

THIS IS NOT A TEST REQUEST FORM. Please complete and submit with the test request form or electronic packing list.

EXOME SEQUENCING INTAKE FORM

Failure to provide required information for exome testing will result in a suboptimal clinical report and delays in testing.

REQUIRED for Exome Sequencing (ARUP test code 3016583):

1. Proband Patient Name: Date of Birth:

Suspected Clinical Diagnosis: ____ 2.

Provide medical records detailing the patient's phenotype/relevant previous testing/family history or complete the Additional Clinical Information section found in this document.

Parental/familial control samples (RECOMMENDED; ARUP test code 3016589). Submit comparator samples within 7 days of the 3. proband's sample.

a. Maternal Last Name, First Name:

Clinically affected?

No
Yes: Date of birth:

b. Paternal Last Name, First Name:

Date of birth:

Clinically affected? \Box No \Box Yes: _

Additional Familial Control (if applicable) Name: C.

Relationship to proband:

__ Clinically affected? \Box No \Box Yes: ____ Date of birth:

Ordering Provider Attestation of Informed Consent (signature required below) 4.

Test Purpose and Description

The purpose of exome sequencing is to identify the gene variant(s) causing a suspected Mendelian genetic condition. Testing requires drawing 2 mL of blood from which the DNA is extracted. DNA codes for genes. Most of the patient's genes will be sequenced.

Thousands of DNA variants will be detected by sequencing. DNA variants may be disease causing or harmless; however, the effect of many DNA variants is currently unknown.

Ordering Considerations

Participation in exome sequencing is voluntary. Genetic counseling is recommended before and following this complex test.

The chance a cause for the patient's medical issue(s) can be determined using this test varies and is influenced by the specific clinical features present. Diagnostic rates are highest when biological parents' samples are included as comparators for exome sequencing. Parental sequence data is used to identify de novo (new) changes in the patient's DNA, not found in either parent, that could explain the patient's disorder.

It is important that the familial biological relationships are correctly stated because undisclosed adoption or uncertain paternity will cause confusion and decrease the chance of identifying the causative disease variant.

Exome sequencing may identify genetic findings unrelated to the original reason for testing such as:

Predict another family member has, is at risk for, or is a carrier of an unsuspected genetic condition

- Reveal nonpaternity (the person stated to be the biological father is not, in fact, the biological father)
- · Indicate the biological parents of the patient are close blood relatives (consanguineous)

Reporting of Results

Results are generally available in 3-4 weeks. Variants that are known or suspected to be causative for the patient's medical issues are reported.

All variants identified in the patient that are related to the patient's primary disorder will be tested in familial controls. The status of all primary variants tested in controls will be indicated on the proband's report.

Candidate variants that are not known to be causative for the patient's medical issues, such as de novo variants (not inherited from either parent) or variants inherited from both parents in the same gene, may be reported even if the function of the gene is unknown.

Variants in genes unrelated to the patient's medical condition are not usually reported except for disease-causing secondary findings (see the Secondary Findings section below).

Because genetic knowledge is advancing at a rapid pace, reanalysis of exome sequencing data should be considered 12-18 months after testing is complete if a cause for the patient's condition was not determined. ARUP will perform reanalysis (ARUP test code 3001457) of exome data. If the report is amended, the ordering client will receive an updated report.

Limitations

Although genetic test results are usually accurate, several sources of error are possible, including clinical misdiagnosis of a condition, sample mislabeling or contamination, transfusion, bone marrow transplantation, maternal cell contamination of cord blood samples, or inaccurate information regarding biological relationships. If biological relationships are inaccurately reported, it could lead to an incorrect diagnosis or inconclusive result. ARUP will contact the referring provider if nonpaternity and/or consanguinity is detected but that information will not be included in the patient's report.

Often, exome sequencing is not able to identify the cause of a patient's medical issues. This does not exclude the possibility that the patient has a genetic condition. Some disease-causing variants are in genes with unknown function while others may not be identifiable using this test. Examples of variants not detectable with this test include large gene deletion/duplications, variants occurring outside of the coding region and intron-exon boundaries, chromosome rearrangements, inversions, methylation abnormalities, and those causing repeat disorders. This test does not include sequencing of the mitochondrial genome.

