

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)
<p>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)</p>

PATIENT INFORMATION			
Name (Last, First, MI)	Sex at Birth <input type="checkbox"/> F <input type="checkbox"/> M	Date of Birth (MM/DD/YY)	MRN
Ethnicity: <input type="checkbox"/> Asian <input type="checkbox"/> Ashkenazi Jewish <input type="checkbox"/> Black/African American <input type="checkbox"/> White <input type="checkbox"/> French Canadian/Cajun <input type="checkbox"/> Hispanic/Latino <input type="checkbox"/> Mediterranean <input type="checkbox"/> Middle Eastern <input type="checkbox"/> Native American <input type="checkbox"/> Pacific Islander <input type="checkbox"/> Portuguese <input type="checkbox"/> Unknown <input type="checkbox"/> Other:			
Address	City	State	Zip
Phone	Email	Preferred Billing <input type="checkbox"/> Insurance <input type="checkbox"/> Self-pay <input type="checkbox"/> Institutional	

SPECIMEN INFORMATION* (Please see ambrygen.com/specimen-requirements for details)	
<input type="checkbox"/> Personal history of allogenic bone marrow or peripheral stem cell transplant	<input type="checkbox"/> Current diagnosis of heme malignancy, Type:
Specimen ID: <input type="checkbox"/> Send saliva kit to patient	Medical Record #
<i>*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</i>	
Collection Assistance: <input type="checkbox"/> Phlebotomy draw** <input type="checkbox"/> Send saliva kit to patient <input type="checkbox"/> Insurance preverification first (available for ExomeNext and SNP array only) <i>** 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.</i>	

ORDERING PHYSICIAN/SENDING FACILITY (Each listed person will receive a copy of the report)				
Facility Name (Facility Code)	Address	City	State/Country	Zip
Ordering Licensed Provider Name (Last, First)(Code)	NPI#	Phone	Fax	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. 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 Signature:	Date:
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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
Special Billing Notes:		Contact Name	
		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 Patient Assistance 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:
 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.

Signature Required For Insurance/Self-Pay Patients and NY Sample Storage Consent:	Date:
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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 MEMBER #1 INFORMATION				
Name (Last, First, MI)		Date of Birth (MM/DD/YY)	Date of Death (if applicable)	Phone Number/Email
Biological Sex: <input type="checkbox"/> F <input type="checkbox"/> M	Ethnicity: <input type="checkbox"/> Asian <input type="checkbox"/> Ashkenazi Jewish <input type="checkbox"/> Jewish (non-Ashkenazi) <input type="checkbox"/> Black/African American <input type="checkbox"/> White <input type="checkbox"/> French Canadian/Cajun <input type="checkbox"/> Hispanic/Latino <input type="checkbox"/> Mediterranean <input type="checkbox"/> Middle Eastern <input type="checkbox"/> Native American <input type="checkbox"/> Pacific Islander <input type="checkbox"/> Portuguese <input type="checkbox"/> Unknown <input type="checkbox"/> Other:			
Address: <input type="checkbox"/> Same as Proband	Address	City	State	Zip
				Relationship to proband
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, Source: _____ <input type="checkbox"/> Other: _____				
<input type="checkbox"/> Personal history of allogenic bone marrow or peripheral stem cell transplant		<input type="checkbox"/> Current diagnosis of heme malignancy, Type: _____		
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 ambrygen.com/specimen-requirements for details.				
Phlebotomy Services Request: <input type="checkbox"/> Phlebotomy draw <input type="checkbox"/> Insurance preverification first <input type="checkbox"/> Send blood kit to patient* <input type="checkbox"/> Send saliva 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.				
CLINICAL INFORMATION				
Does the family member have any features similar to the proband? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Partially <input type="checkbox"/> 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, secondary findings will be reported. <input type="checkbox"/> Opt-out: I choose to decline the ACMG Recommended List of secondary findings.				
FAMILY MEMBER #2 INFORMATION				
Name (Last, First, MI)		Date of Birth (MM/DD/YY)	Date of Death (if applicable)	Phone Number/Email
Biological Sex: <input type="checkbox"/> F <input type="checkbox"/> M	Ethnicity: <input type="checkbox"/> Asian <input type="checkbox"/> Ashkenazi Jewish <input type="checkbox"/> Jewish (non-Ashkenazi) <input type="checkbox"/> Black/African American <input type="checkbox"/> White <input type="checkbox"/> French Canadian/Cajun <input type="checkbox"/> Hispanic/Latino <input type="checkbox"/> Mediterranean <input type="checkbox"/> Middle Eastern <input type="checkbox"/> Native American <input type="checkbox"/> Pacific Islander <input type="checkbox"/> Portuguese <input type="checkbox"/> Unknown <input type="checkbox"/> Other:			
Address: <input type="checkbox"/> Same as Proband	Address	City	State	Zip
				Relationship to proband
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, Source: _____ <input type="checkbox"/> Other: _____				
<input type="checkbox"/> Personal history of allogenic bone marrow or peripheral stem cell transplant		<input type="checkbox"/> Current diagnosis of heme malignancy, Type: _____		
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 ambrygen.com/specimen-requirements for details.				
Phlebotomy Services Request: <input type="checkbox"/> Phlebotomy draw <input type="checkbox"/> Insurance preverification first <input type="checkbox"/> Send blood kit to patient* <input type="checkbox"/> Send saliva 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.				
CLINICAL INFORMATION				
Does the family member have any features similar to the proband? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Partially <input type="checkbox"/> 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, secondary findings will be reported. <input type="checkbox"/> Opt-out: I choose to decline the ACMG Recommended List of secondary findings.				

