

PHONE 1.800.411.4363 FAX 1.800.434.9850

CONNECT





MOLECULAR DIAGNOSTIC TESTING REQUISITION

				/ /
Patient Last Name	Patient First Name		MI	Date of Birth (MM / DD / YYYY)
Address	City	State Patient discharged from the hospital/facility:	Zip Genetic Sex:	Phone
Accession #	Hospital / Medical Record #	Yes No	Female Gender identity (if different) Male () Unknown () () () () () () () () () (
REPORTING RECIPIENTS				
Ordering Physician	Ins	stitution Name		
Email (Required for International Clie	nts) Ph	one	Fax	
ADDITIONAL RECIPIENTS				
Name	En	nail	Fax	
Name	En	nail	Fax	
PAYMENT (FILL OUT ONE OF THE	OPTIONS BELOW)			
SELF PAYMENT				•••••
Pay With Sample	Bill To Patient			
	Ditt 10 1 ditent			
) INSTITUTIONAL BILLING				
nstitution Name	Institution Code Institution	on Contact Name Ir	- PL P - DI	Land Carlot Facility
- nstitution name		on contact Name	stitution Phone	Institution Contact Email
INCHDANCE				
INSURANCE	Intigent in Assert of Out Of Decket Costs (evaluates a			
Do Not Perform Test Until F	latient is Aware of Out-Of-Pocket Costs (excludes p	renatal testing)		
Do Not Perform Test Until F		renatal testing)		gnature of Authorization
Do Not Perform Test Until F	ratient is Aware of Out-Of-Pocket Costs (excludes p of the Front/Back of Insurance Card(s) 2. ICD10 Diagno	renatal testing) psis Code(s) 3. Name of Orderin		gnature of Authorization
Do Not Perform Test Until F	latient is Aware of Out-Of-Pocket Costs (excludes p	renatal testing)	g Physician 4. Insured Si	gnature of Authorization / red Date of Birth (MM / DD / YYYY)
Do Not Perform Test Until F REQUIRED ITEMS 1. Copy of the company	ratient is Aware of Out-Of-Pocket Costs (excludes p of the Front/Back of Insurance Card(s) 2. ICD10 Diagno	renatal testing) psis Code(s) 3. Name of Orderin	g Physician 4. Insured Si Insu	/ /
Do Not Perform Test Until F REQUIRED ITEMS 1. Copy of the Copy of	ratient is Aware of Out-Of-Pocket Costs (excludes p of the Front/Back of Insurance Card(s) 2. ICD10 Diagno /	renatal testing) osis Code(s) 3. Name of Orderin Name of Insured	g Physician 4. Insured Si Insu	red Date of Birth (MM / DD / YYYY)
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Do Not Perform Test Until F REQUIRED ITEMS 1. Copy of Section 1. Copy	Patient is Aware of Out-Of-Pocket Costs (excludes prof the Front/Back of Insurance Card(s) 2. ICD10 Diagnor	renatal testing) pois Code(s) 3. Name of Ordering Name of Insured Patient's Relationship to Address of Insured City Secondary Insurance Co Secondary Member Policer any information necessary, nat the insurance policy dictates restand that I am responsible for	g Physician 4. Insured Si Insured Pho Stat Name Secuent Sec	red Date of Birth (MM / DD / YYYY) ne of Insured E Zip Dondary Insurance Co. Phone Dondary Member Group # r processing my insurance claim not paid by my insurance carrier f any and all payments that I recei
Do Not Perform Test Until F REQUIRED ITEMS 1. Copy of the second of the	ratient is Aware of Out-Of-Pocket Costs (excludes p of the Front/Back of Insurance Card(s) 2. ICD10 Diagno / / / Insured Date of Birth (MM / DD / YYYY) Phone of Insured State Zip Primary Insurance Co. Phone Primary Member Group # e Baylor Genetics to provide my insurance carrie any co-pay, co-insurance, and unmet deductible the non-covered and non-authorized services. I unde	renatal testing) psis Code(s) 3. Name of Ordering Name of Insured Patient's Relationship to Address of Insured City Secondary Insurance Co Secondary Member Policer any information necessary, and the insurance policy dictates restand that I am responsible for edoes not cover routine screen	g Physician 4. Insured Si Insured Pho Stat Name Secuent Sec	red Date of Birth (MM / DD / YYYY) ne of Insured e Zip ondary Insurance Co. Phone ondary Member Group # r processing my insurance claim not paid by my insurance carrier f
Do Not Perform Test Until F REQUIRED ITEMS 1. Copy of Section 1. Copy	Primary Insurance Co. Phone Primary Member Group # Baylor Genetics to provide my insurance carriany co-pay, co-insurance, and unmet deductible thon-covered and non-authorized services. I unde in payment for this test. Please note that Medicar	renatal testing) psis Code(s) 3. Name of Ordering Name of Insured Patient's Relationship to Address of Insured City Secondary Insurance Co Secondary Member Policer any information necessary, and the insurance policy dictates restand that I am responsible for edoes not cover routine screen	g Physician 4. Insured Si Insured Pho Stat Name Secuent Sec	red Date of Birth (MM / DD / YYYY) ne of Insured E Zip Dondary Insurance Co. Phone Dondary Member Group # r processing my insurance claim not paid by my insurance carrier f any and all payments that I recei
Do Not Perform Test Until F REQUIRED ITEMS 1. Copy of Statement's Relationship to Insured Address of Insured City Primary Insurance Co. Name Primary Member Policy # By signing below, I hereby authorize and that I am responsible for easons including, but not limited to, lirectly from my insurance company Patient's Printed Name STATEMENT OF MEDICAL NECESS This test is medically necessary for the patient's medical management and the statement of the stateme	ratient is Aware of Out-Of-Pocket Costs (excludes prof the Front/Back of Insurance Card(s) 2. ICD10 Diagnor	renatal testing) pois Code(s) 3. Name of Ordering Name of Insured Patient's Relationship to Address of Insured City Secondary Insurance Co Secondary Member Police er any information necessary, nat the insurance policy dictates restand that I am responsible force does not cover routine screen sture ature isease, illness, impairment, syrring Physician is authorized by	g Physician 4. Insured Si Insured Pho Stat Name Secuent Secuen	red Date of Birth (MM / DD / YYYY) ne of Insured Zip ondary Insurance Co. Phone ondary Member Group # r processing my insurance claim not paid by my insurance carrier f any and all payments that I recei
Do Not Perform Test Until F REQUIRED ITEMS 1. Copy of Section 1. Copy	Primary Insurance Co. Phone Primary Member Group # Baylor Genetics to provide my insurance carriany co-pay, co-insurance, and unmet deductible thon-covered and non-authorized services. I unde in payment for this test. Please note that Medicar Patient's Signa ITY (REQUIRED) he risk assessment, diagnosis, or detection of a diagnosis.	renatal testing) pois Code(s) 3. Name of Ordering Name of Insured Patient's Relationship to Address of Insured City Secondary Insurance Co Secondary Member Police er any information necessary, nat the insurance policy dictates restand that I am responsible force does not cover routine screen sture ature isease, illness, impairment, syrring Physician is authorized by	g Physician 4. Insured Si Insured Pho Stat Name Secuent Secuen	red Date of Birth (MM / DD / YYYY) ne of Insured Zip ondary Insurance Co. Phone ondary Member Group # r processing my insurance claim not paid by my insurance carrier i any and all payments that I receive in the control of the