Secondary Findings

The American College of Medical Genetics and Genomics (ACMG) recommends reporting disease-causing variants in specific genes that increase the risk for developing cancer, cardiovascular issues, metabolic disorders, problems with anesthesia, retinopathy, and other conditions where monitoring or early treatment may be available. Please refer to the latest version of the ACMG recommendations for reporting secondary findings in clinical exome sequencing for a list of genes and associated disorders tested. Additional medically actionable variants in non-ACMG genes may be reported at ARUP's discretion.

If a patient has symptoms of a condition related to an ACMG recommended gene, separate testing should be ordered, as coverage of ACMG genes may be incomplete. Only variants in ACMG genes identified by routine exome sequencing are reported. Single disease-causing variants in recessive ACMG genes are not reported.

To receive secondary findings about the patient, the patient (or their legal guardian) would need to choose to receive this information by

selecting the "opt in" option on this intake form. Familial controls who undergo exome sequencing and desire a full analysis and report of their secondary findings must select the "opt in" option on this intake form and will incur a separate fee. Secondary findings will be reported for familial controls who elect to receive this information regardless of whether the finding was also identified in the patient. Parental inheritance of secondary findings identified in the patient will only be included in the patient's report if the positive parent also opts to receive secondary findings.

If a disease-causing genetic variant is identified, insurance rates, the ability to obtain disability and life insurance, and employability could be affected. The Genetic Information Nondiscrimination Act of 2008 extends some protections against genetic discrimination (genome.gov/10002328). All test results are released to the ordering healthcare provider and those parties entitled to them by federal, state, and local laws.

Access to Sequence Data/Data Sharing/Sample Storage

ARUP Laboratories will have access to the patient's sequence data from exome sequencing. Your healthcare provider and the hospital that submitted the test to ARUP can also request a copy of the sequence data.

Because ARUP is not a storage facility, most samples are discarded after testing is completed. Some samples may be stored indefinitely for test validation or education purposes after personal identifiers are removed. You may request disposal of your sample by calling ARUP Laboratories at 800-242-2787 ext. 3301.

In cooperation with the National Institutes of Health's effort to improve understanding of specific genetic variants, ARUP submits HIPAA-compliant, deidentified (cannot be traced back to the patient) genetic test results and health information to public databases. The confidentiality of each sample is maintained. If you prefer that your test result not be shared, call ARUP at 800-242- 2787 ext. 3301. Your deidentified information will not be disclosed to public databases after your request is received, but a separate request is required for each genetic test. Additionally, patients have the opportunity to participate in patient registries and research. To learn more, visit aruplab.com/genetics.

Ordering Healthcare Provider, Genetic Counselor: 1) I attest that I am the ordering healthcare provider or certified genetic counselor; 2) I have explained the purpose/benefits and limitations of the test to the patient or their legal guardian and all parental controls; 3) the patient/legal guardian and parental controls were offered copies of this consent document; 4) I have answered all of their questions regarding the purpose of the test, the reporting of primary and secondary findings, the use and retention of samples, and data sharing.

Ordering Provider/Genetic Counselor Printed Name	Signature	Date
Secondary findings identified WILL NOT be examined and reported for fee applies for generating a clinical report with secondary findings for	•	SS the corresponding box is checked. A
□ Opt in to report secondary findings for the PATIENT.		
\Box Opt in to report secondary findings for the MATERNAL CONTROL.		
\Box Opt in to report secondary findings for the PATERNAL CONTROL.		
\square Opt in to report secondary findings for ADDITIONAL FAMILIAL CONT	TROL. (name(s):)

For questions, contact an ARUP genetic counselor at 800-242-2787 ext. 2141.

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Exome Sequencing Additional Clinical Information:

Please provide medical records detailing the patient's phenotype/relevant previous testing/family history or complete the Additional Clinical Information section below. The ability to identify causative variant(s) for the patient's presentation is influenced by the quality of the clinical information provided.

