

Shodair Children's Hospital Genetics Laboratory 2620 Shodair Dr. Helena, MT 59601 Phone (406) 444-7532 Fax (406) 444-1022

Shodair Lab Number

WHOLE EXOME SEQUENCING TEST REQUEST FORM

PATIENT INFORMATION							
Last Name:	First Name:			MI: _	D	OB:	
Sex Assigned at Birth: □ Male □ Fem	nale 🗆	Gender lo	dentity: □ Male □ Fe	male 🗆 Non-bi			
Address:			Ethnicity	☐ Caucasian		☐ Ashkenazi Jewish	
City, State, Zip:			(select all that apply):	☐ Asian☐ Hispanic		☐ Hutterite ☐ American Indian	
Phone: Ema				☐ African American		☐ Other	
ORDERING HEALTH CARE PROFES	SIONAL & AUTHORIZATIO	N					
Name:		NP	·I #:				
Address:		Cit	y, State, Zip:				
Telephone:	Fax:	R	eferring Facility:				
Additional Reports To:							
By submitting this requisition, I confirm valuable for the patient. For patients		-	-		-		
Signature of Ordering Provider:			Date:		_		
ADDITIONAL FAMILY SAMPLES (ED	OTA Whole Blood or Saliva/Bu	iccal Cells)	SAMPLE INFORMA	TION			
Last Name:	First Name:		☐ Whole Blood (≥3r	•	Date of Co	llection:	
DOB: Relationship:	Affected: \square	Yes 🗆 No	☐ Saliva/Buccal Cells ☐ Extracted DNA (≥10ug)			ference #:	
Date of Collection: Sou	ırce:	-					
Last Name:	First Name:		INFORMED CONSE	NT (REOUIREI	D)		
DOB: Relationship:						nd each family member	
Date of Collection: Sou			Informed consent is required for the patient and each family member participating in whole exome sequencing prior to initiating testing. Please complete the Informed Consent section on page 2 of this form.				
PRIOR AUTHORIZATION IS REQUI	RED FOR WHOLE EXOME	SEQUENC	ING				
Prior authorization has been comp	oleted & approved, or is not roproval letter from the insura	required. I a	gree to include a [any with this form.	_		or authorization updates:	
Prior authorization has not been c I agree to provide clinical records of the front & back of the insuran	and/or a letter of medical ne	ecessity, IC	D-10 codes, a copy	Phone:			
INSURANCE BILLING Please provide	le a copy of front & back of c	ard	INSTITUTIONAL BII	LLING			
Name of policy holder:			Institution:				
Policy holder DOB: Relation to patient:			Address:				
Name of Ins. Co:			City, State, Zip:				
Ins. Co Policy #:			Phone: Fax:				
Ins. Co Phone:			Billing Contact:				
MEDICAID / MEDICARE BILLING			SELF-PAY				
Name of policy holder:			Please contact us at 406-	444-7532 for a cop	y of the Patio	ent Self-Pay Agreement.	
Policy holder DOB: Medicaid/Medicare #:			Name of responsible party:				
Address:			Relation to patient: Phone:				
City, State, Zip:			Email:				
Received	Date Received	Tracking	#	Sender	•	Initials	



Patient Name:

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Call 406-444-7532 with questions.

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	, ICD-10 CODES, AND and/or any additional cli	CLINICAL INDICATION nical information (medical records, pid	ctures, family	history) to aid in resul	t interpretation.	
Reason for Testing ☐ Diagnostic ☐ Prenatal ☐ Carrier Screening ☐ Family History ☐ No Family History	ICD-10 Codes (required)	Clinical Indication				
WHOLE EXOME SEQU Acceptable sample: EDTA	ENCING (WES) blood, saliva/buccal cells, ex	ctracted DNA				
☐ Trio Whole Exome	le Exome Sequencing (Patient + two additional family members)			Specific Test Instructions		
☐ Duo Whole Exome	Duo Whole Exome Sequencing (Patient + one additional family member)					
☐ Singleton Whole Exome Sequencing (Patient only)						
☐ Rapid Infant Whole	Exome Sequencing (d	oes not include analysis for secondary fi	ndings)			
☐ Reflex to Full V	Vhole Exome Sequenc	ing				
WHOLE EXOME SEQU	ENCING INFORMED C	ONSENT (REQUIRED)				
our website or contact used under the Sequencing Consent For would like assistance with paperwork and coordinate presentation and family Please mark your selection. Informed consent has exome sequencing content member participating results along with the Informed consent has	s at 406-444-7532 to obtom. Shodair Genetics would the consenting process te a telemedicine visit with history, and will discuss to on below. If you have an onsenting process. I agregin whole exome seque is test request form.	each family member participating in value copies of the Patient Whole Exom the delay of the Patient Whole Exom the happy to provide consenting sers, please indicate below and Shodair's thone of our genetic counselors. Dure whole exome sequencing in detail. Pley questions, please contact our office avoid like to refer the patient for a term to complete the below section provincing if applicable. I agree to provide the patient has completed the Patient Whole Exome Sequencing Consent For	e Sequencing vices to your scheduling to scheduling to see aware at 406-444-7. Elemedicine viding demog clinical reco	Consent Form and the patient and each partie am will contact the fathe genetic counselor that there will be a costa. Visit with a genetic couragnaphic and contact in reds including recent viquencing Consent Form	e Family Member Whole Exome cipating family member. If you mily to complete registration will review the patient's est associated with this visit. Unselor to complete the whole formation for each family sit notes and diagnostic test on, and each participating family	
Family Member Nam	ne DOB	Relationship to Patient	Phone	Number	Best Time to Contact	



