

PHONE 1.800.411.4363 FAX 1.800.434.9850







PATIENT INFORMATION (COMPLETE ON	IE FORM FOR EACH PERSON TESTED)				
Patient Last Name	Patient First Name		MI		Date of Birth (MM / DD / YYYY)
Address	City		State Biological Sex:	Zip	Phone
Accession #	Hospital / Medical Record	#	Female Gender identity (if di	Male fferent from above):	Unknown
REPORTING RECIPIENTS					
Ordering Physician		Institution Name			
Email (Required for International Clients)		Phone		Fax	
ADDITIONAL RECIPIENTS					
Name	<u> </u>	Email		Fax	
Name		Email		Fax	
PAYMENT (FILL OUT ONE OF THE OPTION	ONS BELOW)				
SELF PAYMENT					
	o Patient				
() INSTITUTIONAL BILLING					
O INSTITUTIONAL BILLING					
Institution Name	Institution Code Inst	itution Contact Name	Institution F	Phone	Institution Contact Email
O INSURANCE					
☐ Do Not Perform Test Until Patien	t is Aware of Out-Of-Pocket Costs (exclud	les prenatal testing)			
REQUIRED ITEMS 1. Copy of the I	Front/Back of Insurance Card(s) 2. ICD10 D	Diagnosis Code(s) 3. Name	of Ordering Physician	4. Insured Sign	ature of Authorization
	/ /	_			_ / /
Name of Insured	Insured Date of Birth (MM / DD / YYYY)	Name of Insure	d	Insure	d Date of Birth (MM / DD / YYYY)
Patient's Relationship to Insured	Phone of Insured	Patient's Relation	onship to Insured	Phone	of Insured
Address of Insured		Address of Insu	red		
		_ :			
City	State Zip	City		State	Zip
Primary Insurance Co. Name	Primary Insurance Co. Phone	Secondary Insu	rance Co. Name	Secon	dary Insurance Co. Phone
Primary Member Policy #	Primary Member Group #	Secondary Mem	nber Policy #	Secon	dary Member Group #
By signing below, I hereby authorize Bay understand that I am responsible for any or reasons including, but not limited to, non- directly from my insurance company in pa	o-pay, co-insurance, and unmet deductib covered and non-authorized services. I u	ole that the insurance policy understand that I am respo	/ dictates, as well a nsible for sending l	s any amounts no Baylor Genetics ar	t paid by my insurance carrier for
					//
Patient's Printed Name	Patient's S	Signature			Date (MM / DD / YYYY)
STATEMENT OF MEDICAL NECESSITY (REQUIRED)				
This test is medically necessary for the ris patient's medical management and treatm provided genetic testing information to the	nent decisions. The person listed as the O	Ordering Physician is author			
					/ /
Physician's Printed Name	Physician	's Signature			Date (MM / DD / YYYY)