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

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09.07.23

MOLECULAR DIAGNOSTIC TESTING REQUISITION

			/ /	
Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY) Gene	etic Sex
ETHNICITY				
African American Ashkenazi Jewish East Asian (China, Japan, Korea) Finnish French Canadian	Hispanic American Mennonite Middle Eastern (Saudi Arabia, Qatar, Ira Native American Northern European Caucasian (Scandin		Pacific Islander (Philippines, Micronesia, South Asian (India, Pakistan) Southeast Asian (Vietnam, Cambodia, Southern European Caucasian (Spain, Other (Specify):	Thailand)
SAMPLE				
SAMPLE TYPE			DATE OF COI	
Blood in EDTA-tube (purple-top) Buccal Swab 1	Cord Blood DNA (Specify, Saliva Other (Specify)		(MM/DD/YYY	Υ)
NOTE: Extracted DNA/RNA will only be a meeting equivalent requirements as deter	ccepted if the isolation of nucleic acids for clinical testin mined by the CAP and/or the CMS.	g occurs in a CLIA-cert	ified laboratory or a laboratory	/
Blood should not be sent from patients who I Only accepted for FMR1 CGG Repeat Expans	have had a bone marrow transplant or recent blood transfus sion Analysis (test code 6573)	ion		
INDICATION FOR TESTING (REQUIRED)		MOLECULAR DIA	AGNOSTIC TESTS	
Symptomatic (Summarize below)		MASSIVELY PA	ARALLEL SEQUENCING (BCM-MitomeNGS SM) F	PANELS
		TEST CODE	TEST NAME	SAMPLE TYPE *
		20100	Albinism Panel (13 genes)	BE, BUC, DNA
Symptomatic with Positive Family	/ History	20400	Bardet-Biedl Syndrome Panel (18 genes)	BE, BUC, DNA
Asymptomatic Population Screening	Positive Family History	2105	Cholestasis Panel (7 genes)	BE, BUC, DNA
o repaiding servering	O rosilite running riskery	2100	CoQ10 Panel (PDSS1, PDSS2, COQ2, COQ9, and ADCK3)	BE, BUC, DNA
Disease ICD10 Diagnosis Code(s)	Gene Variant	2120	Cobalamin Metabolism Panel + Severe MTHFR Deficiency (20 genes)	BE, BUC, DNA
		2625	COL1A1/2-Related Disorders (COL1A1 & COL1A2)	BE, BUC, DNA
TESTING OPTIONS		5095	Congenital Disorders of Glycosylation Panel (36 genes)	BE, BUC, DNA
Targeted Sequencing for Known F	Familial Mutation	2095	Fatty Acid Oxidation Deficiency Panel (20 genes)	BE, BUC, DNA
	below and complete section to the right)	2125	Glycogen Storage Disease (GSD) Comprehensive Panel (23 genes)	BE, BUC, DNA
Test Code	Gene	2126	Glycogen Storage Disease (GSD) Muscle Panel (13 genes)	BE, BUC, DNA
Full Gene Sequencing	Deletion/ Duplication Analysis	2127	Glycogen Storage Disease (GSD) Liver Panel (13 genes)	BE, BUC, DNA
FOR TARGETED TESTING SELECTION C	DNLY	2200	High Bone Mass Panel (14 genes)	BE, BUC, DNA
Proband Last Name	Proband First Name	21700	Hyperinsulinism Panel (8 genes)	BE, BUC, DNA
	/	21000	Hypoglycemia Panel (85 genes)	BE, BUC, DNA
Relationship to Proband	Date (MM / DD / YYYY)			
PROBAND TESTING LOCATION (SE	ELECT ONE)			
Baylor Genetics	Another laboratory			
Lab # Family #	Attach a copy of the Proband test results A positive control sample of the Proband is		* Refer to Sample Specific	rations Table (nage 11)
	requested. Please provide, if available.			ontinued on next nage