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INDICATION(S) FOR TESTING		
ICD-10 code(s):	Will medical management change depending upon the results of the test? <input type="checkbox"/> Yes <input type="checkbox"/> No	
PROBAND'S PRIMARY INDICATION FOR TESTING		
Please describe in a few words the main reason for ordering exome testing (<i>Please also provide clinic notes and pedigree</i>):		
PROBAND'S CLINICAL OVERVIEW (Check all that apply)		
<input type="checkbox"/> Audiologic/Otolaryngologic <input type="checkbox"/> Cardiovascular <input type="checkbox"/> Craniofacial <input type="checkbox"/> Dental <input type="checkbox"/> Dysmorphic Features <input type="checkbox"/> Dermatologic <input type="checkbox"/> Endocrine <input type="checkbox"/> Fetal (<i>Please complete and attach "ExomeNext Prenatal Form"</i>) <input type="checkbox"/> Gastrointestinal <input type="checkbox"/> Genitourinary <input type="checkbox"/> Growth Disorders: <input type="checkbox"/> Undergrowth <input type="checkbox"/> Overgrowth <input type="checkbox"/> Failure to thrive	<input type="checkbox"/> Hematologic <input type="checkbox"/> Immunologic/Infectious/Allergy <input type="checkbox"/> Metabolic/Biochemical <input type="checkbox"/> Movement Disorder <input type="checkbox"/> Musculoskeletal/Structural <input type="checkbox"/> Multiple Congenital Anomalies <input type="checkbox"/> Neurologic <input type="checkbox"/> Seizures/Epilepsy <input type="checkbox"/> Autism Spectrum Disorder <input type="checkbox"/> Developmental Delay/Intellectual disability <input type="checkbox"/> Ataxia/Spasticity <input type="checkbox"/> Psychiatric <input type="checkbox"/> Abnormal brain MRI <input type="checkbox"/> Obstetric <input type="checkbox"/> Oncologic	<input type="checkbox"/> Ophthalmologic <input type="checkbox"/> Pulmonary <input type="checkbox"/> Renal <input type="checkbox"/> Tone abnormalities <input type="checkbox"/> Hypotonia <input type="checkbox"/> Hypertonia
ADDITIONAL CLINICAL DETAILS		
Autism: <input type="checkbox"/> no autistic behaviors <input type="checkbox"/> autistic behaviors (describe): _____		
Dysmorphic Features (describe): _____		
Congenital Anomalies (describe): _____		
History of Seizures <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> diagnosed epilepsy Seizure type(s): _____		
Progressive disease <input type="checkbox"/> Yes <input type="checkbox"/> No		
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? <input type="checkbox"/> NO <input type="checkbox"/> YES, please list exact relationship to proband, symptoms and age of onset of symptoms: _____ _____		
Is there any consanguinity (conception between blood relatives) in the family? <input type="checkbox"/> NO <input type="checkbox"/> YES If yes please describe: _____ _____		