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Patient Last Name	Patient First Name	MI	Date of Birth (M	MM / DD / YYYY)	Biological Sex
INSTRUCTIONS FOR ORDERING					
	ered together allow for the most comprehensive a mal Microarray Analysis, mtDNA Analysis or Globa nd WES.				
TRIO WES TEST OPTIONS					
 1600 Trio Whole Exome Sequencing 1532 Trio Whole Exome Sequencing 1722 Critical Trio Whole Exome Sequencing 1533 Critical Trio Whole Exome Sequencing 	+ Comprehensive mtDNA Analysis	(Both Parents	·		
PROBAND WES TEST OPTIONS					
(CMA) (Comprehensive)	ncing + Chromosomal Microarray Analysis		ING PARENTAL TESTS Optional Parental Cont		NOTE: See consent for for the control will be used.
DUO WES TEST OPTIONS					
☐ 1603 Duo Whole Exome Sequencing ☐ 1723 Rapid Duo Whole Exome Sequencing		1550 I	ING PARENTAL TESTS (On Parental WES - Materr Parental WES - Patern Other Relative	nal	NOTE: Please use separate Additional Affected Sibling for Trio requisition for additional family members.
GLOBAL MAPS® TESTS		ADD-ON TE	STS		
Global Metabolomic Assiste Was plasma extracted from Global Metabolomic Assiste	0 0	2055	Chromosomal Microa Comprehensive mtDN Exome Raw Data Rele	NA analysis by NGS	SNP Screen (Comprehensive)
PROBAND SAMPLE(S)					
Please refer to www.baylorgenetics.com for Blood in EDTA Buccal Swab Cord Blood (Call lab for sample specification)	Cultured Skin Fibroblast Extracted DNA from	mt DNA anal Skeleta Liver Tissue	l Muscle (Global MAPS only Plasma from EDTA // Date of Collection MM / DD / YYYY)	Urine
NOTE: Extracted DNA/RNA will only be accepted if the	isolation of nucleic acids for clinical testing occurs in a CLIA-cer	rtified laboratory o	r a laboratory meeting equiv	valent requirements as determin	ed by the CAP and/or the CMS.
BIOLOGICAL PARENTS INFORMATION					
	FOR TRIO WES; Other family members cannot be substitute uthorization on consent.	ed for either pare	ent. Be sure to label parer	ntal samples with full name a	nd date of birth - DO NOT LABEL
MATERNAL INFORMATION		PATERNAI	L INFORMATION		
Asymptomatic Symptomatic (a	Attach summary of findings)	Asymp	otomatic Symp	otomatic (Attach summary o	f findings)
Maternal Last Name Mat	rernal First Name MI	Paternal La	ıst Name	Paternal First Nam	e MI
Maternal Date of Birth (MM / DD / YYYY) Date of Collection	Sample Type: Blood Buccal swab	Paternal Da (MM / DD / '	YYYY)	//	Sample Type: Blood Buccal swab
(MM / DD / YYYY)/	·	(MM / DD / '	YYYY)		



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WES ADVANTAGE REQUISITION

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Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Biological Sex
ITEM CHECKLIST FOR TESTING				
Proband Sample (Required)	Signed WES Cons	ent Form	Indication for	Study
Maternal Sample (Required for Trio)	Clinical Note/Sun			•
Paternal Sample (Required for Trio)	Requisition	,		
INDICATION FOR TESTING (REQUIRED) Please provide the following clinical information reg- number (http://human-phenotype-ontology.github.io the health care provider to be contacted:				
Physician Name	Physician Phone		ICD-10 Diagnosis Code(s)	
PRE/PERINATAL HISTORY	EYE DEFECTS & VIS	ION	MOTOR/COGNIT	VE DEVELOPMENT
0001622 Prematurity - GA at birth	0000505 Visua	l Impairment	0000750 De	elayed Speech & Language Development
0001511 Intrauterine Growth Restricti	ons 0000618 Blinds	ness	0001270 De	elayed Motor Milestones
0001562 Oligohydramnios	0000589 Colob	oma	0002376 De	evelopmental Regression
0001561 Polyhydramnios	0000526 Anirio	lia	☐ Intellectual Di	sability
0000476 Cystic Hygroma		nthalmia	000125	6 Mild
0000776 Congenital Diaphragmatic He		phthalmia	000234	2 Moderate
0001508 Failure to Thrive	0000508 Ptosis		001086	4 Severe
0001539 Omphalocele		ismus		itistic Spectrum Disorder
0002084 Encephalocele		act Congenital Bilateral		and open an District
0010880 Increased Nuchal Translucer	□			
<u> </u>	⊔ <u></u>		⊔	
STRUCTURAL BRAIN ABNORMALITIES	NEUROLOGICAL		····· CRANIOFACIAL	
0001360 Holoprosencephaly	0001284 Arefle	wia.		acrocephaly
0001339 Lissencephaly				icrocephaly
		tic Encephalopathy		. ,
0002084 Encephalocele	0001250 Seizu	res		raniosynostosis
0000238 Hydrocephalus	0002373	Febrile Seizures	=	eft Upper Lip
0002119 Ventriculomegaly	—	Infantile Spasms		eft Palate
0001273 Abnormality of Corpus Callos	o002123	Generalized Myoclonic		/pertelorism
0002539 Cortical Dysplasia		Seizures		/potelorism
0012444 Brain Atrophy	0002069	Generalized Tonic-clonic Seizures	☐ 0008050 AI	onormality of the Palpebral Fissures
0002352 Leukoencephalopathy				oicanthal Folds
0002269 Abnormality of Neuronal Mig		Generalized Tonic Seizure	0000288 AI	onormality of the Philtrum
0002126 Polymicrogyria	0010819	Atonic Seizures	☐ 0010938 A	onormality of the External Nose
0001302 Pachgyria	0002121	Absence Seizures		
0002500 Abnormality of Cerebral White	te Matter 0011169	Generalized Clonic Seizur	es	
0007266 Cerebral Dysmyelination	0001251	Ataxia		
0006808 Cerebral Hypomyelination	O001332	Dystonia		
0002134 Abnormality of the Basal Gar	nglia 0002072	Chorea		
0002363 Abnormality of the Brainsten		Spasticity		
0007360 Aplasia/Hypoplasia of the Ce	rehellum			
0006817 Aplasia/Hypoplasia of the Ce		Neuropathy		
Vermis	닏			
LI				

