

COMPLETE ENTIRE FORM TO AVOID DELAYS

PATIENT INFORMATION					
Name (Last, First, MI)		Date of Birth (MM/DD/YY)	Date of Death (if applicable)	Phone Number/Email	
Address	City	State	Zip	Biological Sex <input type="checkbox"/> F <input type="checkbox"/> M	Ethnicity: <input type="checkbox"/> African American <input type="checkbox"/> Asian <input type="checkbox"/> Caucasian <input type="checkbox"/> Hispanic <input type="checkbox"/> Jewish (Ashkenazi) <input type="checkbox"/> Portuguese <input type="checkbox"/> Other:
SPECIMEN INFORMATION* (For phlebotomy service, select all services you are requesting)					
Type(s) <input type="checkbox"/> Blood (EDTA preferred) <input type="checkbox"/> Saliva <input type="checkbox"/> DNA <input type="checkbox"/> Other:			<input type="checkbox"/> Personal history of allogenic bone marrow or peripheral stem cell transplant		
Collection Date	Specimen ID		Medical Record #		
*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 tissue are preferred. See ambrygen.com/specimen-requirements for details.					
Phlebotomy Services Request: <input type="checkbox"/> Phlebotomy draw <input type="checkbox"/> Insurance preverification first <input type="checkbox"/> Send kit to patient ^As the patient's clinician, I am unaware of any potential for complication or difficulty in drawing blood for the listed patient(s). I understand that the phlebotomist has full authority to refuse to draw any patient if the safety of the phlebotomist and/or patient(s) are in question.					
ORDERING PHYSICIAN/SENDING FACILITY (Each listed person will receive a copy of the report)					
Facility Name (Facility Code)		Address	City	State /Country	Zip Phone
Ordering Licensed Provider Name (Last, First)(Code)		NPI#	Phone	Fax/Email	
ADDITIONAL RESULTS RECIPIENTS					
Genetic Counselor or Other Medical Provider Name (Last, First) (Code)			Phone/Fax/Email		
CONFIRMATION OF INFORMED CONSENT AND MEDICAL NECESSITY FOR GENETIC TESTING					
The undersigned person (or representative thereof) ensures he/she is a licensed medical professional authorized to order genetic testing and confirms that the patient has given appropriate consent. I confirm that testing is medically necessary and that test results may impact medical management for the patient. Furthermore, all information on this TRF is true to the best of my knowledge. My signature applies to the attached letter of medical necessity (unless this box is checked <input type="checkbox"/>).					
Signature Required for Processing Medical Professional Signature:				Date:	
INSURANCE BILLING (Include copy of both sides of insurance card)			INSTITUTIONAL BILLING		
Patient Relation to Policy Holder? <input type="checkbox"/> Self <input type="checkbox"/> Spouse <input type="checkbox"/> Child	Name and DOB of Policy Holder (if not self)		Facility Name <input type="checkbox"/> Send invoice to facility address above		
Insurance Company	Policy #	HMO Auth #	Address		
Ambry Genetics preverifies insurance coverage and will contact the patient after the patient's sample is received if the out-of-pocket amount for testing is estimated to exceed (Nothing checked defaults to >\$100): <input type="checkbox"/> \$100 <input type="checkbox"/> Any amount <input type="checkbox"/> Other \$			Contact Name		
<input type="checkbox"/> Hold order pending patient contact and approval of payment terms regarding out-of-pocket. Patient preferred method of contact regarding out-of-pocket amount: <input type="checkbox"/> Email <input type="checkbox"/> Phone			Phone Number		E-mail/Fax
			<input type="checkbox"/> PATIENT PAYMENT		<input type="checkbox"/> Check (Payable to Ambry Genetics) <input type="checkbox"/> Credit Card (Call 949-900-5795)
Patient Acknowledgement: I acknowledge that the information provided by me is true and correct. For direct insurance billing: I authorize my insurance benefits to be paid directly to Ambry Genetics Corporation (Ambry), authorize Ambry to release medical information concerning my testing to my insurer, to be my designated representative for purposes of appealing any denial of benefits as needed and to request additional medical records for this purpose. I understand that I am financially responsible for any amounts not covered by my insurer and responsible for sending Ambry money received from my health insurance company. For patient payment by credit card: I hereby authorize Ambry Genetics Corporation to bill my credit card as indicated above. In order to expedite consideration for eligibility for Ambry's E.P.I.C. Program, please provide the total annual gross household income: \$_____ and the number of family members in the household supported by the listed income: _____. I authorize Ambry Genetics Corporation to verify the above information for the sole purpose of assessing financial need, including the right to seek supporting documentation.					
FOR NY RESIDENTS:					
<input type="checkbox"/> I am a New York resident and I give Ambry Genetics permission to store my sample for longer than 60 days. NOTE: If left blank, consent is interpreted as "NO".					
Signature Required For Insurance/Self-Pay Patients and NY Sample Storage Consent:				Date:	

