

# Pulmonology Test Requisition Form - Page 1 of 3 **COMPLETE ENTIRE FORM AND SUBMIT PEDIGREE/CLINIC NOTES TO AVOID DELAYS** To submit an order via email, please send the completed test requisition form to info@ambrygen.com

PATIENT INFORMATION										
Legal Name (Last, First, MI)					Sex Assigned	Gender (optional)		D	Date of Birth (MM/DD/YY)	
					at Birth	Man Woman	🗌 Nonb	inary		
					DF DM	Self-described	_			
Genetic Ancestry: Ashkenazi Jewish Asian Black/African American French Canadian/Cajun Hispanic/Latino Mediterranean										
	an Pacific Islander Port	uguese 🗌 Wi		vn 🗌 Other:			<b>CL</b> 1		-7.	
Address			City				State		Zip	
Phone		Email					1	ed Billing	Self-pay 🔲 Institutional	
*										
SPECIMEN INFORMATION*			nents for details	)						
Personal history of allogenic bone marrow or peripheral stem cell transplant										
Collection Date (Required) If date of co	ollection is not provided, three calen eceipt will be used (for specimens s	dar days before tored longer than :	30 Specimen	ID Medical Record #						
Collection Date (Required) If date of collection is not provided, three calendar days before specimen receipt will be used (for specimens stored longer than 30 days, the day of archive retrieval will be used as the date of service)										
* Fetal specimens, cord blood and POC submission test codes.	will have maternal cell contamir	ation studies ad	lded for a charge	e. Maternal and	fetal specimen re	equired. Please see page	e 2 for Ma	ternal Cell	Contamination sample	
				(:(:(:(:(:(:	allahla fan Evran					
Collection Assistance: Phlebotomy								full authori	ity to refuse to draw any	
patient if the safety of the phlebotomist	and/or patient(s) are in question	n.				, 	,		.,	
INDICATION(S) FOR TESTI	NG									
ICD-10 code(s):										
Will medical management change de	pending upon the results of th	ne test? 🗌 Yes	□No	STAT TE	ST: Date results	needed (if known):				
PATIENT HISTORY No per	sonal history of pulmonology	disease		1						
PLEASE SUPPLY CLINIC NOTES AND				Upcoming pro	ocedure date:					
Reason(s) for Testing										
				Relevant Jah	results (include	copies if possible)				
Positive newborn screen				Relevant lab		copies il possible)				
Infections:										
Sweat chloride:mmol/L										
□ Sweat chloride: □<40 □ 40-6 □ Congenital bilateral absence of the v				FAMILY HISTORY						
	as deferens									
Pancreatic insufficiency IRT level:				□ None (maternal) □ Maternal hx unknown □ None (paternal) □ Paternal hx unknown						
Respiratory distress, explain:				Relationship to patient 🗌 Maternal 🗌 Paternal						
Respiratory assistance devices:										
Ultrasound findings:				Relationship to patient 🗌 Maternal 🗌 Paternal						
Other:				Diagnosis Dx age						
ORDERING PHYSICIAN/SEN		to din over on will		of the report)						
Facility Name (Facility Code)	Address	teu person wii	тесете а сору	City		State /Country	Zip		Phone	
	Address			City		State / Country	Σip		THORE	
Ordering Licensed Provider Name (La	ast, First)(Code)	NPI#	F	hone	Fa	x/Email				
Genetic Counselor or Other Medical Provider Name (Last, First) (Code) Phone/Fax/Email										
CONFIRMATION OF INFORMEI						ns that the natient has give	en annronr	iate consen	t. I confirm that testing is	
The undersigned person (or representative medically necessary and that test results m the patient's insurance provider. Furthermo	ay impact medical management for	r the patient. I ag	ree to allow Amb	ry Genetics to faci	ilitate the provision	n of pre-test genetic coun	seling servi	ces by a thi	rd-party service, as required by	
Signature Required for Processing			ly knowledge. Wy	signature applies		ater of medical necessity.		Date:		
■ INSURANCE BILLING (Inclu										
Patient Relation to Policy Holder?	Name and DOB of	ance card)				Facility Name			to facility address above	
Self Spouse Child	Policy Holder (if not self)					raciiity Name		iu invoice i	to facility dataless above	
Insurance	Policy #		НМО			Address				
Company	-		Auth #							
Special Billing Notes:						Contact Name				
					-					
						Phone Number		E-mail/	гах	
								Chack	(Payable to Ambry Genetics)	
						PATIENT PAYI			Card (Call 949-900-5795)	
Patient Acknowledgement: I acknowledge th	nat the information provided by me i	s true and correct.	. For direct insuran	ce billing: I authori	ize my insurance be	enefits to be paid directly to	o Ambry Ge	_		
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.										
I agree to be contacted regarding future research studies for which I may be a candidate. Any future research projects will be subject to a separate informed consent process and participation is voluntary. Learn more about Ambry's								y. Learn more about Ambry's		
privacy practices at https://www.ambrygen.c			Name of the second s	dahara da da da		and an a feature of the feature of the feature of the second second second second second second second second s			Deserver al state	
For patient payment by credit card: I hereby total annual gross household income: \$_	_ and the number of family mer	nbers in the house							e Program, please provide the 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.										