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Please check the box next to the test(s) being ordered below. 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. 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
Chromosomal Microarray Analysis			
<input type="checkbox"/>	SNP Array^^	5490	Chromosomal microarray (>2.6 million copy number probes and 750,000 SNP probes)
<input type="checkbox"/>	Familial targeted microarray^^	5495	Paid option. Only available following SNP Array (5490) completed at Ambyr. Incidental findings unrelated to the variant(s) detected in the proband, will NOT be reported. Name of proband tested at Ambyr: _____
Exome Sequencing			
<input type="checkbox"/>	ExomeNext®-Proband	9993	Proband only exome sequencing Secondary Findings ¹ : <input type="checkbox"/> Opt-out
<input type="checkbox"/>	ExomeNext®-Proband plus mtDNA	9994	Proband only exome sequencing plus mtDNA sequencing Secondary Findings ¹ : <input type="checkbox"/> Opt-out
<input type="checkbox"/>	ExomeNext®-Duo	9991	Duo exome sequencing Secondary Findings ¹ : <input type="checkbox"/> Opt-out
<input type="checkbox"/>	ExomeNext®-Duo plus mtDNA	9992	Duo exome sequencing plus mtDNA sequencing Secondary Findings ¹ : <input type="checkbox"/> Opt-out
<input type="checkbox"/>	ExomeNext®-Trio	9995	Trio exome sequencing Secondary Findings ¹ : <input type="checkbox"/> Opt-out
<input type="checkbox"/>	ExomeNext®-Trio plus mtDNA	9996	Trio exome sequencing plus mtDNA sequencing Secondary Findings ¹ : <input type="checkbox"/> Opt-out
<input type="checkbox"/>	ExomeNext-Rapid® (Institutional billing or patient payment only)	9999R	Rapid Trio exome sequencing plus mtDNA sequencing Secondary Findings ¹ : <input type="checkbox"/> Opt-out
Order through Ambyr-Port®	ExomeNext®-Select	9500	Up to 500 gene custom exome sequencing test
^^ Buccal swab accepted for SNP Array ¹ Secondary Findings: If box is left unchecked, the ACMG recommended list of Secondary Findings will be reported. Secondary Findings are not available for ExomeNext-Select orders.			

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: will be provided already at Ambyr not available

FOR PRENATAL SPECIMENS, POC OR CORD BLOOD: MATERNAL CELL CONTAMINATION ANALYSIS REQUIRED
 Both test codes required for fetal specimens.

- 1260 MCC for amniotic fluid culture or CVS
 1262 MCC Reference for maternal blood sample (No Charge)

OTHER ORDER

 Please visit ambrygen.com/tests for details.
 Test Code: _____ Test Name: _____

 Notes: _____

ORDERING CHECKLIST (Required¹)

- Proband specimen
 Clinical Genomics TRF with patient & clinician signatures
 Clinical history (attach clinic notes)
 Medical Necessity Form (insurance orders only) (see page 5)
 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)

- 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.

AmbyrPort 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 AmbyrPort during the ordering process or after order submission. Please visit portal.ambrygen.com/signup to sign up.

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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 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 4 for Maternal Cell Contamination sample submission test codes.

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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)
- Family history strongly implicating a genetic etiology (please specify findings and relationships): _____
- 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: _____