REQUIRED ORDERING CHECKLIST

- Clinic notes (with pedigree if available)
- ICD-10 code(s)
- Clinician & patient signatures
- Insurer-specific forms (i.e. ABN), if applicable
- Front/back copy of insurance card(s)

Cardiovascular Test Requisition Form - Page 3 of 3

Please check the box next to the test(s) being ordered below. All tests include gene sequence and deletion/duplication analyses unless otherwise indicated. If this TRF is sent to Ambyr 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.

For multiple test orders, testing will be run concurrently (multiple tests initiated at the same time) unless otherwise specified. To order reflexive testing (second test starts pending first test outcome), please clearly indicate the order of reflexive tests in the notes section or next to the test check box. For reflex test orders, any positive findings (pathogenic/likely pathogenic) in the first test will be reported out to the clinician, and the requested second test will be canceled; all other findings will automatically reflex (including VUS).

Check to order	Test Name	Test Code	Description
Cardiomyopathy Panels			
<input type="checkbox"/>	HCMFirst	8935	First tier test of 2 most common genes for hypertrophic cardiomyopathy (MYBPC3, MYH7)
<input type="checkbox"/>	HCMNext	8936	27 genes for hypertrophic cardiomyopathy
<input type="checkbox"/>	HCMNext Reflex	8883	HCMFirst reflex to HCMNext
<input type="checkbox"/>	DCMNext	8884	36 genes for dilated cardiomyopathy
<input type="checkbox"/>	CMNext without TTN	8886	54 genes for hereditary cardiomyopathy
<input type="checkbox"/>	CMNext with TTN	8887	55 genes for hereditary cardiomyopathy
<input type="checkbox"/>	ARVDNext	8904	9 genes for arrhythmogenic right ventricular dysplasia
<input type="checkbox"/>	LVNCNext	8906	8 genes for left ventricular non-compaction
Comprehensive Cardiovascular Panels			
<input type="checkbox"/>	CardioNext without TTN	8910	84 genes for hereditary cardiomyopathies and arrhythmias
<input type="checkbox"/>	CardioNext with TTN	8911	85 genes for hereditary cardiomyopathies and arrhythmias
<input type="checkbox"/>	CustomNext-Cardio	9520	Up to 106 gene custom hereditary cardiomyopathies and arrhythmias test*
Arrhythmia, Long QT, and Brugada Panels			
<input type="checkbox"/>	RhythmFirst	8888	12 genes for long QT, Brugada, and short QT syndromes
<input type="checkbox"/>	RhythmNext	8900	36 genes for long QT syndrome, Brugada syndrome, and other inherited arrhythmias
<input type="checkbox"/>	RhythmNext Reflex	8901	RhythmFirst with reflex to RhythmNext
<input type="checkbox"/>	CPVTNext	8902	6 genes for catecholaminergic polymorphic ventricular tachycardia
Aneurysms and Related Disorders			
<input type="checkbox"/>	TAADNext	8789	22 genes for thoracic aortic aneurysms/dissections
<input type="checkbox"/>	Marfan syndrome	8781	FBN1
<input type="checkbox"/>	Marfan reflex to TAADNext	8783	FBN1 reflex to TAADNext
<input type="checkbox"/>	Ehlers-Danlos vascular type (EDS IV)	8790	COL3A1
<input type="checkbox"/>	Ehlers-Danlos reflex to TAADNext	8791	COL3A1 reflex to TAADNext
*Required: completed CustomNext-Cardio supplemental form. ambrygen.com/forms			
SINGLE SITE ANALYSIS (Please include a copy of relative's report)			
Gene(s): _____ Mutation(s): _____			
Relative Name: _____			
Relationship to Relative: _____			
Accession # (If tested at Ambyr): _____			
Positive control sample: <input type="checkbox"/> will be provided <input type="checkbox"/> already at Ambyr <input type="checkbox"/> not available			

