

PHONE 1.800.411.4363 FAX 1.800.434.9850

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### **NGS PANEL REQUISITION**

Patient Last Name	Patient First Name				Date of Birth (MM / DD / YYYY
Address	City			Zip	Phone
Addiess	City		Genetic Sex:		
Accession #	Hospital / Medical Record #		Female Gender identity (if diff	Male (ferent from above):	Unknown
ORDERING PHYSICIAN		ADDITIONAL REPORT	S		
Ordering Physician	Institution Code	Name		Name	
Institution Name		Email		Email	
Email (Required for International Client	ris)	Phone		Phone	
Phone	Fax	Fax Note: Reports will be sent	by FAX except for intern	Fax national recipients	
PAYMENT (FILL OUT ONE OF THE O	PTIONS BELOW)				
SELF PAYMENT					
Pay With Sample	Bill To Patient				
O INSTITUTIONAL BILLING					
nstitution Name	Institution Code Institu	ition Contact Name	Institution Pt	hone	Institution Contact Email
INSURANCE					
Do Not Perform Test Until Pa	tient is Aware of Out-Of-Pocket Costs (excludes	prenatal testing)			
	-	gnosis Code(s) gnature of Authorization		ICD10	Diagnosis Code(s) (Required)
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 #
Name of Insured	Insured Date of Birth (MM / DD / YYYY)	Name of Insure	d	Insure	ed Date of Birth (MM / DD / YYY
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
m responsible for any co-pay, co-insura Senetics as outlined in the Good Faith Est	lor Genetics to provide my insurance carrier any i ince, and unmet deductible that the insurance pol imate I received. I understand that I am responsibl licare may not cover certain screening tests.	icy dictates. If self-pay is s	selected, I agree to pa	ay for the cost of	testing ordered and billed by Ba
Patient's Printed Name	Patient's Sig	nature			// Date (MM / DD / YYYY)
STATEMENT OF MEDICAL NECESSIT	TY (REQUIRED)				
This requisition hereby incorporates the entities, https://www.baylorgenetics.com/lisease, illness, impairment, symptom, sy	e Terms and Conditions of the Laboratory Service 1/terms-conditions-of-the-laboratory-services-int 1/ndrome, or disorder. The results will determine m 1/ndrome equested herein. I confirm that I have provided generated	ernational/. This test is m y patient's medical manage	nedically necessary fement and treatment	or the risk asses decisions. The per	ssment, diagnosis, or detection rson listed as the Ordering Phys
Physician's Printed Name	Physician's S	Signatura			Data (MM / DD / 2000)
cuvsician's Printen Name	Physician's S	Siunature			Date (MM / DD / YYYY)



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# **NGS PANEL REQUISITION**

Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex
INSTRUCTIONS FOR ORDERING				
Listed below are testing options th	at allow for comprehensive assessment to i	ncrease the diagnostic vielo	for natients with a focused disease phenot	vne Any combination of nane

testing or Chromosomal Microarray Analysis (CMA) can be ordered. Please note that the turnaround time may differ for each test. DISEASE-SPECIFIC PANELS | 1300 NEUROMUSCULAR PANELS ..... NEUROLOGY PANELS ..... BG-1300-P142-1 Neuromuscular Disorders Panel BG-1300-P397-1 Epilepsy Panel BG-1300-P76-1 STAT Epilepsy Panel IMMUNOLOGY PANELS ..... BG-1300-P236-1 Neurodevelopmental Disorders Panel BG-1300-P419-1 Cerebral Palsy Spectrum Disorders Panel BG-1300-P463-1 Primary Immunodeficiency Panel SKELETAL PANELS ..... If panel result is negative, reflex to test code 1520, Whole Exome Sequencing Reflex. ☐ BG-1300-P354-1 Skeletal Disorders Panel CONNECTIVE TISSUE PANELS ..... BG-1300-P92-1 Connective Tissue Disorders Panel **ADDITIONAL TESTING OPTIONS** 8665 Chromosomal Microarray Analysis (CMA)-HR+SNP Screen (Comprehensive) 2055 Comprehensive mtDNA analysis by NGS 6573 FMR1 CCG Repeat Expansion Analysis 6350 DMD Deletion/Duplication Analysis 6006 Angelman Syndrome Methylation Analysis **SAMPLE TYPES** Please refer to www.baylorgenetics.com for full sample requirements. Blood in EDTA Cultured Skin Fibroblast Date of Collection (MM / DD / YYYY) Buccal Swab Saliva Extracted DNA from Skin Biopsy<sup>†</sup> NOTE: Extracted DNA/RNA will only be accepted if the isolation of nucleic acids for clinical testing occurs in a CLIA-certified laboratory or a laboratory meeting equivalent requirements as determined by the CAP and/or the CMS.