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MOLECULAR DIAGNOSTIC TESTING REQUISITION

				/ /	
Patient Last Nam	e Patient First Nar	ne	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex
MOLECULAR DIA	GNOSTIC TESTS				
MASSIVELY PA	RALLEL SEQUENCING (BCM-MitomeNG	S sm)			······
TEST CODE	TEST NAME	SAMPLE TYPE *	TEST CODE	TEST NAME	SAMPLE TYPE *
2090	Low Bone Mass Panel (23 genes)	BE, BUC, DNA	24001	Noonan Spectrum Disorders Panel (26	genes) BE, BUC, DNA, SA
32870	Maple Syrup Urine Disease (MSUD) Panel (BCKHDA, BCKHDB, DBT, and DBD)	BE, BUC, DNA	22100	Peroxisomal Disorders Panel (22 gene	
21900	Maturity-Onset Diabetes of the Young (MODY) Panel (25 genes)	BE, BUC, DNA	5274	Proximal Urea Cycle Disordersz (PUCD prehensive (Seq. & Del/Dup) (CPS1, NAC	BE BUILDING
2300	Myopathy/Rhabdomyolysis Panel (25 gene	BE, BUC, DNA	2190	Retinitis Pigmentosa + RPGR orf15 by (66 genes)	NGS BE, BUC, DNA
20200	Nephronophthisis Panel (NPHP1, INVS/ NPHP2, NPHP3, and NPHP4)	BE, BUC, DNA	2110	Urea Cycle Disorders (UCD) and Hyperammonemia by NGS (8 genes)	BE, BUC, DNA
SINGLE GENE	ANALYSIS				
	nd on this form, please obtain the test code	rom our website (www BM)	GL com) and write in t	the helow space(s)	
4 1001 10 1101 100	:		2	:	
Test Code	Gene	Test Code	Gene	Test Code	Gene
Test Name		Test Name		: Test Name	
TEST CODE	TEST NAME		DISORDER	•	CAMDIE TYPE *
5044	TEST NAME HSD17B10 Comprehensive (Seq & Del/Dup	Analysis)		xybutyryl-CoA Dehydrogenase Deficiency	SAMPLE TYPE * BE, DNA
5064	HMGCL Comprehensive (Seq & Del/Dup Ana	<u> </u>		ylglutaryl CoA Lyase Deficiency	BE, DNA
2874	MCCC1 and MCCC2 Comprehensive (Seq &	•	3-Methylcrontonyl-	BE, DNA	
3639	MCCC1 Comprehensive (Seg & Del/Dup Ana			-CoA-Carboxylase Deficiency	BE, DNA
3644	MCCC2 Comprehensive (Seq & Del/Dup Ana	<u>, </u>		-CoA-Carboxylase Deficiency	BE, DNA
3914	AUH Comprehensive (Seq & Del/Dup Analys	sis)	3-Methylglutaconic	c Aciduria Type I	BE, DNA
6603	ABCA4 Comprehensive (Seq & Del/Dup Ana	lysis)	ABCA4-Related Dis	sorders	BE, DNA
3284	LPIN1 Comprehensive (Seq & Del/Dup Anal	ysis)	Acute Recurrent M	lyoglobinuria (LPIN1-Related Disorders)	BE, DNA
2034	ACADSB Comprehensive (Seq & Del/Dup Ar	nalysis)	Acyl-CoA Dehydrog	BE, DNA	
2825	APRT Sequence Analysis		Adenine Phosphor	ibosyltransferase Deficiency	BE, DNA
5010	ADA Sequence Analysis		Adenosine Deamin	BE, DNA	
3699	ADSL Comprehensive (Seq & Del/Dup Analy	/sis)	Adenylosuccinase	Deficiency	BE, DNA
5279	ABCD1 Comprehensive (Seq & Del/Dup Ana	lysis)	Adrenoleukodystro	ophy	BE, DNA
3759	JAG1 Comprehensive (Seq & Del/Dup Analy	rsis)	Alagille Syndrome	BE, DNA	
2254	ALPL Comprehensive (Seq & Del/Dup Analy	rsis)	ALPL-Related Diso	BE, DNA	
6490	AR Sequence Analysis		Androgen Insensiti	ivity Syndrome	BE, DNA
6006	Angelman Syndrome (UBE3A) Methylation	Analysis	Angelman Syndror	BE, DNA	
3429	ARG1 Comprehensive (Seq & Del/Dup Anal	ysis)	Arginase Deficienc	BE, DNA	
3459	GATM Comprehensive (Seq & Del/Dup Anal	ysis)	Arginine: Glycine A	Amidinotransferase Deficiency	BE, DNA
6360	ASL Sequence Analysis		Argininosuccinate	Lyase Deficiency	BE, DNA
6742	ARX Comprehensive (Seg & Del/Dup Analys	is)	ARX-Related Disor	ders	BE. DNA

