

PHONE 1.800.411.4363 FAX 1.800.434.9850

CONNECT





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atient Last Name	Patient First Name		MI	Date of Birth (MM / DD / YYY	
ddress	City	State		Phone	
	,		·		
cession #	BG Lab #		Biological Sex: Female Gender identity (if diffe	Male Unknown	
spital / Medical Record #	BG Family #		•		
DERING PHYSICIAN		ADDITIONAL REPORTS			
dering Physician*		Name	Na	ame	
stitution Name		Email	Er	nail	
nail (Required for International Clie	nts)	Phone	Ph	none	
none		Fax		x	
different from original order, complete Re om Individual's Records, pg. 2)	equest for and Consent to Release Information	Note: Reports will be sent by FAX except for international recipients			
AYMENT (FILL OUT ONE OF THE (ODTIONS BELOW)				
SELF PAYMENT ·····			• • • • • • • • • • • • • • • • • • • •		
Pay With Sample	Bill To Patient				
) INSTITUTIONAL BILLING					
stitution Name	Institution Code Instit	tian Cantant Nama			
	ilistitution code ilistit	cution Contact Name Ins	titution Phone	Institution Contact Email	
) INSURANCE			ititution Phone	Institution Contact Email	
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Do Not Perform Test Until P REQUIRED ITEMS 1. Copy of	Patient is Aware of Out-Of-Pocket Costs (exclude	s prenatal testing)	Physician 4. Insur	ed Signature of Authorization	
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Do Not Perform Test Until P REQUIRED ITEMS 1. Copy of me of Insured tient's Relationship to Insured dress of Insured y imary Insurance Co. Name imary Member Policy # signing below, I hereby authorize erstand that I am responsible for a sons including, but not limited to, r	Patient is Aware of Out-Of-Pocket Costs (exclude of the Front/Back of Insurance Card(s) 2. ICD10 Diagram of Birth (MM / DD / YYYY) Phone of Insured State Zip Primary Insurance Co. Phone Primary Member Group # Baylor Genetics to provide my insurance call and concovered and non-authorized services. I uni-covered and non-authorized services. I uni-	s prenatal testing) agnosis Code(s) 3. Name of Ordering Name of Insured Patient's Relationship to I Address of Insured City Secondary Insurance Co. I Secondary Member Policy rrier any information necessary, in a that the insurance policy dictates, a derstand that I am responsible for secondary in the content of	Physician 4. Insured Insured Name cluding test results as well as any amoulending Baylor Gene	ed Signature of Authorization /	
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Do Not Perform Test Until P REQUIRED ITEMS 1. Copy of the second of the	Patient is Aware of Out-Of-Pocket Costs (exclude of the Front/Back of Insurance Card(s)	s prenatal testing) agnosis Code(s) 3. Name of Ordering Name of Insured Patient's Relationship to I Address of Insured City Secondary Insurance Co. I Secondary Member Policy rrier any information necessary, in that the insurance policy dictates, a derstand that I am responsible for sare does not cover routine screenin gnature https://www.baylorgenetics.com/lab-term test is medically necessary for the risk as: treatment decisions. The person listed as	Physician 4. Insur-	ed Signature of Authorization /	

Physician's Printed Name

Physician's Signature

Date (MM / DD / YYYY)



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REQUEST FOR AND CONSENT TO RELEASE OF INFORMATION FROM INDIVIDUAL'S RECORDSPlease complete if reanalysis is being requested by a different provider than original ordered genome.