ITEM CHECKLIST FOR TESTING		
<ul><li>□ Proband Sample (Required)</li><li>□ Signed Consent Form</li></ul>	☐ Clinical Note/Summary ☐ Requisition	☐ Indication for Study ☐ Pedigree

<sup>\*</sup> This sample type incurs an additional fee and typically adds 14 days to the turnaround time, depending on sample quality. † Baylor Genetics will store this sample for up to 14 days after the report is issued, allowing for follow-up testing if needed.



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# **NGS PANEL REQUISITION**

			/ /
Patient Last Na	me Patient First Na	me MI	Date of Birth (MM / DD / YYYY) Genetic Sex
INDICATION F	OR TESTING (REQUIRED)		
Please provide th	e following clinical information regarding the patient t		if available. Phenotypes listed are in HPO terms with the corresponding HPO ratory requires additional information, please indicate the health care provider to
Physician Nam	е	Physician Phone	ICD-10 Diagnosis Code(s)
PRE/PERINA	TAL HISTORY	EYE DEFECTS & VISION	MOTOR/COGNITIVE DEVELOPMENT
0001622	Prematurity - GA at birth	0000505 Visual Impairment	0000750 Delayed Speech & Language Development
0001511	Intrauterine Growth Restrictions	0000618 Blindness	0001270 Delayed Motor Milestones
0001562	Oligohydramnios	0000589 Coloboma	0002376 Developmental Regression
0001561	Polyhydramnios	0000526 Aniridia	☐ Intellectual Disability
0000476	Cystic Hygroma	0000528 Anophthalmia	0001256 Mild
0000776	Congenital Diaphragmatic Hernia	0000568 Microphthalmia	0002342 Moderate
0001508	Failure to Thrive	0000508 Ptosis	
0001539	Omphalocele	0000486 Strabismus	
0002084	Encephalocele	0000519 Cataract Congenital Bilateral	0000729 Autistic Spectrum Disorder
0010880	Increased Nuchal Translucency	<u> </u>	📙
		L.	L
STRUCTURAL	BRAIN ABNORMALITIES	NEUROLOGICAL	····· CRANIOFACIAL ·····
0001360	Holoprosencephaly	0001284 Areflexia	0000256 Macrocephaly
0001339	Lissencephaly	0200134 Epileptic Encephalopathy	0000252 Microcephaly
0002084	Encephalocele	0001250 Seizures	0001363 Craniosynostosis
0000238	Hydrocephalus	0002373 Febrile Seizures	0000204 Cleft Upper Lip
0002119	Ventriculomegaly	0012469 Infantile Spasms	0000175 Cleft Palate
0001273	Abnormality of Corpus Callosum		0000316 Hypertelorism
0002539	Cortical Dysplasia	O002123 Generalized Myoclonic Seizures	0000601 Hypotelorism
0012444	Brain Atrophy	Generalized Tonic-clonic	0008050 Abnormality of the Palpebral Fissures
0002352	Leukoencephalopathy	Seizures	0000286 Epicanthal Folds
0002269	Abnormality of Neuronal Migration	0010818 Generalized Tonic Seizur	es 0000288 Abnormality of the Philtrum
0002126	Polymicrogyria	0010819 Atonic Seizures	0010938 Abnormality of the External Nose
 	Pachgyria	0002121 Absence Seizures	
0002500	Abnormality of Cerebral White Matter	0011169 Generalized Clonic Seizu	res $\square$
0007266	Cerebral Dysmyelination		
0006808	Cerebral Hypomyelination	0001332 Dystonia	
0002134	Abnormality of the Basal Ganglia		
0002363	Abnormality of the Brainstem	0002072	
0007360	Aplasia/Hypoplasia of the Cerebellum	0001257 Spasticity	
_	Aplasia/Hypoplasia of the Cerebellar	0009830 Neuropathy	
0006817	Vermis		