Indications continued on next page



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INDICATION F	OR TESTING (REQUIRED) - CONTINUED					
HAIR & SKIN		CARDIAC		G	ENITOURIN	ARY
0000957	Cafe-Au-Lait Spots	0001631	Atria Septal Defect	Γ	0000113	Polycystic Kidney Dysplasia
0001034	Hypermelanotic Macule	0001629	Ventricular Septal Defect	Γ	0000107	Renal Cyst
0001010	Hypopigmentation of the Skin		·	-	0008738	Partially Duplicated Kidney
0008066	Abnormal Blistering of the Skin	0001655	Patent Foramen Ovale		0000730	Renal Agenesis
0008064	Ichthyosis	0001713	Abnormality of Cardiac Ventric	cle L	_	•
0000988	Skin Rash	0001636	Tetralogy of Fallot	L	0000085	Horseshoe Kidney
0001581	Recurrent Skin Infections	0001680	Coarctation of Aorta	L	0000069	Abnormality of the Ureter
0005306	Capillary Hemangiomas	0001647	Bicuspid Aortic Valve	L	0000795	Abnormality of the Urethra
0001597	Abnormality of the Nail	0002616	Aortic Root Dilatation		0000047	Hypospadias
0004554	Generalized Hypertrichosis				0000028	Cryptorchidism
0001596	Alopecia Coarse Hair	0001638	Cardiomyopathy		0000035	Abnormality of the Testis
0002208	Brittle Hair	0011675	Arrhythmia	[0000062	Ambiguous Genitalia
	Brittle Hall					
<u> </u>						
ш		<u> </u>				
RESPIRATOR	γ	METABOLIC		N	IUSCULOSK	ELETAL
0002093	Respiratory Insufficiency	0001946	Ketosis		0011398	Hypotonia
0002878	Respiratory Failure	0003074	Hyperglycemia		0001276	Hypertonia
	Apnea	0001943	Hypoglycemia	[0000098	Tall Stature
0002791	Hypoventilation	0001941	Acidosis		0004322	Short Stature
_					0001382	Joint Hypermobility
0002883	Hyperventilation Recurrent Upper Respiratory Tract	0003128	Lactic Acidosis		0001371	Flexion Contracture
0002788	Infections	0003215	Dicarboxylic Aciduria		0002804	Arthrogryposis Multiplex Congenita
Ш		0002490	Increased CSF lactate		0001161	Hand Polydactly
		0001992	Organic Aciduria	[0001829	Foot Polydactly
		0030085	Abnormal CSF Lactate Level	[0006101	Finger Syndactly
		00003542	! Increased Serum Pyruvate	[0001770	Toe Syndactly
GASTROINTE	STINAL	0003535	3-Methylglutaconic aciduria	اِ	0100490	Camptodactyly of Finger
0002021	Pyloric Stenosis	0001942	Metabolic acidosis	Ĺ	0012165	Oligodactyly
0002575	Tracheoesophogeal Fistula			Ĺ	0001762	Talipes Equinovarus
0002032	Esophageal Atresia	0100493	Hypoammonemia 	L	0002757	Recurrent Fractures
0002020	Gastroesophageal Reflux	0001987	Hyperammonemia	L	0002650	Scoliosis
0001733	Pancreatitis	0004923	Hyperphenylalaninemia	Ĺ	0002808	Kyphosis
0002014	Diarrhea	0003234	Decreased Plasma Carnitine	Ĺ	0003307	Hyperlordosis
0002019	Constipation	0003236	Elevated Serum Creatine Phosphokinase	L	0001528	Hemihypertrophy
0002037	Inflammatory Bowel Disease Intestinal Pseudo-Obstruction	Abnormal	Newborn Screen	L	0001513	Obesity
0004389	Hepatic Failure	Unusual C	Color/Odor	L	0001548	Overgrowth
0001377	Episodic Vomiting			L	0002652 □	Skeletal Dysplasia
0001744	Splenomegaly	<u> </u>		[┦───	
0002240	Hepatomegaly	Ш				
0001508	Postnatal Failure to Thrive					
0002578	Gastroparesis					