Check to order	Test Name	Test Code	Description
Familial Hypercholesterolemia			
<input type="checkbox"/>	FHNNext	8680	4 genes (APOB, LDLR, LDLRAP1, PCSK9)
<input type="checkbox"/> Check this box if you would like to have the SLC01B1 c.521T>C polymorphism reported, which has been associated in medical literature with statin-induced myopathies			
Hereditary Hemorrhagic Telangiectasia (HHT)			
<input type="checkbox"/>	HHTFirst	8673	First tier test of 3 most common genes for HHT (ACVRL1, ENG, SMAD4)
<input type="checkbox"/>	HHTNext	8672	5 genes for HHT
<input type="checkbox"/>	HHTReflex	8671	HHTFirst reflex to HHTNext
<input type="checkbox"/>	GDF2 and RASA1	8674	GDF2, RASA1
Noonan and Related Syndromes			
<input type="checkbox"/>	Noonan syndrome	8402	PTPN11, SOS1, KRAS, RAF1
<input type="checkbox"/>	PTPN11 - Noonan	2280	
<input type="checkbox"/>	SOS1 - Noonan	2300	
<input type="checkbox"/>	RAF1 - Noonan	2320	
<input type="checkbox"/>	KRAS - Noonan	2340	
Other Cardiovascular Genetic Tests			
<input type="checkbox"/>	CHARGE syndrome	2380	CHD7 gene sequence
<input type="checkbox"/>	Alagille syndrome	1640	JAG1
<input type="checkbox"/>	Transthyretin amyloidosis	1560	TTR
<input type="checkbox"/>	PCDNext	8122	21 genes for primary ciliary dyskinesia
Clinical Genomics			
<input type="checkbox"/>	Karyotype	3660	Chromosome analysis (requires green-top sodium-heparin tube)
<input type="checkbox"/>	Karyotype, rule out mosaic	3662	Chromosome analysis (requires green-top sodium-heparin tube)
<input type="checkbox"/>	SNP Array	5490	Chromosomal microarray (>2.6 million copy number probes and 750,000 SNP probes)
<input type="checkbox"/>	Follow-up parental FISH studies - ONLY available following SNP Array (5490) completed at Ambyr	3750	Sodium heparin tube, submit proband sample for positive control. Name of proband tested at Ambyr: _____
<input type="checkbox"/>	ExomeNext	9999	<input type="checkbox"/> Opt-out of analysis and reporting of Novel Genetic Etiologies
<input type="checkbox"/>	ExomeNext-Rapid	9999R	<input type="checkbox"/> Opt-out of analysis and reporting of Novel Genetic Etiologies
<input type="checkbox"/>	ExomeNext-Select	9500	Up to 500 gene custom exome sequencing test
*AP2 is AmbyrPort 2.0, our online portal ambrygen.com/ap2			
If ordering ExomeNext/ExomeNext-Rapid, please complete: Secondary Findings Report: Check below to order the ACMG Recommended List of secondary findings. If neither box is checked secondary findings will not be reported. Secondary findings results are issued in a separate report. (For expanded secondary findings options and pricing please complete the "ExomeNext Expanded Secondary Findings Request Form" and submit with sample). <input type="checkbox"/> Yes: I choose to receive the ACMG Recommended List of secondary findings <input type="checkbox"/> No: I choose to decline the ACMG Recommended List of secondary findings			
OTHER ORDER			
Please visit ambrygen.com/tests for details.			
Test Code: _____ Test Name: _____			