

Clinical Genomics Test Requisition Form - Page 1 of 6 (Exome Sequencing and Microarray)

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

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)

2. PATIENT INFORMATION									
Legal Name (Last, First, MI)					Date of Birth (MM/DD/YY)	Sex Assign	ed Ger	nder (optional)	
	Tame (Last, First, Mi) Sex Assign at Birth						Man Woman Nonbinary		
Genetic Ancestry: 🗌 Ashkenazi Jewi	ish □Asian □Black/Africar	n American	French Canadian/C	ajun		lediterranean	I	MRN	
☐ Middle Eastern ☐ Native Americ	an 🗌 Pacific Islander 🗌 Portu	ıguese □W	/hite 🗌 Unknown 🗌	Other	:				
Address			City				State		Zip
Mobile # Email							Preferrec	l Billing nce □Self-pay [Institutional
SPECIMEN INFORMATION*			nents for details)						
Personal history of allogenic bone	marrow or peripheral stem cell	transplant		Cu	irrent diagnosis of heme r	nalignancy, T	ype:		
Specimen ID:			Medical Record #						
*Fetal specimens, cord blood and POC v sample submission test codes	vill have maternal cell contamina	tion studies ac	dded for a charge. Mate	rnal an	d fetal specimen required.	Please see pag	e 4 for Ma	ternal Cell Contami	nation
Collection Assistance: Phlebotomy ** As the patient's clinician, I am unawa patient if the safety of the phlebotomist	are of any potential for complicat	ion or difficulty							
ORDERING PHYSICIAN/SEN			I receive a copy of the	report	t)				
Facility Name (Facility Code)	Address		City			e /Country	Zip	Phor	1e
	, (dd. 655						p		
Ordering Licensed Provider Name (La	ast, First)(Code) N	IPI#	Phone		Fax	Fax/I	Email		
ADDITIONAL RESULTS RECI	PIENTS								
Genetic Counselor or Other Medical	Provider Name (Last, First) (Co	ode)	Phone/Fa	x/Ema	ail				
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. I agree to allow Ambry Genetics to facilitate the provision of pre-test genetic counseling services by a third-party service, as required by the patient's insurance provider. Furthermore, all information on this TRF is true to the best of my knowledge. My signature applies to the attached letter of medical necessity.									
Signature Required for Processing	Medical Professional Sign	nature:					l	Date:	
INSURANCE BILLING (Inclu	de copy of both sides of insura	ince card)				TITUTION	IAL BILL	.ING	
Patient Relation to Policy Holder?	Name and DOB of				Facility	Name	Sei	nd invoice to facility	y address above
□Self □Spouse □Child	Policy Holder (if not self)								
Insurance Company	Policy #		HMO Auth #		Addres	S			
Special Billing Notes:					Contac	t Name			
					Phone	Number		E-mail/Fax	
					□ PA	ΓΙΕΝΤ ΡΑΥ	MENT		e to Ambry Genetics) all 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. 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 privacy practices at https://www.ambrygen.com/lega/notice-of-privacy-practices. 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 Patient Assistance Program, please provide the total annual gross household income: more about 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.									



Clinical Genomics Test Requisition Form - Page 2 of 6

ONLY COMPLETE FOR EXOMENEXT-DUO/TRIO ORDERS OR IF FAMILY MEMBERS WILL BE SUBMITTED FOR CO-SEGREGATION.

All family member specimens must be received within 4 weeks of order. Otherwise test will be run as proband only.

FAMILY MI	ember #1 info	RMATION						
Legal Name (Last, First, MI)			Date of Birt	1 (MM/DD/YY) Da	te of Death (If applicable	Phone Number/Email	I	
Sex Assigned at Birth: □F□M	Gender (optional)	□ Nonbinary	Genetic Ancestry: Ash Mediterranean Mic Other:					
Address: 🗌 S	ame as Proband	Address		City		State	Zip	Relationship to proband
SPECIMEN	INFORMATION	*(Please see am	brygen.com/specimen-requirem	ents for details)				
Personal his	tory of allogenic bone	e marrow or per	ripheral stem cell transplant	Current diagnosis	of heme maligr	nancy, Type:		
Collection Date	е	SI	pecimen ID				Medical Record #	
*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.								
Collection Assistance: Phlebotomy draw** Send saliva kit to patient Send buccal kit to patient I Insurance preverification first (available for ExomeNext and SNP array only) ** 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.								
CLINICAL INFORMATION								
Does the family member have any features similar to the proband? Yes No Partially Possibly								
Describe:								
SECONDARY FINDINGS								
	Secondary findings results are available for each family member sequenced as part of the trio. Check below to opt-out of the ACMG Recommended List of secondary findings. If left unchecked, sec- ondary findings will be reported.							

Opt-out: I choose to decline the ACMG Recommended List of secondary findings.