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INDICATION F	OR TESTING (REQUIRED) - CONTINUED				
ENDOCRINE		HEMATOLOGY ·····	0	THER	
0000819 0000873 0000821 0000829 0000834 0001738 0002721	Diabetes Mellitus Diabetes Insipidus Hypothyroidism Hypoparathyroidism Abnormality of the Adrenal Glands Exocrine Pancreatic Insufficiency Immunodeficiency	0001875 Neutropenia 0005549 Congenital Chronic Cyclic 0001873 Thrombocytopenia 0040185 Macrothrombocyt 0005537 Decreased Mean F 0005518 Erythrocyte Macro 0004444 Spherocytosis 0012410 Pure Red Cell Apla	ppenia [Platelet Volume [pcytosis [0001954 Episodic Fer 0004313 Hypogamm 0010701 Abnormal Ir 0002721 Immunodefi 0012088 Abnormal u 0012537 Food intoler	aglobulinemia nmunoglobulins iciency rinary odor
EAR DEFECTS 0000407 0000 0000405	S & HEARING Sensorineural Hearing Impairment 8619 Bilateral Conductive Hearing Impairment	Aplastic Hypoplastic 0001903 Anemia 0005528 Bone Marrow Hyp		Abnormal Movements Family History of Simila 0001254 Lethargy 0002415 Leukodystro	r Disorder
 □ 0000410 Mixed Hearing Impairment □ 0004467 Preauricular Pit □ 0000384 Preauricular Skin Tag □ 0000369 Low-set Ears □ 000037 Abnormality of the Pinna 		CANCER Type of Cancer Age of Diagnosis Family History of Cancer and		ENES OF INTEREST -	
ADDITIONAL C	CLINICAL INFORMATION	DIF	FERENTIAL DIAGNOSIS		

Consent on next page



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INFORMATION AND CONSENT FOR TESTING

DESCRIPTION OF WHOLE EXOME SEQUENCING TEST

The WES test is a highly complex test that is developed to identify changes in an individual's DNA that cause or contribute to their medical concerns. The exome refers to-the portion of the human genome that contains functionally important sequences of DNA that direct the body to make proteins essential for the body to work properly. These regions of DNA are called exons. It is known that most of the errors that occur in DNA sequences that then lead to genetic disorders are located in the exons. In contrast to other sequencing tests that analyze one gene or small groups of related genes at a time, WES will analyze the important regions of tens of thousands of genes at the same time. Therefore, sequencing of the exome is thought to be an efficient method of analyzing a person's DNA to discover the underlying genetic cause of diseases or disabilities. However, it is possible that even if WES identifies the underlying genetic cause for the disorder in your family this information may not help in predicting prognosis or change medical management or treatment of disease.

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TESTING REPORTING

When your exome sequence is compared to a normal reference sequence, many variations or differences are expected to be found. Based on currently available medical and scientific information, we will decide whether any of these variations are predicted to be causative or related to your medical concerns. The report will contain results that may explain the cause of your current medical problems. It may also contain information about genes and diseases that have clear and immediate medical significance to your health or the health of family members, whether or not they relate to your current symptoms.