Indications continued on next page



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# **NGS PANEL REQUISITION**

				/ /	
Patient Last Na	me Patient First I	Name	MI Da	ate of Birth (MM / DD / YY	YY) Genetic Sex
INDICATION F	OR TESTING (REQUIRED) - CONTINUED				
HAIR & SKIN		CARDIAC		···· GENITOURINA	ARY ·····
0000957	Cafe-Au-Lait Spots	0001631	Atria Septal Defect	0000113	Polycystic Kidney Dysplasia
0001034	Hypermelanotic Macule	 	Ventricular Septal Defect	0000107	Renal Cyst
0001010	Hypopigmentation of the Skin			0008738	Partially Duplicated Kidney
0008066	Abnormal Blistering of the Skin	0001655 	Patent Foramen Ovale	0000104	Renal Agenesis
0008064	Ichthyosis	0001713	Abnormality of Cardiac Ventricle	_	-
0000988	Skin Rash	0001636	Tetralogy of Fallot	0000085	Horseshoe Kidney
0001581	Recurrent Skin Infections	0001680	Coarctation of Aorta	U 0000069	Abnormality of the Ureter
0005306	Capillary Hemangiomas	0001647	Bicuspid Aortic Valve	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
0002299	Brittle Hair	0011675	Arrhythmia	0000062	Ambiguous Genitalia
	Drittle Hall				
				_	
<u> </u>					
RESPIRATOR	γ	METABOLIC		···· MUSCULOSKI	ELETAL
0002093	Respiratory Insufficiency	0001946	Ketosis	0011398	Hypotonia
0002878	Respiratory Failure	0003074	Hyperglycemia	0001276	Hypertonia
0002104	Apnea	0001943	Hypoglycemia	0000098	Tall Stature
_	·	0001743		0004322	Short Stature
0002791	Hypoventilation		Acidosis	0001382	Joint Hypermobility
0002883	Hyperventilation Recurrent Upper Respiratory Tract	U 0003128	Lactic Acidosis	0001371	Flexion Contracture
0002788	Infections	0003215	Dicarboxylic Aciduria	0002804	Arthrogryposis Multiplex Congenita
		0002490	Increased CSF lactate	0001161	Hand Polydactyly
		0001992	Organic Aciduria	0001829	Foot Polydactyly
		0030085	Abnormal CSF Lactate Level	0006101	Finger Syndactyly
		00003542	Increased Serum Pyruvate	0001770	Toe Syndactyly
GASTROINTE	STINAL	0003535	3-Methylglutaconic aciduria	0100490	Camptodactyly of Finger
0002021	Pyloric Stenosis	0001942	Metabolic acidosis	U 0012165	Oligodactyly
0002575	Tracheoesophogeal Fistula			0001762	Talipes Equinovarus
0002032	Esophageal Atresia	U 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 Phosphokinase	0001528	Hemihypertrophy
0002037	Inflammatory Bowel Disease	Abnormal	Newborn Screen	0001513	Obesity
0004389	Intestinal Pseudo-Obstruction	Unusual Co	olor/Odor	0001548	Overgrowth
0001377	Hepatic Failure Episodic Vomiting				Skeletal Dysplasia
0002372	Splenomegaly	⊢——		_	
0001744	Hepatomegaly	⊔		⊔	
0001508	Postnatal Failure to Thrive				
0002578	Gastroparesis				



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### **NGS PANEL REQUISITION**