FAMILY ME	MBER #2 INFO	RMATION						
Legal Name (Last, First, MI)				D	ate of Birth (MM/DD/YY)	Date of Death (If applicable	Phone Number/Email	
Sex Assigned	Gender (optional)		Genetic Ancestry: 🗌 Ash	etic Ancestry: 🗌 Ashkenazi Jewish 📋 Asian 🔲 Black/African American 📄 French Canadian/Cajun 🗌 Hispanic/Latino				
at Birth:	🗌 Man 🗌 Woman	□ Nonbinary	☐ Mediterranean ☐ Mic	□Mediterranean □Middle Eastern □Native American □Pacific Islander □Portuguese □White □Unknown				
DF DM	Self-described		□Other:					
Address: Same as Proband Address				City Sta		State	Zip	Relationship to proband
SPECIMEN INFORMATION* (Please see ambrygen.com/specimen-requirements for details)								
Personal history of allogenic bone marrow or peripheral stem cell transplant								
Collection Date Specimen ID						Medical Record #		
*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 <u>ambrygen.com/specimen-requirements</u> for details.								
Collection Assistance: Phlebotomy draw** Send saliva kit to patient Send buccal kit to patient I Insurance preverification first (available for ExomeNext and SNP array only)								

** 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.

CLINICAL INFORMATION

Does the family member have any features similar to the proband?

Describe:

SECONDARY FINDINGS

Secondary findings results are available for each family member sequenced as part of the trio. 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.

Note: Additional relatives may be submitted for co-segregation analysis, free of charge. Please complete "Clinical Genomics Family Member TRF" if additional relatives will be included.



Clinical Genomics Test Requisition Form - Page 3 of 6

INDICATION(S) FOR TESTING						
ICD-10 code(s):	Will medical management cha ☐ Yes ☐ No	ange depending upon the results of the test?				
PROBAND'S PRIMARY INDICATION FOR TESTING						
Please describe in a few words the main reason for ordering	exome testing (Please also provide clinic notes and pedigree):					
PROBAND'S CLINICAL OVERVIEW (Check yes for	all that apply)					
 Yes □ No Audiologic/Otolaryngologic Yes □ No Cardiovascular Yes □ No Craniofacial Yes □ No Dental Yes □ No Dermatologic Yes □ No Endocrine Yes □ No Fetal (<i>Please complete and attach "ExomeNext Prenatal Form</i>") Yes □ No Gastrointestinal Yes □ No Genitourinary Yes □ No Growth Disorders: □ Yes □ No Overgrowth □ Yes □ No Failure to thrive 	 Yes □ No Hematologic Yes □ No Immunologic/Infectious/Allergy Yes □ No Metabolic/Biochemical Yes □ No Movement Disorder Yes □ No Musculoskeletal/Structural Yes □ No Multiple Congenital Anomalies Yes □ No Neurologic □ Yes □ No Seizures/Epilepsy □ Yes □ No Autism Spectrum Disorder □ Yes □ No Autism Spectrum Disorder □ Yes □ No Autia/Spasticity □ Yes □ No Abnormal brain MRI □ Yes □ No Obstetric □ Yes □ No Oncologic 	 Yes □ No Ophthalmologic Yes □ No Pulmonary Yes □ No Renal Yes □ No Tone abnormalities □ Yes □ No Hypotonia □ Yes □ No Hypertonia 				
ADDITIONAL CLINICAL DETAILS						
Dysmorphic Features (describe):	viors (describe): pilepsy Seizure type(s):					
Previous Studies						
MRI/CT studies (findings): Chromosome analysis: Microarray analysis: Other molecular studies:						
Growth Indices (current): Head circumference:	% Weight:% Height:	%				
Differential diagnosis/Genes of interest:						
FAMILY HISTORY (Please attach pedigree)						
Is anyone in the family affected with a similar phenotype as the proband? NO YES, please list exact relationship to proband, symptoms and age of onset of symptoms:						