You may receive any of the following types of results:

- · Positive: Positive or "abnormal" results mean there is a change in the genetic material related to your medical issues.
- Negative: Negative or "normal" results mean no relevant genetic change could be detected using WES. This does not mean there is no genetic change, but it may
 mean that WES could not detect it.
- Results of Unclear Significance: WES can detect change(s) in DNA that do not have clear meaning. These alterations are also referred to to as variants of uncertain significance (VUS). Additional studies may be indicated if a VUS is identified in a gene that may be associated with your medical concerns.
- SecondaryFindings: WES testing can sometimes detect a change in a person's DNA unrelated to the reason for testing. If this change has medical significance, it is called a secondary finding.

SECONDARY FINDINGS

You have the choice to OPT-IN or OPT-OUT of the following category of secondary findings:

Category I: Medically Actionable

The report may also contain information regarding genes and diseases that are considered medically actionable because they have clear and immediate medical significance to your health or the health of family members, whether or not they relate to your current symptoms. The American College of Medical Genetics (ACMG) has published guidelines for the reporting of theses types of medically actionable or secondary findings (PMID: 34012068). These guidelines include a list of genes (updated periodically) that are considered medically actionable and thus, laboratories should seek and report pathogenic (disease causing) findings in these genes. In accordance with an update to this policy statement (PMID: 25356965), there is the choice to opt-out of recieving this information.

ADDITIONAL REPORTING

The report will NOT include findings in genes causing adult onset dementia syndromes for which there is presently no prevention or cure. If the reason for testing includes features that clearly indicate such a disorder, we recommend pursuing targeted testing based on specific symptoms and not WES testing. However, if the reason for testing includes a clinical presentation that could include such a disorder or a mixed neurological phenotypes, then results may be reported in the proband (patient) and the parents for genes that have an allelic association with dementia or is a component of the phenotype.

The interpretation of the variants is based on information available at the time of testing and may change in the future as medical knowledge advances. As determined necessary by the laboratory, the proband's sample will have the findings confirmed by a second methodology (Sanger sequencing). We expect to find hundreds of variations when comparing the DNA to the reference sequence, Most of these do not relate to disease and therefore, will not be reported. The raw sequence data generated by WES is available for request once a WES report has been issued. Pleasde see our website for further information regarding this.

Additional reporting for Trio WES (test codes 1600, 1722, 1532, 1533): As part of the Trio WES analysis, we will report findings in genes that have occurred in the affected individual, but not in the asymptomatic parents. This category of results caused by new (de novo) findings may be significant in determining the cause of your medical condition. Thus, this category of changes will be reported for genes with or without a known current association with disease. We will also report compound heterozygous, homozygous and hemizygous variants in genes where each parent has one change and the affected individual has inherited both changes, for genes with or without a known association with disease. It is important to note that the Trio WES report may contain information about diseases and genes that do not relate to your current condition, or may develop many years from now, or do not have any known link to disease, according to current knowledge. As part of the Trio WES test, blood samples from the biological parents of the proband are required. Trio WES will be performed on the proband and parental samples at the same time and the sequence data will be analyzed in the context of the family relationships. The parental data will be used to help interpret the proband's data. Custom Family Sequence Analysis (test code 1580) is available for family members at an additional charge. Free testing for variants of unknown significance is available with prior approval. A separate parental report will be issued regarding the two categories of secondary findings.

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INFORMATION AND CONSENT FOR TESTING

Additional reporting for Duo WES (test codes 1603, 1623): As part of the Duo WES analysis, we will report findings in genes that have occurred in the affected individual, but not in the asymptomatic parent. This category of results caused by new (de novo) findings may be significant in determining the cause of your medical condition. Thus, this category of changes will be reported for genes with or without a known current association with disease. We will also report compound heterozygous, homozygous, and hemizygous variants in genes where the parent has one change and the affected individual has inherited the change, for genes with or without a known association with disease. It is important to note that the Duo WES report may contain information about diseases and genes that do not relate to your current condition, or may develop many years from now, or do not have any known link to disease, according to current knowledge. As part of the Duo WES test, blood samples from the biological parent of the proband are required. Duo WES will be performed on the proband and parental sample at the same time and the sequence data will be analyzed in the context of the family relationships. The parental data will be used to help interpret the proband's data. Custom Family Sequence Analysis (test code 1580) is available for family members at an additional charge. Free testing for variants of unknown significance is available with prior approval. A separate parental report will be issued regarding the category of secondary findings.