BACKOROUND IN ORMATION			
NOTE: The execution of this form does not author	orize the release of information other than that spec	cifically described below. This form authorizes th	e release of information that you specify in
accordance with 5 U.S.C., Section 5701 and 7332	2; and 45 C.F.R., parts 160 and 164.		
			//
Individual/Patient Last Name	Individual/Patient First Name	MI	Date of Birth (MM / DD / YYYY)
BG Lab #	BG Family #	Ordering Physician Phone	Ordering Physician Fax
Individual or Organization's Name to Whom Ir	nformation is Being Released		
Address		City	State Zip
,		o.v,	2.5
Information Requested:		Purpose(s) or need for which information is	to be used by Organization of Individual
		to whom information is to be released:	
I want my original provider to receive the res	ults of the analysis: Yes	○ No	
want my originat provider to receive the res	und of the unutysis.	<u> </u>	
AUTHORIZATION AND CERTIFICATION			
I certify that this request has been made freely,	voluntarily, and without coercion and that the inform	mation given above is accurate and complete to tl	ne best of my knowledge. I understand this
release may not be obtained or offered as condi	tion for treatment, payment, or other eligibility for I	benefits upon my signing this authorization. I may	y revoke this authorization at any time in
writing, except to the extent that this action has	already been taken to comply with it. Written revoc	ation is effective upon receipt by the facility hous	ing the records. Upon release, my records
will no longer be protected, and re-disclosure by	y those receiving the information may be accomplis	shed without my further authorization. Without m	y express revocation, the authorization will
automatically expire upon satisfaction of the nec	ed for disclosure, under the conditions listed below	, or upon this date (sup	oplied by individual/patient).
			/ /
Individual/Patient Signature			,,,,
			/ /
Personal Representative Signature, if not sign	ned by patient*		Date (MM / DD / YYYY)

*[NOTE: ATTACH DOCUMENTS DEMONSTRATING YOUR AUTHORITY TO ACT ON BEHALF OF THE PATIENT.] PLEASE FAX COMPLETED FORM TO: 713.798.2787

OP.FR 6 Authorization For Release of Protected Health Information



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GENOME REANALYSIS REQUISITION

Patient Last Na	ame Patient First Nan	me MI Date of	Birth (MM / DD / YYYY) Biological Sex		
GENOME REA	NALYSIS TEST OPTIONS	PHENOTYPE INFORMATION (REQUIRED)			
1897	Genome Reanalysis	YES Phenotype has changed and I have included (Please complete the "Indication for Testing"	an updated clinical note and/or other documentation. 'section.)		
		NO Phenotype has NOT changed. I have reviewe summary section of the WGS report and I ag	ed the phenotype information included in the clinical gree this is currently accurate information.		
OPT-IN TESTI	NG OPTIONS				
Opt-in for RNA	Sequencing (RNASeq) as reflex to genome real	nalvsis			
	Reanalysis identifies a qualified variant that m from this patient to perform RNAseq.	night be reclassified through RNA Sequencing, please notify	me about this variant. A new blood sample will need to be		
INDICATION F	OR TESTING (REQUIRED IF SELECTED "YES	S" FOR THE "PHENOTYPE INFORMATION" SECTION.)			
number (http://h		o be tested. Please also submit a clinic note and pedigree, if available is needed to facilitate interpretation of whole genome sequencing re	t. Phenotypes listed are in HPO terms with the corresponding HPO ssults. If the laboratory requires additional information, please indicate		
Physician Nam	e	Physician Phone ICD-10 I	Diagnosis Code(s)		
PRE/PERINA	TAL HISTORY ·····	EYE DEFECTS & VISION	MOTOR/COGNITIVE DEVELOPMENT		
0001360 0001339 0002084 0000238 0002119 0001273 0002539 0012444 00002352	Prematurity - GA at birth Intrauterine Growth Restrictions Oligohydramnios Polyhydramnios Cystic Hygroma Congenital Diaphragmatic Hernia Failure to Thrive Omphalocele Encephalocele Increased Nuchal Translucency BRAIN ABNORMALITIES Holoprosencephaly Lissencephaly Encephalocele Hydrocephalus Ventriculomegaly Abnormality of Corpus Callosum Cortical Dysplasia Brain Atrophy Leukoencephalopathy	0000505	□ 0000750 Delayed Speech & Language Development □ 0001270 Delayed Motor Milestones □ 0002376 Developmental Regression □ Intellectual Disability □ 0001256 □ 00002342 Moderate □ 0010864 Severe □ 0000729 Autistic Spectrum Disorder □ 0000729 Autistic Spectrum Disorder □ 0000256 Macrocephaly □ 0000252 Microcephaly □ 0001363 Craniosynostosis □ 0000204 Cleft Upper Lip □ 0000175 Cleft Palate □ 0000316 Hypertelorism □ 00008050 Abnormality of the Palpebral Fissures □ 0000286 Epicanthal Folds		
 □ 0002269 □ 0002126 □ 0001302 □ 0002500 □ 0007266 □ 0006808 □ 0002134 □ 0002363 □ 0007360 □ 0006817 	Abnormality of Neuronal Migration Polymicrogyria Pachgyria Abnormality of Cerebral White Matter Cerebral Dysmyelination Cerebral Hypomyelination Abnormality of the Basal Ganglia Abnormality of the Brainstem Aplasia/Hypoplasia of the Cerebellum Aplasia/Hypoplasia of the Cerebellar Vermis	□ 0010819 Atonic Seizures □ 0002121 Absence Seizures □ 0011169 Generalized Clonic Seizures □ 0001251 Ataxia □ 0001332 Dystonia □ 0002072 Chorea □ 0001257 Spasticity □ 0009830 Neuropathy	O000288 Abnormality of the Philtrum O010938 Abnormality of the External Nose		