* Refer to Sample Specifications Table (page 11)



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MOLECULAR DIAGNOSTIC TESTING REQUISITION

B .:			
Patient Last Na	ame Patient First Name	MI Date of Birth (MM / DD / YYYY)	Genetic Sex
MOLECULAR D	DIAGNOSTIC TESTS		
SINGLE GEN	E ANALYSIS		
TEST CODE	TEST NAME	DISORDER	SAMPLE TYPE *
2205	AGA Sequence Analysis	Aspartylglycosaminuria	BE, DNA
6195	AIRE Sequence Analysis	Autoimmune Polyendocrinopathy 1	BE, DNA
3299	B4GALT7 Comprehensive (Seq & Del/Dup Analysis)	B4GALT7-Related Disorders	BE, DNA
3614	TAZ Comprehensive (Seq & Del/Dup Analysis)	Barth Syndrome (TAZ-Related Disorders)	BE, DNA
3499	BTD Comprehensive (Seq & Del/Dup Analysis)	Biotinidase Deficiency	BE, DNA
6012	Ashkenazic Mutation Panel (BLM)	Bloom Syndrome	BE, DNA
2429	LEMD3 Comprehensive (Seq & Del/Dup Analysis)	Buschke-Ollendorff Syndrome	BE, DNA
2589	TGFB1 Comprehensive (Seq & Del/Dup Analysis)	Camurati-Engelmann Disease	BE, DNA
6910	BRAF Sequence Analysis	Cardiofaciocutaneous Syndrome/ Costello Syndrome	BE, DNA
3439	SLC25A20 (CACT) Comprehensive (Seq & Del/Dup Analysis)	Carnitine Acylcarnitine Translocase Deficiency	BE, DNA
3364	SLC22A5 (OCTN2) Comprehensive (Seq & Del/Dup Analysis)	Carnitine Deficiency, Systemic	BE, DNA
3369	CPT1A Comprehensive (Seq & Del/Dup Analysis)	Carnitine Palmitoyltransferase IA Deficiency	BE, DNA
3374	CPT1B Comprehensive (Seq & Del/Dup Analysis)	Carnitine Palmitoyltransferase IB (CPT1B-Related Disorders)	BE, DNA
3164	CPT2 Comprehensive (Seq & Del/Dup Analysis)	Carnitine Palmitoyltransferase II Deficiency	BE, DNA
6125	RMRP Sequence Analysis	Cartilage Hair Hypoplasia (RMRP-Related Disorders)	BE, DNA
6733	CDKL5 Comprehensive (Seq & Del/Dup Analysis)	CDKL5-Related Disorders	BE, DNA
6376	CFTR Comprehensive Analysis (Seq, Del/Dup & 5T)	CFTR-Related Disorders (Cystic Fibrosis)	BE, DNA
6174	CHD7 Comprehensive (Seq & Del/Dup Analysis)	CHD7-Related Disorders (CHARGE Syndrome)	BE, DNA
6680	CHRNA7 Sequence Analysis	CHRNA7-Related Disorders	BE, DNA
3159	SLC25A13 (CTLN2) Comprehensive (Seq & Del/Dup Analysis)	Citrin Deficiency	BE, DNA
6180	ASS1 Sequence Analysis	Citrullinemia Type 1	BE, DNA
6150	RUNX2 Sequence Analysis	Cleidocranial Dysplasia	BE, DNA
3854	CABC1 (ADCK3) Comprehensive (Seq & Del/Dup Analysis)	Coenzyme Q10 Deficiency	BE, DNA
3419	COQ2 Comprehensive (Seq & Del/Dup Analysis)	Coenzyme Q10 Deficiency	BE, DNA
3414	PDSS2 Comprehensive (Seq & Del/Dup Analysis)	Coenzyme Q10 Deficiency	BE, DNA
7521	COL2A1 Comprehensive (Seq & Del/Dup Analysis)	COL2A1-Related Disorders	BE, DNA
6585	COL5A1 Sequence Analysis	COL5A1-Related Disorders	BE, DNA
	COL5A2 Sequence Analysis	COL5A2-Related Disorders	BE, DNA
3180	SDHA Sequence Analysis	Complex II Deficiency	BE, DNA
3185	SDHB Sequence Analysis	Complex II Deficiency	BE, DNA
3190	SDHC Sequence Analysis	Complex II Deficiency	BE, DNA
3195	SDHD Sequence Analysis	Complex II Deficiency	BE, DNA
2069	CYP17A1 Comprehensive (Seq & Del/Dup Analysis)	Congenital Adrenal Hyperplasia	BE, DNA
3259	CDG1A (PMM2) Comprehensive (Seq & Del/Dup Analysis)	Congenital Disorders of Glycosylation	BE, DNA
3454	CDG1B (MPI) Comprehensive (Seq & Del/Dup Analysis)	Congenital Disorders of Glycosylation	BE, DNA