information provided.					
Ordering Provider:		Provider's P	hone:		
Practice Specialty:		Provider's F	ax:		
Genetic Counselor		Counselor's	Phone:		
Ethnicity/Ancestry:	erican/Black	🗆 Asian	🗆 Hispanic	🗆 White	Other
Genes of Interest:					
Family History:					
PRE/PERINATAL	□ 0002126	Polymicrogyria		□ 0000750 Delayed spee	ch and
0000776 Congenital diaphragmatic			language development		
hernia		· ····································		0002376 Development	
0001627 Congenital heart defect				□ 0001263 Global develo	
0000476 Cystic hygroma	NEUROLOGI			0001249 Intellectual di	• •
0002084 Encephalocele		Abnormality of move	ement	0002187 Profound	
0010945 Fetal pyelectasis				0010864 Severe	
0007430 Generalized edema				0002342 Moderate	
0001789 Hydrops fetalis				0001256 Mild	
0010880 Increased nuchal				0001270 Motor delay	
translucency		Dyskinesia Epileptic encephalop	ath.	D	
0001511 Intrauterine growth		Encephalopathy	Jatny	MUSCULOSKELETAL	
restriction		Hyperreflexia		0002804 Arthrogryposi	e multiplov
0002475 Myelomeningocele/		Involuntary moveme	nte	congenita	s multiplex
spina bifida				0003199 Decreased mi	iecle mass
0001562 Oligohydramnios		21 Absence		□ 0001371 Flexion contra	
0001539 Omphalocele				□ 0001528 Hemihypertro	
0001561 Polyhydramnios				□ 0001252 Hypotonia	5,
0001622 Prematurity-GA at birth		23 Generalized myoc	lonic	□ 0001276 Hypertonia	
		18 Generalized tonic		□ 0001382 Joint hyperme	obility
0003026 Short long bones		59 Generalized clonic		□ 0002808 Kyphosis	·····,
□ 0001518 Small for gestational age		59 Infantile	-	□ 0040064 Limb abnorma	ality
□		59 Tonic-clonic		0001324 Muscle weak	•
STRUCTURAL BRAIN ABNORMALITIES				0003198 Myopathy	
0002134 Abnormality of				0010442 Polydactyly	
basal ganglia				0002757 Recurrent frac	ctures
0002363 Abnormality of the		Abnormal CK		0002650 Scoliosis	
brain stem			- mitin -	0004322 Short stature	
0001273 Abnormality of		Decreased plasma c Hyperalaninemia	amune	0002652 Skeletal dysp	lasia
corpus callosum		Hypoglycemia		0001257 Spasticity	
0002269 Abnormality of		Hyperammonemia		0001159 Syndactyly	
neuronal migration		Increased CSF lacta	to	0001762 Talipes equine	ovarus
0007360 Aplasia/hypoplasia of the equal allows		Increased serum py		0000098 Tall stature	
the cerebellum			luvate	0000925 Vertebral colu	Imn
 0012444 Brain atrophy 0007266 Cerebral dysmyelination 		Lactic acidosis		abnormality	
		Metabolic acidosis			
 0006808 Cerebral hypomyelination 0002500 Cerebral white 		Organic aciduria		CRANIOFACIAL	
matter abnormality		l newborn screen		0000271 Abnormal fac	ies
0002539 Cortical dysplasia				0000306 Abnormality of the second secon	
□ 0002282 Heterotopia				□ 0000290 Abnormality of	
□ 0001360 Holoprosencephaly				□ 0000175 Cleft palate	
□ 0000238 Hydrocephalus		ENTAL/BEHAVIORA	L	□ 0410030 Cleft lip	
0002352 Leukoencephalopathy		Attention deficit vity disorder		0001363 Craniosynost	osis
□ 0001339 Lissencephaly		Autistic spectrum di	isorder	0000286 Epicanthus	
·····/	. 00007297	nutione opectiviti u		0000316 Hypertelorisn	ı

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EXOME SEQUENCING INTAKE FORM

- □ 0000601 Hypotelorism
- □ 0000256 Macrocephaly
- 0000252 Microcephaly

EYE AND VISION

- 🗆 0000526 Aniridia
- 0000528 Anophthalmia
- 0000618 Blindness
- □ 0000589 Coloboma
- 0000519 Congenital cataract
- D 0000568 Microphthalmia
- □ 0000639 Nystagmus
- □ 0000648 Optic atrophy
- □ 0000508 Ptosis
- 0009919 Retinoblastoma
- □ 0000486 Strabismus
- □ 0000505 Visual impairment
- EAR AND HEARING
- D 0000377 Abnormal external ear
- 0000405 Conductive hearing impairment
- 0000410 Mixed hearing impairment
- 0000407 Sensorineural hearing
- impairment
- ۰ ____

CARDIAC

- 0001713 Abnormal cardiac ventricle
- □ 0002616 Aortic root dilatation
- 0011675 Arrhythmia
- □ 0001631 Atrial septal defect
- □ 0001647 Bicuspid aortic valve
- □ 0001638 Cardiomyopathy
- □ 0001680 Coarctation of aorta
- □ 0001642 Pulmonary stenosis
- □ 0001636 Tetralogy of Fallot
- □ 0001629 Ventricular septal defect