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GENOME REANALYSIS REQUISITION

Patient Last Na	me Patient First Na	ume		Date of Birth (MM / DD / YY	YY) Biological Sex
		iiiie	IVII	Date of Biltil (MM) / BB / 11	Try Biological Sex
INDICATION F	OR TESTING (REQUIRED) - CONTINUED				
HAIR & SKIN		CARDIAC		····· GENITOURIN	ARY
0000957	Cafe-Au-Lait Spots	0001631	Atria Septal Defect	0000113	Polycystic Kidney Dysplasia
0001034	Hypermelanotic Macule	0001629	Ventricular Septal Defect	0000107	Renal Cyst
0001010	Hypopigmentation of the Skin	0001655	Patent Foramen Ovale	0008738	Partially Duplicated Kidney
0008066	Abnormal Blistering of the Skin			0000104	Renal Agenesis
0008064	Ichthyosis	0001713	Abnormality of Cardiac Ventricle	0000085	Horseshoe Kidney
0000988	Skin Rash Recurrent Skin Infections	0001636	Tetralogy of Fallot		Abnormality of the Ureter
0005306	Capillary Hemangiomas	0001680	Coarctation of Aorta	0000795	Abnormality of the Urethra
0001597	Abnormality of the Nail	0001647	Bicuspid Aortic Valve	0000047	Hypospadias
0004554	Generalized Hypertrichosis	0002616	Aortic Root Dilatation	0000028	
0001596	Alopecia	0001638	Cardiomyopathy		Cryptorchidism
0002208	Coarse Hair	0011675	Arrhythmia		Abnormality of the Testis
0002299	Brittle Hair		Arriyanina		Ambiguous Genitalia
		<u> </u>			
		Ш		L	
RESPIRATOR	γ	METABOLIC		····· MUSCULOSK	ELETAL
0002093	Respiratory Insufficiency	0001946	Ketosis	0011398	Hypotonia
0002878	Respiratory Failure	0003074	Hyperglycemia	0001276	Hypertonia
0002104	Apnea	0001943	Hypoglycemia	0000098	Tall Stature
0002791	Hypoventilation	0001941	Acidosis	0004322	Short Stature
0002883	Hyperventilation	0003128	Lactic Acidosis	0001382	Joint Hypermobility
0002788	Recurrent Upper Respiratory Tract	0003215	Dicarboxylic Aciduria		Flexion Contracture
	Infections	0002490	Increased CSF lactate	0002804	Arthrogryposis Multiplex Congenita
H		_		0001161 0001829	Hand Polydactly
L		0001992	Organic Aciduria	0001829	Foot Polydactly
		0030085	Abnormal CSF Lactate Level	0008101	Finger Syndactly Toe Syndactly
GASTROINTE	STINAL	00003542	2 Increased Serum Pyruvate	0100490	Camptodactyly of Finger
		0003535	3-Methylglutaconic aciduria	0012165	Oligodactyly
0002021	Pyloric Stenosis Tracheoesophogeal Fistula	0001942	Metabolic acidosis	0001762	Talipes Equinovarus
0002373	Esophageal Atresia	0100493	Hypoammonemia	0002757	Recurrent Fractures
0002020	Gastroesophageal Reflux	0001987	Hyperammonemia	0002650	Scoliosis
0001733	Pancreatitis	0004923	Hyperphenylalaninemia	0002808	Kyphosis
0002014	Diarrhea	0003234	Decreased Plasma Carnitine	0003307	Hyperlordosis
0002019	Constipation	0003236	Elevated Serum Creatine	0001528	Hemihypertrophy
0002037	Inflammatory Bowel Disease	_	Phosphokinase	0001513	Obesity
0004389	Intestinal Pseudo-Obstruction	_	l Newborn Screen	0001548	Overgrowth
0001399	Hepatic Failure	Unusual (Color/Odor	0002652	Skeletal Dysplasia
0002572	Episodic Vomiting				
0001744	Splenomegaly				
0002240	Hepatomegaly				
0001508	Postnatal Failure to Thrive				
	Gastroparesis				