* Refer to Sample Specifications Table (page 11)



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MOLECULAR DIAGNOSTIC TESTING REQUISITION

Patient Last Na	me Patient First Name	/	Genetic Sex
		MI Date OF BIT (IT (MIM / DD / TTTT)	Genetic Sex
MOLECULAR D	IAGNOSTIC TESTS		
SINGLE GENE	E ANALYSIS		•••••••••••••••••••••••••••••••••••••••
TEST CODE	TEST NAME	DISORDER	SAMPLE TYPE *
5119	CDG1M (DOLK) Comprehensive (Seq & Del/Dup Analysis)	Congenital Disorders of Glycosylation	BE, DNA
6545	HRAS Sequence Analysis	Costello Syndrome	BE, DNA
3150	SLC6A8 (CT1) Sequence Analysis	Creatine Transporter (CRTR) Deficiency-Related Disorders	BE, DNA
6949	RPS19 Comprehensive (Seq & Del/Dup Analysis)	Diamond Blackfan Anemia-RPS19 Related Disorders	BE, DNA
5310	TBX1 Sequence Analysis	DiGeorge Syndrome	BE, DNA
3464	DLD Comprehensive (Seq & Del/Dup Analysis)	Dihydrolipoamide Dehydrogense Deficiency	BE, DNA
6350	DMD Deletion/Duplication Analysis	DMD-Related Disorders	BE, DNA
2634	SLC39A13 Comprehensive (Seq & Del/Dup Analysis)	Ehlers-Danlos Syndrome, Spondylodysplastic Type 3	BE, DNA
6930	Type 4, STXBP1 Sequence Analysis	Epileptic Encephalopathy, Early Infantile	BE, DNA
7110	Type 7, KCNQ2 Sequence Analysis	Epileptic Encephalopathy, Early Infantile	BE, DNA
3749	ETHE1 Comprehensive (Seq & Del/Dup Analysis)	Ethylmalonic Encephalopathy	BE, DNA
6011	GLA Comprehensive (Seq & Del/Dup Analysis)	Fabry Disease	BE, DNA
2579	FAM20C Comprehensive (Seq & Del/Dup Analysis)	FAM20C-Related Disorders	BE, DNA
6740	LDLR Comprehensive (Seq & Del/Dup Analysis)	Familial Hypercholesterolemia	BE, DNA
6520	RUNX1 Sequence Analysis	Familial Platelet Disorder w/ Associated Myeloid Malignancy	BE, DNA
6573	FMR1 CGG Repeat Expansion	FMR1-Related Disorders (Fragile X Syndrome)	BE, BUC, DNA, SA
6570	FMR1 Sequence Analysis	FMR1-Related Disorders (Fragile X Syndrome)	BE, DNA
6345	PORCN Sequence Analysis	Focal Dermal Hypoplasia	BE, DNA
6690	FOXF1 Sequence Analysis	FOXF1-Related Disorders	BE, DNA
6031	Friedreich Ataxia Repeat Expansion Analysis	Friedreich Ataxia Syndrome	BE, DNA
6365	FXN Sequence Analysis	Friedreich Ataxia Syndrome	BE, DNA
3939	FBP1 Comprehensive (Seq & Del/Dup Analysis)	Fructose 1,6 Bisphosphatase Deficiency	BE, DNA
3740	FH Sequence Analysis	Fumarate Hydratase Deficiency (FH-Related Disorders)	BE, DNA
3279	GALE Comprehensive (Seq & Del/Dup Analysis)	Galactosemia	BE, DNA
3249	GALT Comprehensive (Seq & Del/Dup Analysis)	Galactosemia	BE, DNA
3799	GALK1 Comprehensive (Seq & Del/Dup Analysis)	Galactokinase Deficiency	BE, DNA
6955	SLC2A1 (GLUT1) Sequence Analysis	Glucose Transporter Type 1 Deficiency Syndrome	BE, DNA
3689	Type 1, GCDH Comprehensive (Seq & Del/Dup Analysis)	Glutaric Acidemia	BE, DNA
2044	Type 3, C7orf10 Comprehensive (Seq & Del/Dup Analysis)	Glutaric Acidemia	BE, DNA
5034	AMT Comprehensive (Seq & Del/Dup Analysis)	Glycine Encephalopathy	BE, DNA
3534	Type 0 Liver Isoform, GYS2 Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA
3839	Type 0 Muscle Isoform, GYS1 Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA
3134	Type 1a (GSD1A), G6PC Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA

* Refer to Sample Specifications Table (page 11)



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Patient Last Name Patient First Name		MI Date of Birth (MM / DD / YYYY) Get	Senetic Sex	
MOLECULAR DI	AGNOSTIC TESTS			
SINGLE GENE	ANALYSIS			
TEST CODE	TEST NAME	DISORDER	SAMPLE TYPE	
3834	Type 1 (b,c,d), SLC37A4 (GSD1B) Comprehensive (Seq & Del/ Dup Analysis)	Glycogen Storage Disease	BE, DNA	
3404	Type II, GAA Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA	
3829	Type IV (GSDIV), GBE1 Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA	
3804	Type V (GSDV), PYGM Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA	
3794	Type VI (GSDVI), PYGL Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA	
3824	Type VII (GSDVII), PFKM Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA	
3984	Type IX (GSDIX), PHKG2 Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA	
3809	Type X (GSDX), PGAM2 Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA	
2529	Type XIII (GSDXIII), ENO3 Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA	
2524	Type XIV (GSDXIV), PGM1 Comprehensive (Seq & Del/Dup Analysis)	Glycogen Storage Disease	BE, DNA	
5129	GNE Comprehensive (Seq & Del/Dup Analysis)	GNE-Related Disorders	BE, DNA	
3149	GAMT Comprehensive (Seq & Del/Dup Analysis)	Guanidinoacetate Methyltransferase Deficiency	BE, DNA	
6019	Connexin 26 - GJB2 Sequence Analysis	Hearing Loss	BE, DNA	
6355	Connexin 30 - GJB6 (232kb and 309kb) Deletion/Duplication Analysis	Hearing Loss	BE, DNA	
3030	Mitochondrial Nonsyndromic Hearing Loss and Deafness Mutation Panel (MT-RNR1, MT-TS1, MT-TS2, MTRNR1)	Hearing Loss	BE, DNA	
6395	MY07A Sequence Analysis	Hearing Loss	BE, DNA	
6655	CDH23 Sequence Analysis	Hearing Loss	BE, DNA	
6670	POU3F4 Sequence Analysis	Hearing Loss	BE, DNA	
3344	TIMM8A Comprehensive (Seq & Del/Dup Analysis)	Hearing Loss	BE, DNA	
5405	Hemochromatosis Panel by Sanger Sequencing (HAMP, HFE, HFE2, SLC40A1, TFR2)	Hemochromatosis	BE, DNA	
3129	ALDOB Comprehensive (Seq & Del/Dup Analysis)	Hereditary Fructose Intolerance	BE, DNA	
3784	ALDOB, FBP1, GYS2, & PC Sequence Analysis	Hereditary Fructose Intolerance	BE, DNA	
2145	SEPT9 Targeted Mutation Analysis	Hereditary Neuralgic Amyotrophy (HNA)	BE, DNA	
6925	HEXA Sequence Analysis	Hexosaminidase A Deficiency/ Tay-Sachs Disease	BE, DNA	
5390	HNRNPA1 Sequence Analysis	HNRNPA1-Related Disorders	BE, DNA	
3544	HLCS Comprehensive (Seq & Del/Dup Analysis)	Holocarboxylase Synthetase Deficiency	BE, DNA	
3974	CBS Comprehensive (Seq & Del/Dup Analysis)	Homocystinuria Caused by Cystathionine Beta-Synthase Deficiency	BE, DNA	
2075	HPD Sequence Analysis	HPD-Related Disorders	BE, DNA	
6034	Huntington Disease Repeat Expansion Analysis	Huntington Disease (Disease Specific Consent Required)	BE, DNA	
5285	GLUD1 Sequence Analysis	Hyperinsulinism-Hyperammonemia Syndrome	BE, DNA	
2070	GNMT Sequence Analysis	Hypermethioninemia	BE, DNA	
2135	AHCY Sequence Analysis	Hypermethioninemia with S-Adenosylhomocysteine Hydrolase Deficiency	BE, DNA	