RESPIRATORY

- □ 0002104 Apnea
- □ 0002883 Hyperventilation
- □ 0002791 Hypoventilation
- □ 0008751 Laryngeal cleft
- 0001601 Laryngomalacia

- 0002205 Recurrent respiratory infections
- 0002878 Respiratory failure
- 0002093 Respiratory insufficiency
- 0002107 Pneumothorax
- 0002206 Pulmonary fibrosis
- 0002575 Tracheoesophageal fistula
- □ 0002779 Tracheomalacia
- _____

GASTROINTESTINAL

- 0002251 Aganglionic megacolon
- 0002910 Elevated hepatic transaminase
- □ 0001508 Failure to thrive
- D 0001543 Gastroschisis
- □ 0001399 Hepatic failure
- 0002240 Hepatomegaly
- □ 0002021 Pyloric stenosis
- □ 0001744 Splenomegaly
- 0002013 Vomiting

GENITOURINARY

- 0000812 Abnormal internal genitalia
- 0000062 Ambiguous genitalia
- □ 0000028 Cryptorchidism
- 0000085 Horseshoe kidney
- □ 0000126 Hydronephrosis
- □ 0000047 Hypospadias
- 0008738 Partially duplicated kidney
- 🗆 0000113 Polycystic kidney dysplasia
- 0000107 Renal cyst
- 0000104 Renal agenesis
- 0000089 Renal hypoplasia
- 0000069 Ureter abnormality
- 0000795 Urethra abnormality

SKIN AND HAIR

- 🗆 0008066 Blistering of skin
- 🗆 0000957 Café-au-lait spot
- □ 0005306 Capillary hemangioma
- 0001595 Hair abnormality
- □ 0000974 Hyperextensible skin
- □ 0000953 Hyperpigmentation of skin
- □ 0000998 Hypertrichosis
- □ 0001010 Hypopigmentation of skin

ARUP LABORATORIES | 500 Chipeta Way, Salt Lake City, UT 84108 | phone: 801-583-2787 | toll free: 800-242-2787 | fax: 801-584-5249 | aruplab.com

D 0008066 Ichthyosis

- 0001597 Nail abnormality
- 0001581 Recurrent skin infections

HEMATOLOGY AND IMMUNOLOGY

- □ 0001928 Abnormality of coagulation
- 0004432 Agammaglobulinemia
- □ 0001903 Anemia
- 0031020 Bone marrow hypercellularity
- □ 0001878 Hemolytic anemia
- □ 0002721 Immunodeficiency
- 🗆 0001888 Lymphopenia
- □ 0001875 Neutropenia
- 0001876 Pancytopenia

immunodeficiency

П

CANCER

ENDOCRINE

glands

hyperplasia

insufficiency

□ Type of cancer

relatives

OTHER

□ Age of diagnosis

0002719 Recurrent infections
 0004430 Severe combined

0001873 Thrombocytopenia

□ 0000834 Abnormality of adrenal

0008226 Androgen insufficiency

0008258 Congenital adrenal

□ 0000819 Diabetes mellitus

0000873 Diabetes insipidus

□ 0000821 Hypothyroidism

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0001738 Exocrine pancreatic

0000829 Hypoparathyroidism

□ Family history of cancer and affected

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		Shorman coun)
Echocardiogram:	□ Not performed	□ Normal	□ Abnormal _
EEG:	□ Not performed	Normal	□ Abnormal _
EMG/NCV:	□ Not performed	□ Normal	□ Abnormal _
Gene testing:		□ Normal	□ Abnormal _
•		□ Normal	□ Abnormal
Karyotype:	□ Not performed	□ Normal	□ Abnormal _
Prenatal genomic	microarray:	□ Normal	□ Abnormal _
Postnatal genomi	c microarray: □ Not performed	□ Normal	□ Abnormal _
MRI (brain):	Not performed	Normal	□ Abnormal _
MRI (other):	□ Not performed	Normal	□ Abnormal _
CT (brain):	□ Not performed	Normal	□ Abnormal _
CT (other):	□ Not performed	Normal	□ Abnormal _
Muscle biopsy:	□ Not performed	□ Normal	□ Abnormal _
Ultrasound:	□ Not performed	Normal	□ Abnormal _
X-ray:	□ Not performed	Normal	□ Abnormal _
Other test:		□ Normal	□ Abnormal _
		□ Normal	□ Abnormal _
		Normal	
			□ Abnormal _

PREVIOUS TESTING (provide copy of abnormal results)

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