Pati	ents Mee	ting Colorectal C	ancer Sy	ndrome Testing Criteria (Lynch)	□ Other:					
		est: 🗆 MLH1, MSH		,					ny genetic testing criteria)	
Select a	1 -	al Supplemental	Test (Pe	r payer policy, all tests in this section will	be process	1	oilled separately; te	ests may	v 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	
<u> </u>			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®	andad®	40 gene pan-cancer test		0511	Notes:		up to 90 gene custom test	
	8875	CancerNext-Exp		77 gene pan-cancer test e (Additional 8 genes)		9511			Gene content is required. Use CustomNext- Cancer supplemental form for guidance.	
		Add on: Pancre			-					
Other S	uppleme	I		<u>_</u>						
□+RNA	Other Supplemental Test Options (Select if applicable) □ +RNAinsight® (Not available with BRCAplus, or STAT orders; PAXgene® tube required for RNA)									
Order	Test Nam		Test Code	Description	Order	Test Na		Test Code	Description	
		arian Cancer	1	Γ		1	l Cancer (Cont.)		I	
	ATM		9014	Ataxia-telangiectasia		MLH1		8508	Lynch syndrome	
	BRCA1/2		8838	Hereditary breast and ovarian cancer			EPCAM del/dup	8510	Includes MSH2 inversion	
		Ashkenazi Jew- mutation panel	5892				nversion	2226	Lynch syndrome	
	CHEK2		9016			MSH6		8512	Lynch syndrome	
	DICER1		5260			MUTYH	1	4661	MUTYH-associated polyposis	
	PALB2		2366			PMS2		4646	Lynch syndrome	
	DTEN		2107	PTEN-related disorders		STK11		2766	Peutz-Jeghers syndrome	
	PTEN	2106		(including Cowden syndrome)	Genito	ourinary Cancer		1	1	
	TP53		2866	Li-Fraumeni syndrome		BAP1		9044		
Endocri	ne Tumor	S		1		FH		6301	Hereditary leiomyomatosis and renal cell cancer	
	MEN1		2646	Multiple endocrine neoplasia type 1		FLCN		5921	Birt-Hogg-Dubé syndrome	
	RET gene	sequence	2680	Multiple endocrine neoplasia type 2		VHL		2606	Von-Hippel Lindau disease	
Gastroi	ntestinal (Cancer				TSC1 ar	nd TSC2	5904	Tuberous sclerosis complex	
	APC		3040	Familial adenomatous polyposis	Skin Ca	ancer/Me	elanoma		1	
	APC and concurre	nt	8726	Adenomatous polyposis		CDKN2 concurr	A and CDK4 rent	4708	Familial atypical multiple mole melanoma (FAMMM)	
	BMPR1A concurre	and SMAD4 nt	8604	Juvenile polyposis syndrome		PTCH1		5684	Gorlin syndrome	
	CDH1		4726	Hereditary diffuse gastric cancer	Other I	Hereditar	ry Cancer Testing			
	EPCAM c	lel/dup	8519	Lynch syndrome		NF1		5704	Neurofibromatosis type 1	
	Lynch sy		8517	MLH1, MSH2, MSH6, PMS2 + EPCAM del/		NF2		9024	Neurofibromatosis type 2	
	(concurr	ent)	0.517	dup		RB1		5426	Hereditary retinoblastoma	
						SMARC	CB1	7180	Schwannomatosis	
					Other S	Single Sy	ndrome Orders		·	
						Please	visit ambrygen.com	/heredit	ary-cancer-single-gene-tests for details.	

Test Code(s):

Gene/Test Name(s):