Additional reporting for Proband WES (test codes 1500, 1530,1531): We will also include variants in possible candidate disease genes that might potentially contribute to patient phenotype. Further research studies are needed to clarify the clinical relevance of those variants/genes. Biological parental samples may help facilitate interpretation of Proband WES results. After the proband report is issued, the parental samples received will be tested by whole exome sequencing (test code 1551) for the entire exome, or will be tested by targeted methods such as Sanger sequencing (test code 1580) for changes in genes that are highly likely to be causative of disease (related to patient indication for testing) to confirm mode of inheritance, de novo status, etc. as determined necessary by the laboratory Additionally, if opted-in to receive medically actionable findings, this information will be issued in a separate parental report. Testing of parental status will ONLY be initiated if there is a variant identified in the proband. For targeted testing on the variants detected in the proband's exome data, test code 1580 is available for all family members. Free testing for variants of unknown significance in the immediate family members is available if approved by Baylor Genetics.

Your physician may order additional tests along with WES. Further test code specific information is as follows:

Test codes 1531, 1532 and 1533: In addition to WES analysis as detailed above, this order will also include a separate analysis of the mitochondrial DNA.

Test code 2055: This is the evaluation of the entire mitochondrial genome for point mutations and deletions. This will be reported separately from the WES results with a turnaround time of 14-28 days. If an mtDNA change is identified, the report will indicate recommendations for familial follow-up. Baylor Genetics will NOT automatically initiate testing on the maternal sample. If this is desired, please contact client services for assistance.

Test code 1530: This order will also include a separate analysis for detection of deletions and duplications plus a screen for detection of uniparental disomy (UPD) andabsence of heterozygosity (AOH).

Test code 8665: This will be reported separately from the WES results with a turnaround time of 14 days. If a copy number change is identified, the report will indicate recommendations for familial follow-up. Baylor Genetics will NOT automatically initiate testing on the parental smple(s). If this is desired, please contact client services for assistance.

Test codes 4900 and 4901 (Global MAPS): This is a large scale, semi-quantitative screening test that looks at changes in both individual analytes and pathways related to biochemical abnormalities, including (but not limited to) amino acid, organic acid, lipid and nucleotide metabolism. It should be used as a screening tool for individuals who have an undifferentiated phenotype or as supportive evidence in individuals with equivocal mutations in genes related to metabolic processes. It is not intended to supplant current diagnostic testing for specific conditions, nor is it intended for monitoring therapy. Any abnormalities detected using Global MAPS should be confirmed by diagnostic biochemical or molecular diagnostic testing. Consent for testing below is for WES and does not need to be completed if only Chromosomal Microarray Analysis, mtDNA Analysis or Global MAPS is ordered. Please visit our website for further information about these tests.