* Refer to Sample Specifications Table (page 11)



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Patient Last Na	ame Patient First Name	MI Date of Birth (MM / DD / YYYY) G	enetic Sex
MOLECULAR D	IAGNOSTIC TESTS		
SINGLE GEN	E ANALYSIS		
TEST CODE	TEST NAME	DISORDER	SAMPLE TYPE *
3239	SLC25A15 (HHH) Comprehensive (Seq & Del/Dup Analysis)	Hyperornithinemia-Hyperammonemia-Homocitrullinuria (HHH) Syndrome	BE, DNA
5139	ALDH4A1 Comprehensive (Seq & Del/Dup Analysis)	Hyperprolinemia Type II	BE, DNA
2654	SLC34A1 (NPT2) Comprehensive (Seq & Del/Dup Analysis)	Hypophosphatemic Nephrolithiasis/Osteoporosis, 1	BE, DNA
5045	IYD Sequence Analysis	Hypothyroidism, Congenital	BE, DNA
5395	HNRNPA2B1 Sequence Analysis	Inclusion Body Myopathy with Early-Onset Paget Disease with or without Frontotemporal Dementia 2	BE, DNA
6036	Incontentia Pigmenti Common Deletion Analysis	Incontinentia Pigmenti (IKBKG-Related Disorders)	BE, DNA
7100	IKBKG Sequence Analysis	Incontinentia Pigmenti (IKBKG-Related Disorders)	BE, DNA
3314	ABCB11 Comprehensive (Seq & Del/Dup Analysis)	Intrahepatic Cholestasis	BE, DNA
3319	ABCB4 Comprehensive (Seq & Del/Dup Analysis)	Intrahepatic Cholestasis	BE, DNA
3309	ATP8B1 Comprehensive (Seq & Del/Dup Analysis)	Intrahepatic Cholestasis	BE, DNA
2029	ACAD8 Comprehensive (Seq & Del/Dup Analysis)	Isobutyryl-CoA Dehydrogenase Deficiency	BE, DNA
3684	IVD Comprehensive (Seq & Del/Dup Analysis)	Isovaleric Acidemia	BE, DNA
6037	Kennedy Disease Repeat Expansion Analysis	Kennedy Disease	BE, DNA
5370	KIF11 Sequence Analysis	KIF11-Related Disorders	BE, DNA
6415	GALC Sequence Analysis	Krabbe Disease	BE, DNA
3389	ACADL Comprehensive (Seq & Del/Dup Analysis)	LCAD Deficiency	BE, DNA
3124	HADHA Comprehensive (Seq & Del/Dup Analysis)	LCHAD Deficiency (HADHA-Related Disorders)	BE, DNA
6065	PTPN11 Sequence Analysis	LEOPARD Syndrome	BE, DNA
6475	RAF1 Sequence Analysis	LEOPARD Syndrome	BE, DNA
6240	HPRT Sequence Analysis	Lesch-Nyhan Syndrome	BE, DNA
3719	DARS2 Comprehensive (Seq & Del/Dup Analysis)	Leukoencephalopathy	BE, DNA
3819	TRMU Comprehensive (Seq & Del/Dup Analysis)	Liver Failure, Acute Infantile	BE, DNA
6039	OCRL Sequence Analysis	Lowe Syndrome	BE, DNA
2039	ACSF3 Comprehensive (Seq & Del/Dup Analysis)	Malonic & Methylmalonic Aciduria, Combined	BE, DNA
2774	Type 1A, BCKDHA Comprehensive (Seq & Del/Dup Analysis)	Maple Syrup Urine Disease	BE, DNA
2884	Type 1B, BCKDHB Comprehensive (Seq & Del/Dup Analysis)	Maple Syrup Urine Disease	BE, DNA
3869	Type 2, DBT Comprehensive (Seq & Del/Dup Analysis)	Maple Syrup Urine Disease	BE, DNA
3119	ACADM Comprehensive (Seq & Del/Dup Analysis)	MCAD Deficiency	BE, DNA
6380	ARSA Sequence Analysis	Metachromatic Leukodystrophy (Arylsulfatase A Deficiency)	BE, DNA
2569	cblE Type, MTRR Comprehensive (Seq & Del/Dup Analysis)	Methylcobalamin Deficiency	BE, DNA
2054	cblG Type, MTR Comprehensive (Seq & Del/Dup Analysis)	Methylcobalamin Deficiency	BE, DNA
3602	Methylmalonic Acidemia Comprehensive Panel (MUT, MMAA, MMAB)	Methylmalonic Acidemia	BE, DNA
3399	MCEE Comprehensive (Seq & Del/Dup Analysis)	Methylmalonic Acidemia	BE, DNA
3579	MMAA Comprehensive (Seq & Del/Dup Analysis)	Methylmalonic Acidemia	BE, DNA