Patient Last Na	me Patient First N	ame	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex
INDICATION F	OR TESTING (REQUIRED) - CONTINUED				
ENDOCRINE		HEMATOLOGY	•••••	OTHER	
0000819	Diabetes Mellitus	0001875 N	leutropenia	Organomegal	у
0000873	Diabetes Insipidus	0005	549 Congenital	Chronic Infect	tions
0000821	Hypothyroidism	Chron	nic	0004311 A	bnormality of Macrophages
0000829	Hypoparathyroidism	Cyclic		0001954 E	pisodic Fever
0000834	Abnormality of the Adrenal Glands	0001873 T	hrombocytopenia	0004313 H	ypogammaglobulinemia
0001738	Exocrine Pancreatic Insufficiency	0040185 M	Macrothrombocytopenia (	0010701 A	bnormal Immunoglobulins
0002721	Immunodeficiency	=	ecreased Mean Platelet Volu	=	nmunodeficiency
Ц		=	rythrocyte Macrocytosis	_	bnormal urinary odor
		=	pherocytosis		ood intolerance
			Pure Red Cell Aplasia		bnormally lax or hyperextensible skir
EAD DEEECT	S & HEARING ······	☐ Aplas		☐ Abnormal Mo	
	S & HEARING		plastic	=	y of Similar Disorder
0000407	Sensorineural Hearing Impairment	=	nemia	=	ethargy eukodystrophy
000	8619 Bilateral	0005526 B	Bone Marrow Hypocellularity		eukouystropny
0000405	Conductive Hearing Impairment				
0000410	Mixed Hearing Impairment	<u> </u>			
0004467	Preauricular Pit				
0000384	Preauricular Skin Tag	CANCER		GENES OF INTE	REST
0000369	Low-set Ears	Type of Cance	er		
000037		Age of Diagno	osis		
	Abnormality of the Pinna	Family Histor	ry of Cancer and Affected Re	latives	
<u> </u>					
Ш	· · · · · · · · · · · · · · · · · · ·				
ADDITIONAL	CLINICAL INFORMATION		DIFFERENTIAL	DIAGNOSIS	

Consent on next page



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### **NGS PANEL REQUISITION**

			//	
Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex

### INFORMED CONSENT FOR GENETIC TESTING ON THE NGS PANEL REQUISITION

# TEST INFORMATION .....

This consent form will provide you with information regarding genetic testing, which you should discuss with your healthcare provider or a genetic counselor. To assist you in understanding the reason for this testing, we have provided information about the testing process and potential results below.

The purpose of genetic testing is to determine if a genetic disease may be present or if there is an increased risk for a genetic disease to occur in a patient or their family. DNA is the genetic material that we receive from our parents. Genes are made of DNA and are the instructions for maintaining the health of our body. Each person has a unique set of DNA and most of the differences in our DNA do not impact our health. Genetic testing analyzes DNA to find any abnormal changes (mutations also called variants) that might cause disease, make it more likely to develop disease, and/or increase the chance of having a child affected by disease.

Most of the tests on this requisition, if ordered by your healthcare provider, will start being performed once your sample arrives at the laboratory. If your healthcare provider orders the whole exome sequencing (WES) test, this will only be performed if the results of other testing ordered do not explain the symptoms being experienced (WES will be run as a "reflex"). This testing provides a comprehensive analysis of the exome, which is the part of the human genome that helps the body make important proteins. The WES test will analyze the important regions of thousands of genes at the same time. Based on the symptoms that are known, genes with changes associated with these symptoms will be reported. It is possible that even if WES identifies the underlying genetic cause for a disease in a family this information may not help in predicting medical outcomes or changing medical management or treatment of disease. WES testing may also identify information about genes and diseases that have clear and immediate medical significance to your health or the health of your family members, even if that information is not related to currently known symptoms. You may consider discussing the significance of your results with your healthcare provider or genetic counselor.

The testing ordered by your healthcare provider can determine if you or your child have a variant associated with a genetic disease.

Depending on why genetic testing is needed, you might be tested for:

- · A known variant that has already been found in your family
- · A single gene or variant that causes a specific, suspected disease.
- · Multiple genes at the same time. These genes might cause similar diseases or might cause diseases that are unrelated to each other.
- · Multiple types of testing that each test for different variants.