POTENTIAL RISKS, LIMITATIONS, AND DISCOMFORTS

- 1. It is possible that you could have a variant in a gene included in the WES test, but the WES test was unable to detect the variant. Therefore, it is possible that you may be affected with one of the conditions tested by WES, but that the test did not detect the condition.
- 2. The WES test does not analyze 100% of the genes in the human genome. There are some genes that cannot be included in the test due to technical reasons.
- 3. Results may be unclear or indicate the need for further testing on other family members.
- 4. It is possible that additional information may come to light during these studies regarding family relationships. For example, data may suggest that family relationships are not as reported, such as non-paternity (the father of the individual is not the biological father) or consanguinity (marriage or reproductive partners are blood relatives). Since the accurate assignment of family relationships is critical to the analysis of WES, we may perform a separate genetic test to confim that the samples that were submitted from the parents were correctely identified. If a discrepancy is identified, we will proceed with testing for the individual(s) who are correctly identified.
- 5. If you sign the consent form, but you no longer wish to have your samples tested by WES, you can contact your doctor to cancel the test. If testing is complete, but you have not received your results yet, you can inform your doctor that you no longer wish to recieve the results. However, if you withdraw consent for testing after 5p.m. CST, the next business day following sample receipt by the laboratory, you will be charged for the full cost of the test.
- 6. Information including results, indications for testing and clinical status obtained from the WES test may be shared with health care providers, scientists and health care databases or used in scientific publications or presentations, but the personal identifying information of all persons studied will not be revealed in such data sharing or publications/presentations.
- 7. Variants identified by WES may also be submitted to public databases, such as ClinVar, to contribute knowledge to the medical profession. Usually limited clinical information is also required for the submission. However, it is unlikely that contents of the database submissions will include any information that will identify you personally.
- 8. Due to the fact that many different genes and conditions are being analyzed, there is a risk that you will learn genetic information about yourself or your family that is not directly related to the reason for ordering the WES. This information might relate to diseases with symptoms that may develop in the future in yourself or other family members as well as conditions that have no current treatment.
- 9. It is possible that even if WES identifies the underlying genetic cause for the disorder in your family, this information may not help in predicting prognosis or change management or treatment of disease.



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WES ADVANTAGE REQUISITION

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Patient Las	st Name	:	Patient First Nan	ne	MI	Date of Birth (MM / DD / YYYY)	Biol	ogical Sex
INFORMAT	TION A	ND CONS	SENT FOR TESTING					
PROBAND	REPO	RTING	PTIONS AND AUTHORIZATION	•••••				
			atements carefully and check th			e nature of the methodology of this to	esting we are u	nable to guaran
						ported under Sequential Trio Whole pand WES protocol in the consent be		cing (test code
For Option	s 1: If n	either bo	ox is checked, or if form is not sign	ned, the lab will de	fault to the NO/ do not i	report option.		
INITIAL	1.	MEDICA	LLY ACTIONABLE					
			nic variants in genes included ir d as medically actionable on the		statement regarding	recommendations for reporting of ir	cidental findin	gs will be
	\bigcirc	YES	Please report pathogenic va	ariants in genes de	etermined to be medic	cally actionable by the ACMG policy s	tatement.	
	\bigcirc	NO	Please do NOT report patho	genic variants in g	genes included in the A	ACMG policy statement.		
For option	2: if ne	ither box	is checked, or the form is not sig	ned, the lab will de	fault to the YES/ releas	se updated report option.		
INITIAL	2.	OPTION 1	O ALLOW RELEASE OF UPDATED R	ESULTS				
		diagnosi	s can be made with this informa	ition we would like	e to issue an updated	ng the significance of changes in a p report to the physician who ordered NOT include a complete review of all	your WES test.	
	\bigcirc		f new information is known rega vould like for you to issue an upo			n that may not have previously been ed this WES testing.	included in my	WES report I
	\bigcirc		Please do NOT issue an updated previously reported.	report if there is r	new information regar	rding the clinical significance of my V	VES data that n	nay not have be
l hereby a	uthoriz	ze Baylo	r Genetics to conduct genetic tes	sting for myself (o	r my child) for the Wh	ole Exome Sequencing test as recon	nmended by my	physician.
							1	/
Printed Na	me			Signatu	re		Date (MI	M / DD / YYYY)
							/	/
Relationsh	ip to Pa	tient		Probanc	d Name		Proband D	OB (MM/DD/YY)
							/_	/
Physician's	s/Couns	elor's Sig	gnature				Date (MI	M / DD / YYYY)
FOR SAMI	PLES S	SUBMIT	TED FROM NEW YORK STATE					
INITIAL		at the er	nd of testing or not more than 60	days after the sai	mple was taken. How	rformed on my biological sample, an ever, by initialing here, I hereby auth cy for internal laboratory quality ass	orize the lab to	retain my
			n testing.		,			·