Clinical Genomics Test Requisition Form - Page 4 of 6

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 Reflex	or Concurrent Testing:					
Test 1:						
	□ Concurrent with □ Concurrent with					
See Reflex	or Concurrent Testing section		upplemental Information page.			
Check	Test Name	Test Code	Description	SINGLE SITE ANALYSIS (Please include a copy of relative's report)		
Exome	1	1	1	Gene(s): Mutation(s):		
REC	UIRED: Select a Primary Test	Order		Relative Name:		
			Proband only exome sequencing	Relationship to Relative: Accession # (If tested at Ambry):		
	ExomeNext®-Proband	9993	Secondary Findings*: Opt-out	Positive control sample: 🗌 will be provided 🔲 already at Ambry 🗌 not available		
	ExomeNext®-Proband	9994	Proband only exome sequencing plus mtDNA sequencing	FOR PRENATAL SPECIMENS, POC OR CORD BLOOD: MATERNAL CELL CONTAMINATION ANALYSIS REQUIRED		
	plus mtDNA		Secondary Findings*:	Both test codes required for fetal specimens.		
	ExomeNext®-Duo	9991	Duo exome sequencing Secondary Findings*: ☐ Opt-out	 1260 MCC for amniotic fluid culture or CVS 1262 MCC Reference for maternal blood sample (No Charge) 		
	ExomeNext®-Duo plus		Duo exome sequencing plus mtDNA	OTHER ORDER		
	mtDNA	9992	sequencing Secondary Findings*:	Please visit ambrygen.com/tests for details.		
			Trio exome sequencing	Test Code:Test Name:		
	ExomeNext®-Trio	t®- <i>Trio</i> 9995 Secondary Findings*: □ O		Notes:		
	ExomeNext®- <i>Trio</i> plus mtDNA	9996	Trio exome sequencing plus mtDNA sequencing			
		Secondary Findings*:				
	ExomeNext-Rapid® (Institutional billing or	9999R	Rapid Trio exome sequencing plus mtDNA sequencing			
	patient payment only)		Secondary Findings*: Opt-out			
ExomeNe	ext Supplemental Test Opti	ons				
	ExomeReveal™	9990	RNA analysis available with all ExomeNext orders except for ExomeNext- <i>Rapid</i> , EDTA and PAX- gene RNA tubes required			
Fragile X	syndrome and Chromoson	nal Microa	array			
	Fragile X syndrome	4544	FMR1 repeat expansion analysis and			
			methylation studies	ORDERING CHECKLIST (Required')		
	SNP Array	5490	Chromosomal microarray (>2.6 million copy number probes and	Proband specimen		
			750,000 SNP probes)	Clinical Genomics TRF with patient & clinician signatures		
			Paid option. Only available following SNP Array (5490) completed at	Clinical history (attach clinic notes)		
	Familial targeted		Ambry. Incidental findings unrelated	Medical Necessity Form (insurance orders only) (see page 5)		
	microarray	5495	to the variant(s) detected in the proband, will NOT be reported. Name of proband tested at Ambry:	Copy of Insurance Card (insurance orders only)		
				Orders with missing requirements will be placed on hold until all requirements are received.		
				ORDERING CHECKLIST (Highly Recommended)		
	*Secondary Findings: If box is left unchecked, the ACMG recommended list of Secondary Findings will be reported.			Family member specimens Please send all first degree and other informative relatives within 4 weeks of the order.		
				☐ Family history or pedigree		
				Previous test results		

CONTACT INFORMATION

For ExomeNext preverification requests please send the Medical Necessity Form and Clinical Genomics TRF to preverification@ambrygen.com or fax to 949-900-5501.

All other documents can be secure uploaded at ambrygen.com/secure-upload, or faxed to 949-900-5501.

AmbryPort is a secure client portal that allows order submission, test status updates, insurance authorization status and report downloads. All required documents can be completed and directly uploaded through AmbryPort during the ordering process or after order submission. Please visit portal.ambrygen.com/signup to sign up.



Supplemental Information - Page 5 of 6

Specimen Requirements

Blood/saliva/buccal swab sample from patients with a history of allogenic bone marrow or stem cell transplant cannot be used for genetic testing. Blood/saliva/buccal swab sample from patients with active hematological disease is not recommended. An alternative specimen may be needed. See <u>ambrygen.com/specimen-requirements</u> 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 4 for Maternal Cell Contamination sample submission test codes.

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.



ExomeNext Medical Necessity Form - Page 6 of 6

REQUIRED FOR INSURANCE ORDERS ONLY (NOT REQUIRED FOR CIGNA MEMBERS)

This form is required if you are ordering Exome testing and wish to have the patient's insurance billed. Please complete and submit with the TRF and a copy of clinical notes. This form replaces the Letter of Medical Necessity.

1. Has the patient had previous Whole Exome Sequencing (WES) performed?

Yes, date performed: ______

🗌 No

2. Does this patient have a clinical presentation consistent with the following (select all that apply):

□ Multiple abnormalities affecting unrelated organ systems (please specify): ____

OR two of the following:

□ Abnormality affecting a single organ system(specify): ___

Significant intellectual disability, symptoms of a complex neurodevelopmental disorder (i.e. self-injurious behavior, reverse sleep-wake cycle, or seizure/epilepsy), or severe neuropsychiatric condition (e.g. schizophrenia, bipolar, Tourette syndrome)

□ Period of unexplained developmental regression (unrelated to autism or epilepsy)

3. Are the results of this WES test expected to directly influence this patient's medical management recommendations and clinical outcome?

☐ Yes (please describe): ____

🗌 No

4. Please describe the genetic tests that would be indicated if WES were NOT performed (i.e., single gene tests, gene panels, etc.):

Chromosomal microarray		
□ Single gene test(s):	 	
Multigene panel(s):	 	
Other genetic test(s):	 	

5. Please describe follow-up procedures & frequency that would be needed if WES were NOT performed (i.e., lumbar puncture, imaging studies, brain MRI, etc.):

Imaging study:	
□ Surgery:	
□ Biopsy:	
□ Other:	