# RESULTS

There are several types of test results that may be reported including:

- Positive: Positive or "abnormal" results mean there is a change in the DNA found that is related to your/your child's medical issues or that you/your child are at an increased risk of developing a disease in the future. It is possible to test positive for more than one variant. Positive results might include pathogenic variants (variants known to be associated with disease) and likely pathogenic variants (variants that are likely to be associated with disease).
- Negative: Negative or "normal" results mean no relevant variants related to your/your child's medical issues were detected or that you/your child are not expected to be at an increased risk for developing a disease in the future. This might indicate that there are no variants associated with disease in the gene(s) tested. Genetic testing, while highly accurate, might not detect a variant present in the gene(s) tested. This can be due to limitations of the information available about the gene(s) being tested, or limitations of the testing technology.
- Variant of Uncertain Significance: Testing can detect variant(s) in DNA which we do not yet fully understand. These are also referred to as variants of uncertain significance (VUS). Additional testing may be recommended for you or your family if a VUS is identified in a gene that may be associated with your/your child's medical condition.
- Secondary / Incidental Findings: Testing can sometimes detect a variant in a person's DNA unrelated to the reason for testing. If this variant is expected to have medical or reproductive significance, it is called a secondary or incidental finding.

# CONSIDERATIONS AND LIMITATIONS ......

- This consent form should only be used with the NGS Panel requisition. Consent forms for other requisitions are located at https://www.baylorgenetics.com/consent/.
- Results may indicate you have a genetic disease, are at increased risk to develop a genetic disease, and/or be at an increased risk to have a child with a genetic disease. It is important to understand that genetic tests, even if negative, cannot rule out every variant. It is not possible to exclude risks for all genetic diseases for you and your family members.
- Depending on the type of genetic testing performed and the results, additional genetic testing or other testing may be needed to fully understand the
  likelihood of your developing the disease or the severity of the disease. This additional testing might be needed for you/your child or other members of
  your family.
- It is recommended that you discuss genetic testing with your healthcare provider or genetic counselor before signing this consent and again after results are made available.
- It may not always be possible to complete testing. as sometimes the sample does not have enough DNA to perform testing or other reasons. In these cases, another sample may need to be sent to the laboratory to perform testing.



Patient Last Name

BAYLOR GENETICS 2450 HOLCOMBE BLVD. GRAND BLVD. RECEIVING DOCK HOUSTON, TX 77021-2024 **PHONE**1.800.411.4363 **FAX**1.800.434.9850

Date of Birth (MM / DD / YYYY)

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Genetic Sex



# NGS PANEL REQUISITION

### INFORMED CONSENT FOR GENETIC TESTING ON THE NGS PANEL REQUISITION

Patient First Name

### CONSIDERATIONS AND LIMITATIONS CONTINUED ......

• If several family members are tested, the correct interpretation of the results depends on the information provided about the relationships amongst family members. In rare cases, genetic testing can reveal that the true biological relationships in a family are not as they were reported. If a difference is identified, it may be necessary to share this information with the healthcare provider who ordered the testing.