Consent authorization on next page



PHONE 1 800 411 4363 FAX 1.800.434.9850 CONNECT



WES ADVANTAGE REQUISITION

Date of Birth (MM / DD / YYYY) Patient Last Name Patient First Name MI Biological Sex INFORMATION AND CONSENT FOR TESTING Trio WES: (test codes 1600, 1722, 1532, 1533) We understand that our samples will be subjected to Trio WES, and will be analyzed to help interpret the sequence data of our child. A separate parental report will be issued regarding incidental findings. Testing of parental status for this category of results will be initiated independent of the proband's data. It may be possible to infer information about family member's results based on the proband's or other family member's results. Turnaround time to receive this report is up to 8 weeks. Duo WES: (test codes 1603, 1723) I understand that my sample will be subjected to Duo WES, and will be analyzed to help interpret the sequence data of my child. A separate parental report will be issued regarding incidental findings. Testing of parental status for this category of results will be initiated independent of the proband's data. It may be possible to infer information about family member's results based on the proband's or other family member's results. Turnaround time to receive this report is up to 8 weeks. Proband WES (test codes 1500, 1530, 1531) We understand that our samples will be subjected to targeted testing only (such as Sanger sequencing) and will NOT have WES testing. The laboratory will decide which changes will need parental studies. Testing of the parental status for the category of incidental findings will ONLY be initiated if there is a variant identified in the proband. Please read the below statements carefully and check the appropriate box and initial. Due to the nature of the methodology of this testing we are unable to guarantee that all pathogenic variants in each option will be detected by the WES testing. Note, if neither box is checked, or the form is not signed, the lab will default to the NO/ do NOT report option. MATERNAL REPORTING OPTIONS AND AUTHORIZATION INITIAL 1. MEDICALLY ACTIONABLE Pathogenic variants in genes included in the ACMG policy statement regarding recommendations for reporting of incidental findings will be reported as medically actionable on the WES report. Please report pathogenic variants in genes determined to be medically actionable by the ACMG policy statement. Please do NOT report pathogenic variants in genes included in the ACMG policy statement. Date of Birth (MM / DD / YYYY) Mother's Printed Name Mother's Signature PATERNAL REPORTING OPTIONS AND AUTHORIZATION INITIAL 1. MEDICALLY ACTIONABLE Pathogenic variants in genes included in the ACMG policy statement regarding recommendations for reporting of incidental findings will be reported as medically actionable on the WES report. Please report pathogenic variants in genes determined to be medically actionable by the ACMG policy statement. Please do NOT report pathogenic variants in genes included in the ACMG policy statement. Date of Birth (MM / DD / YYYY) Father's Printed Name Father's Signature FOR SAMPLES SUBMITTED FROM NEW YORK STATE MOTHER'S FATHER'S I understand that no genetic test other than those I have authorized shall be performed on my biological sample, and the sample INITIAL INITIAL will be destroyed at the end of testing or not more than 60 days after the sample was taken. However, by initialing here, I hereby authorize the lab to retain my sample(s) for longer retention in accordance with the laboratory retention policy for internal

laboratory quality assurance studies and possible research testing.



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Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Biological Sex	
RAW DATA CONSENT					
By checking this box, I agree t request, to me, my physician,	o allow Baylor Genetics to provide the or the requesting laboratory.	raw data such as FASTQ	or VCF sequencing files from my	genetic test, only upon	
RESEARCH & RECONTACT CONSE	NT				
For more information on research	at Baylor Genetics, please visit baylo	rgenetics.com. Please re	ad the below statements carefully	y and check the appropriate	box.
Note: If left blank, consent is inter	preted as "NO."				
☐ Lagrage to use of my do_identif	ied specimen for research to improve	gonotic testing for all na	tionts and contribute to scientific	rosoarch	
				research.	
in addition to agreeing above,	I agree to be contacted by Baylor Gen	etics regarding research	opportunities.		
CONTACT INFORMATION					
Phone #	Alternative Phon	e #	Email		
Address			ity	State Zip	
Preferred Method of Contact:	Email Mail	Phone			
NO IDO NOT	wish to be contacted regarding partici	pation in research studie	S.		
INITIAL	3 31	•			
PATIENT AUTHORIZATION					
The state of the s					
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Printed Name	Signa	ature		Date (MM / DD / YYY	Ύ)
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Relationship to Patient	Patie	nt Name		Patient Date of Birth (MM/	/DD/YY