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

## Pulmonology Test Requisition Form - Page 2 of 3

Please check the box next to the test(s) being ordered below. 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.

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	Check to order			Description		
Congenital Central Hypoventilation Syndrome				Clinical Genomics					
	Congenital central hypoventilation syndrome	1580	PHOX2B gene sequence		SNP Array	5490	Chromosomal microarray (>2.6 mil- lion copy number probes and 750,000 SNP probes)		
Cystic Fibrosis						Paid option. Only available following SNP Array (5490) completed at Ambry.			
	508 FIRST®	1002	☐ Report poly T/TG status Test for CFTR deltaF508 mutation with reflex to CFTR gene sequence and deletion/duplication		Familial targeted microarray	5495	Incidental findings unrelated to the variant(s) detected in the proband, will NOT be reported. Name of proband tested at Ambry:		
	CFTR gene sequence and deletion/duplication analysis	1007	Report poly T/TG status		ExomeNext®-Proband	9993	Proband only exome sequencing Secondary Findings†: □ Opt-out		
Primary C	Ciliary Dyskinesia	8122	Report poly T/TG status		ExomeNext®-Proband plus mtDNA	9994	Proband only exome sequencing plus mtDNA sequencing Secondary Findings†:  Opt-out		
Pulmonar	Pulmonary Fibrosis				ExomeNext®-Duo	9991	Duo exome sequencing Secondary Findings†:		
Respirato	Telomere-related pulmonary fibrosis pry Distress Syndrome	8140	TERT, TERC		ExomeNext®- <i>Duo</i> plus mtDNA	9992	Duo exome sequencing Secondary Findings†: Opt-out[See ExomeNext-Trio]		
	Surfactant dysfunction panel	8100	ABCA3, SFTPB, SFTPC gene sequence		ExomeNext®-Trio	9995	Trio exome sequencing Secondary Findings†:□ Opt-out		
FOR PRENATAL SPECIMENS, POC OR CORD BLOOD: MATERNAL CELL CONTAMINATION ANALYSIS REQURED. Both test codes required for fetal specimens				ExomeNext®- <i>Trio</i> plus mtDNA	9996	Trio exome sequencing plus mtDNA sequencing Secondary Findings†:□Opt-out			
1262	1262  MCC Reference for maternal blood sample (No Charge)				ExomeNext- <i>Rapid®††</i>	9999R	Trio exome sequencing plus mtDNA sequencing Secondary Findings†:		
	SPECIFIC SITE ANALYSIS (Please include a copy of relative's report)    Gene(s):			Order through Ambry-	ExomeNext®-Select	9500	Up to 500 gene custom exome sequencing test		
Relative Name:			Port®						
	·		Iready at Ambry 🔲 not available						



### Supplemental Information - Page 3 of 3

#### **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. 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 page 2 for Maternal Cell Contamination sample submission test codes.