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Order	Test Name	Test Code	Description	Order	Test Name	Test Code	Description	
CARDI	OLOGY							
Compre	ehensive Cardiovascular Pa	inels		Familial	Hypercholesterolemia			
	CardioNext®	8911	92 genes for hereditary cardiomyopathies and arrhythmias		FHNext®	8680	4 genes (APOB, LDLR, LDLRAP1, PCSK9)	
			Up to 167 genes related to hereditary cardiomyopathies, arrhythmias, TAAD,				LC01B1 c.521T>C polymorphism reported with iterature with statin-induced myopathies	
	CustomNext-Cardio®	9520	HHT, Noonan, and lipidemias. Required: completed CustomNext- <i>Cardio</i> supplemental form. ambrygen.com/forms		FCSNext (Familial Chylo- micronemia Syndrome)	8920	APOA5, APOC2, GPIHBP1, LMF1, LPL	
Arrhyth	Imia Panels	<u> </u>	supplementarionn. and ygen.com/ torns		Sitosterolemia	8930	ABCG5, ABCG8	
	LongQTNext™	8890	17 genes for long QT, Brugada and short	Aneury	sms and Related Disorders	;		
	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		Marfan reflex to TAADNext	0702		
		0902	polymorphic ventricular tachycardia			8783	FBN1 reflex to TAADNext	
	nyopathy Panels	1			ary Hemorrhagic Telangied	1		
	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	1	1		
	ARVCNext™	8904	11 genes for arrhythmogenic right ventricular cardiomyopathy		Transthyretin amyloidosis	1560	TTR	
CLINIC	AL GENOMICS							
	ex or Concurrent Testing:							
	0							
Test 1: _				☐ Reflex to				
	Concurrent with							
	See Reflex or Concurrent Testing section of the Supplemental Information page.							
Chromo	osomal Microarray	1				1		
	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	•							
R	REQUIRED: Select a Primary Test Order							
	ExomeNext®-Proband	9993	Proband only exome sequencing		ExomeNext- <i>Trio</i>	9995	Trio exome sequencing	
	ExomeNext-Proband plus mtDNA	9994	Proband only exome sequencing plus mtDNA sequencing		ExomeNext- <i>Trio</i> plus	9996	Trio exome sequencing plus mtDNA	
	ExomeNext-Duo	9991	Duo exome sequencing		mtDNA		sequencing	
	ExomeNext <i>-Duo</i> plus mtDNA	9992	Duo exome sequencing plus mtDNA sequencing		ExomeNext- <i>Rapid</i> ®	9999R	Rapid Trio exome sequencing plus mtDNA sequencing (Institutional billing or patient payment only)	
If orderin	ng ExomeNext/ExomeNext-Ra	pid, pleas	e complete:		1			
Seconda	ary Findings Report: Check b	pelow to	opt-out of the ACMG Recommended List of s commended List of secondary findings	econdary f	indings. If left unchecked, se	condary	findings will be reported.	
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					
ENDO	RINOLOGY							
	Hereditary				Multiple endocrine			
	leiomyomatosis renal cell carcinoma	6301	FH		neoplasia type 2 and familial medullary thyroid cancer (FMTC)	2680	RET gene sequence	
	Multiple endocrine neoplasia type l	2646	MEN1		Neurofibromatosis type 1	5704	NF1	
					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				Pancreatitis	8022	CASR, CFTR, CPA1, CTRC, PRSS1, SPINK1	
	Hirschsprung disease (<i>RET</i> -related)	2680	RET gene sequence		Peutz-Jeghers syndrome	2766	STK11	
HEMA	TOLOGY/ONCOLOGY							
	Shwachman-Diamond syndrome	1440	SBDS					
NEURO	DLOGY		· · · · · · · · · · · · · · · · · · ·					
For patie you do r Paren Cosegre	 Opt-in to Reporting of Variants of Unknown Significance (VUS) For patients undergoing an epilepsy or neurodevelopmental disorder panel, checking this box indicates that VUS identified on the test(s) ordered below will be reported for this patient. If you do not check this box, VUS will NOT be reported. Parental samples provided for cosegregation Cosegregation testing of family members is available for the following panels: EpilepsyNext, EpilepsyNext-Expanded, AutismNext, NeurodevelopmentNext 							
	ex or Concurrent Testing:							
Test 1: _		Reflex to] Reflex to] Concurr				
See Ref			ne Supplemental Information page.					
Order	Test Name	Test	Description	Order	Test Name	Test	Description	
Epileps		Code			levelopmental Disorders	Code		
	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 methylation studies	
	Familial transthyretin amyloidosis	1560	TTR		NeurodevelopmentNext™	6861	202 genes known to cause developmental delay, intellectual disability and/or autism spectrum disorders	
Neuroc	Neurocutaneous/Neuro-Oncology Disorders							
	Ataxia-telangiectasia	9014	ATM		Neurofibromatosis 2	9024	NF2	
	HHTNext®	8672	ACVRL1, ENG, EPHB4, GDF2, RASA1, SMAD4		Nevoid basal cell carcinoma syndrome/	5684	РТСН1	
	Legius syndrome	5724	SPRED1		Gorlin syndrome Tuberous sclerosis complex	5904	TSC1, TSC2	
	Li-Fraumeni syndrome Neurofibromatosis 1	2866 5704	TP53 NF1		von Hippel-Lindau disease		VHL	
		5704						
	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 I	ibrosis			Pulmonary Fibrosis				
	CFTR gene sequence and deletion/duplication analysis	1007	Report poly T/TG status		Telomere-related pulmonary fibrosis	8140	TERT, TERC	
	allalysis			Respiratory Distress Syndrome				
					Surfactant dysfunction (respiratory distress syndrome)	8100	ABCA3, SFTPB, SFTPC gene sequence	
VASCU						1		
	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:							
Positive control sample: 🗌 will be provided 📄 already at Ambry 📄 not available								
FOR PRENATAL SPECIMENS, POC OR CORD BLOOD: MATERNAL CELL CONTAMINATION ANALYSIS REQUIRED Both maternal and fetal specimens are required.								
1260 MCC for fetal specimen or cord blood 1262 MCC Reference for maternal blood sample (No Charge)								



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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 genes w/ add-on)	8821	APC, AXIN2, BMPR1A, CDH1, EPCAM, GREM1, MBD4, MLH1, MSH2, MSH3, MSH6, MUTYH, NTHL1, PMS2, POLD1, POLE, PTEN, RPS20, SMAD4, STK11, TP53
		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



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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 ambrygen.com/specimen-requirements for details.

Fetal specimens, cord blood and POC will have maternal cell contamination studies added for a charge. Maternal and fetal specimen required. Please see bottom of page 5 for Maternal Cell Contamination sample submission test codes.

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.

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.