МІ

- Genetic testing is highly accurate, however in rare cases, inaccurate results may occur. Reasons for this include, but are not limited to, mislabeled samples, inaccurate reporting of clinical/medical information, or rare technical errors.
- If you sign this consent form, but you no longer wish to have your sample(s) tested, you can contact the healthcare provider who ordered the test to
  cancel the test. If you wish to cancel testing, the laboratory must be notified of the cancellation request before 5 PM CST the business day after the
  sample has begun testing. If the laboratory is not notified of your cancellation request until after this time, you will be charged for the full cost of the
  test(s).
- Only Baylor Genetics and Baylor Genetics contracted partners will have access to the sample(s) provided to conduct the requested testing. Results
  will only be released to the following person(s): (i) a licensed healthcare provider, (ii) those authorized in writing, (iii) the patient or their personal
  representative, and (iv) those allowed access to test results by law. I understand that I have the right to access any test results directly from Baylor
  Genetics by providing a written request. I also understand that laboratory raw data, while not routinely released as part of the testing process, can be
  requested by providing a written request or HIPAA Authorization Form.
- In rare cases, persons with genetic diagnoses have experienced problems with insurance coverage and employment. The U.S. Federal Government has enacted several laws that prohibit discrimination based on genetic test results by health insurance companies and employers. In addition, these laws prohibit unauthorized disclosure of this information. For more information, you can visit www.genome.gov/10002077.
- Samples will be retained in the laboratory in accordance with the laboratory retention policy.
- After testing is complete, the de-identified submitted specimen may be used for test development and improvement, internal validation, quality
  assurance, and training purposes. DNA specimens are not returned to individuals or to referring heath care providers unless specific prior
  arrangements have been made.
- Samples from residents of New York State will not be included in research studies without your written consent and will not be retained for more than 60 days after receipt of the sample. No tests other than those authorized shall be performed on the biological sample.
- By signing this consent form, I understand and agree that variants identified may also be submitted to public databases, such as ClinVar. Such
  submission serves to contribute knowledge to the medical community. I understand that limited clinical information is also required for the submission
  of information to ClinVar's database and further that the contents of this limited clinical information may, although unlikely, include information that
  may identify me personally.
- It is possible that even if the test identifies the underlying genetic cause for the disease in your family, this information may not help in predicting the progression of disease or change management or treatment of disease.

## FOR WHOLE EXOME SEQUENCING (WES) REFLEX ORDERS ONLY

# INCIDENTAL FINDINGS .....

This test may also find changes in genes that cause symptoms or diseases not related to the reason for having the test. These are called Secondary Findings and are associated with clear and immediate medical significance to your health or the health of your family members.

### Category I: ACMG Secondary Findings

The American College of Medical Genetics (ACMG) has published guidelines for the reporting of these types of medically actionable or secondary findings (PMID: 34012068). These guidelines include a list of genes, which are updated occasionally, that are considered medically actionable and indicate laboratories should report pathogenic (disease-causing) findings in these genes. In accordance with an update to this policy statement (PMID: 25356965), you may choose to opt-in to receive this information.

### Category II: Other Incidental Findings

Medically actionable variants are changes found in genes known to be associated with disease but not associated with your current symptoms or clinical presentation. These variants are reported as they may cause severe, early-onset disease or may have implications for treatment and prognosis. You may choose to opt-in to receive this information.

# OTHER CONSIDERATIONS ·····

The report will not include findings in genes causing adult-onset progressive neurological diseases for which there is presently no prevention or cure. If the reason for testing includes symptoms that clearly indicate such a disease, we recommend pursuing targeted testing based on specific symptoms and not WES testing. However, if the reason for testing includes a clinical presentation for such a disease, then genes relating to this presentation may be reported.

Samples from biological parents may help facilitate interpretation of WES results. After the proband report is issued, parental samples received can be tested for just the changes in genes that are highly likely to be causative of disease in the affected individual. This follow-up testing for family members is available at an additional charge. Free testing for variants of unknown significance in the immediate family members is available with prior approval.



healthcare provider and billed to me by Baylor Genetics.

Patient Name

**BAYLOR GENETICS** 2450 HOLCOMBE BLVD GRAND BLVD. RECEIVING DOCK HOUSTON, TX 77021-2024

**PHONE** 1 800 411 4363 FAX 1.800.434.9850 CONNECT



Date Signed (MM / DD / YYYY)