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Patient Name:	DOB:	DOB:				
CLINICAL INDICATIONS (Please check all the app Please include page 3 and/or any additional clini	oly) ical information (medical records, pictures, family history) i	o aid in result interpretation.				
Perinatal history	Cardiac/Congenital Heart Malformations	Skeletal/Limb Abnormalities				
□ Prematurity	□ ASD	☐ Contractures				
☐ Intrauterine growth retardation	□ VSD	☐ Club foot				
☐ Oligohydramnios	☐ Coarctation of aorta	☐ Polydactyly				
□ Polyhydramnios	☐ Hypoplastic left heart	☐ Syndactyly				
☐ Cystic hygroma/increased NT	☐ Tetralogy of Fallot	☐ Scoliosis				
☐ Shortened fetal long bones	☐ Cardiomyopathy	☐ Vertebral anomaly				
☐ Choroid plexus cyst	☐ Arrhythmia/conduction defect	☐ Other:				
☐ Ventriculomegaly	☐ Other:	Genitourinary Abnormalities				
☐ Echogenic bowel	Skin/Hair/Nail Abnormalities	☐ Ambiguous genitalia				
☐ Fetal pyelectasis/fetal renal pelvic dilatation	☐ Abnormal nails	☐ Hypospadias				
☐ Single umbilical artery (SUA)	☐ Abnormal pigmentation	☐ Hydronephrosis				
☐ Maternal diabetes mellitus	☐ Abnormal connective tissue	☐ Undescended testis				
Growth	☐ Blistering	☐ Kidney malformation				
☐ Failure to thrive	☐ Ichthyosis	☐ Renal agenesis				
☐ Growth retardation/short stature	☐ Skin tumors/malignancies	☐ Renal tubulopathy				
☐ Overgrowth	☐ Other:	☐ Polycystic kidneys				
☐ Macrocephaly	Brain Malformations/Abnormal Imaging	☐ Multicystic kidneys				
☐ Microcephaly	☐ Agenesis of the corpus callosum	☐ Other:				
Physical/Cognitive Development	☐ Holoprosencephaly	Endocrine				
☐ Fine motor delay	☐ Lissencephaly	☐ Diabetes mellitus				
☐ Gross motor delay	☐ Cortical dysplasia	☐ Type I				
☐ Speech delay	☐ Heterotopia	☐ Type II				
☐ Intellectual disability	☐ Hydrocephalus	☐ Hypothyroidism				
☐ Learning disability	☐ Brain atrophy	☐ Hypoparathyroidism				
☐ Developmental regression	☐ Periventricular leukomalacia	☐ Pheochromocytoma/paraganglioma				
Behavioral	☐ Hemimegalencephaly	Metabolic				
☐ Autism spectrum disorder	☐ Abnormalities of basal ganglia	☐ Ketosis				
☐ Autistic features	☐ Other:	☐ Lactic acidemia/high CSF lactate				
☐ Obsessive-compulsive disorder	Neurological/Muscular	☐ Elevated pyruvate				
☐ Stereotypic behaviors	☐ Ataxia	☐ Elevated alanine				
☐ Other psychiatric symptoms	☐ Chorea	Organic aciduria				
Craniofacial/Ophthalmologic/Auditory	☐ Dystonia	Low plasma carnitine				
Cataracts	☐ Hypotonia	☐ Elevated CPK				
☐ Cleft lip/palate	☐ Hypertonia	☐ Hypoglycemia				
☐ Coloboma of eye	☐ Seizures (type:)	Hematologic/Immunologic				
☐ CPEO (ophthalmoplegia)	☐ Spasticity	Recurrent fever				
□ Ptosis	☐ Exercise intolerance/easy fatigue	☐ Anemia/neutropenia/pancytopenia				
☐ Blindness	☐ Muscle weakness	☐ Immunodeficiency type:				
Optic atrophy	☐ Stroke/stroke-like episodes	Other:				
Retinitis pigmentosis	☐ Recurrent headache/migraine	Additional Chaire Ladienties				
Hearing loss	Gastrointestinal	Additional Clinical Indications				
Ototoxicity (aminoglycoside-induced)	☐ Gastroschisis/omphalocele					
☐ External ear malformation	Pyloric stenosis					
☐ Facial dysmorphism	☐ Tracheoesophageal fistula					
Describe:	☐ Delayed gastric emptying☐ Eosinophilic esophagitis					
	☐ Gastrointestinal reflux					
	☐ Recurrent vomiting ☐ Chronic diarrhea					
	☐ Constipation					
	☐ Chronic intestinal pseudo-obstruction					
	☐ Hirschsprung disease					
	☐ Hepatic failure					
	☐ Flevated transaminases					