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GENOME REANALYSIS REQUISITION

			/ /	
Patient Last Name Patien	t First Name	МІ	Date of Birth (MM / DD / YYYY)	Biological Sex
INDICATION FOR TESTING (REQUIRED) - CONTI	NUED			
ENDOCRINE	····· HEMATOLOGY ····		OTHER	
O000819 Diabetes Mellitus O000873 Diabetes Insipidus Hypothyroidism O000829 Hypoparathyroidism O000834 Abnormality of the Adrenal Glands Exocrine Pancreatic Insufficiency O002721 Immunodeficiency	Chronic Cyclic O001873 Throm O040185 Macrot O005537 Decrea O005518 Erythro O004444 Sphero O012410 Pure R	penia Congenital bocytopenia chrombocytopenia used Mean Platelet Volu ocyte Macrocytosis ocytosis ed Cell Aplasia	0001954	maglobulinemia I Immunoglobulins eficiency I urinary odor erance Ily lax or hyperextensible skin
EAR DEFECTS & HEARING 0000407 Sensorineural Hearing Impairment 0000405 Conductive Hearing Impairment 0000410 Mixed Hearing Impairment 00004467 Preauricular Pit 0000384 Preauricular Skin Tag 0000369 Low-set Ears 000037 Abnormality of the Pinna	CANCER		atives	ilar Disorder

Consent on next page



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GENOME REANALYSIS REQUISITION

			//	/		
Patient Last Name	Patient First Name	MI	Date of Birth	(MM / DD / YYYY)	Biolo	gical Sex
INFORMATION AND CONSEN	T FOR TESTING					
reported, will be interpreted widiagnosis to the patients. Variar	evious whole genome sequencing are a th current knowledge, and updated cli nts in the original report may be remo nosis for the patient. The healthcare p of genetic counseling.	nical indications if provided. (ved in the updated report due	Clinical report will be issued to variant re-classification i	listing variants that ar nto benign/likely benig	e highly likel gn, or variant	y to provide no longer
Patient's preference of the med order.	dical actionable findings and carrier fir	ndings is presumed to be unc	hanged, unless indicated in a	new consent form sul	bmitted with	the reanalysis
					/	/
Printed Name		Signature			Date (MM	1 / DD / YYYY)
					/	/
Relationship to Patient		Proband Name			Proband D	OB (MM/DD/YY)
					/	/
Physician's/Counselor's Signati	ure				Date (MM	1 / DD / YYYY)