# **NGS PANEL REQUISITION** Date of Birth (MM / DD / YYYY) МІ Patient Last Name Patient First Name Genetic Sex FOR WHOLE EXOME SEQUENCING (WES) REFLEX ORDERS ONLY REPORTING CHOICES ..... Please read the statements below 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 (disease-causing) variants in each option will be detected by WES. For all options below: If neither box is checked, or if the form is not signed, the consent for these options will be interpreted as "NO." SECONDARY FINDINGS ..... Pathogenic or likely pathogenic variants in genes included in the ACMG policy statement regarding recommendations for reporting of secondary findings will be reported as medically actionable on the WES report. YES Please report pathogenic or likely pathogenic variants in genes determined to be secondary findings by the ACMG policy statement. ☐ NO Please do NOT report pathogenic or likely pathogenic variants in genes included in the ACMG policy statement INCIDENTAL FINDINGS ..... Pathogenic or likely pathogenic variants in genes known to be associated with disease but not associated with your current symptoms or clinical presentation will be reported as medically actionable on the WES report. YES Please report pathogenic or likely pathogenic variants in genes known to be associated with disease but not associated with your current symptoms or clinical presentation NO Please do NOT report pathogenic or likely pathogenic variants in genes known to be associated with disease but not associated with your current symptoms or clinical presentation If a possible diagnosis can be made with new information, we would like to issue an updated report to the physician who ordered your WES. This review does NOT include a complete review of all of your data. YES If new information regarding the clinical significance of changes in my WES becomes known, I would like Baylor Genetics to issue an updated report which includes this information to my physician who ordered this WES testing. NO Please do NOT issue an updated report if there is new information regarding the clinical significance of my WES that becomes known. FINANCIAL AGREEMENT AND GUARANTEE By signing this consent form, I accept full and complete financial responsibility for all genetic testing ordered by my healthcare provider. For insurance billing, I hereby authorize Baylor Genetics to bill my health insurance plan on my behalf, and further authorize Baylor Genetics to release any information to my insurance carrier which is reasonably required for billing. I additionally designate Baylor Genetics as my designated representative for purposes of appealing any denial of benefits by my insurance carrier. I irrevocably assign associated payment to Baylor Genetics, and direct that payment be made directly to Baylor Genetics. I understand that my out-of-pocket costs may be different than the estimated amount indicated to me by Baylor Genetics as part of a verification of benefits investigation. I agree to be financially responsible for all amounts as indicated on the explanation of benefits issued by my health insurance plan. If my insurance provider sends a payment directly to me for unpaid services performed by Baylor Genetics on my behalf, I agree to endorse the insurance check as appropriate and forward such check to Baylor Genetics within thirty (30) days of receipt thereof, as payment towards Baylor Genetics' claim for services rendered. If I do not have health insurance, I agree to pay for the full cost of the genetic testing that was ordered by my

I understand that a completed Advance Beneficiary Notice (ABN) is required for Medicare patients if the service is deemed not medically necessary.

Patient Signature



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# **NGS PANEL REQUISITION**

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Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YY	YY) Genetic Sex
RESEARCH & RECONTACT CO	NSENT			
contact patients or their pr research involving the san	rovider(s) directly as part of th nple(s) and/or information ass	th, disease prevention, drug deve is research. I agree to allow Bayl ociated with this testing. I unders arch at Baylor Genetics, please vi	or Genetics to contact me or netainstand that patients generally re	ny provider(s) about possible
If I wish to opt out of being	recontacted for research purp	poses by Baylor Genetics, I under	stand that I may check the box	c below:
Please do not contact	me regarding any research th	nat uses information obtained fro	m this testing.	
For any research I may be will be made via secure en	contacted about, I prefer conta nail if possible):	act through the following method	s (please check all that apply	– if no choices are selected, contact
☐ Email ☐ Ma	il Phone			
PATIENT AUTHORIZATION				i i
received appropriate expla healthcare provider about	anations from my healthcare p the availability and importanc ticist who can provide such co	I have read, understand, and her provider about the planned genetice of genetic counseling and have tunseling services. All my quest	c test(s) and possible results. been provided with written in	I have been informed by my
I hereby give permission to	o Baylor Genetics to conduct g	enetic testing as recommended b	y my physician.	
				1 1
Patient Name		Patient Signature		Date Signed (MM / DD / YYYY)
				1
Patient's Parent / Personal Repr	resentative* Name	Patient's Parent / Personal Rep	resentative* Signature	Date Signed (MM/DD/YY)
				///
Relationship of Personal Repres	entative* to the Patient	Ordering Provider's Signature		Date Signed (MM/DD/YY)

<sup>\*</sup> If you are signing on behalf of the patient as the parent(s) and/or person with legal authority to act on behalf of the patient, you may be required to provide evidence of your authority.