* Refer to Sample Specifications Table (page 11)



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MOLECULAR DIAGNOSTIC TESTING REQUISITION

Patient Last Name Patient First Name		MI Date of Birth (MM / DD / YYYY)	Genetic Sex
MOLECULAR DI	AGNOSTIC TESTS		
SINGLE GENE	ANALYSIS		
TEST CODE	TEST NAME	DISORDER	SAMPLE TYPE
3584	MMAB Comprehensive (Seq & Del/Dup Analysis)	Methylmalonic Acidemia	BE, DNA
3444	MMACHC Comprehensive (Seq & Del/Dup Analysis)	Methylmalonic Acidemia	BE, DNA
3889	MMADHC Comprehensive (Seq & Del/Dup Analysis)	Methylmalonic Acidemia	BE, DNA
3589	MUT Comprehensive (Seq & Del/Dup Analysis)	Methylmalonic Acidemia	BE, DNA
2564	cblF Type, LMBRD1 Comprehensive (Seq & Del/Dup Analysis)	Methylmalonic Aciduria and Homocystinuria	BE, DNA
3064	TYMP Comprehensive (Seq & Del/Dup Analysis)	MNGIE Syndrome	BE, DNA
3599	MOCS1 Comprehensive (Seq & Del/Dup Analysis)	Molybdenum Cofactor Deficiency	BE, DNA
3619	MOCS2 Comprehensive (Seq & Del/Dup Analysis)	Molybdenum Cofactor Deficiency	BE, DNA
6385	Type I (MPS I), IDUA Sequence Analysis	Mucopolysaccharidosis	BE
6814	Type II (MPS II), IDS Comprehensive (Seq & Del/Dup w/Inv Analysis)	Mucopolysaccharidosis	BE, DNA
3604	Multiple Acyl-CoA Dehydrogenase Deficiency (MADD) Comprehensive Panel (Seq & Del/Dup Analysis) (ETFA, ETFB, ETFDH)	Multiple Acyl-CoA Dehydrogenase Deficiency (MADD)	BE, DNA
3859	ETFA Comprehensive (Seq & Del/Dup Analysis)	Multiple Acyl-CoA Dehydrogenase Deficiency (MADD)	BE, DNA
3864	ETFB Comprehensive (Seq & Del/Dup Analysis)	Multiple Acyl-CoA Dehydrogenase Deficiency (MADD)	BE, DNA
3844	ETFDH Comprehensive (Seq & Del/Dup Analysis)	Multiple Acyl-CoA Dehydrogenase Deficiency (MADD)	BE, DNA
6041	Myotonic Dystrophy Type 1 Repeat Expansion Analysis	Myotonic Dystrophy Type 1	BE, DNA
3354	NAGS Comprehensive (Seq & Del/Dup Analysis)	N-Acetyglutamate Synthase (NAGS) Deficiency	BE, DNA
7523	LMX1B Comprehensive (Seq & Del/Dup Analysis)	Nail-Patella Syndrome	BE, DNA
6900	SHOC2 Sequence Analysis	Noonan-like Syndrome	BE, DNA
6845	LEP Sequence Analysis	Obesity, Monogenic Nonsyndromic	BE, DNA
6850	LEPR Sequence Analysis	Obesity, Monogenic Nonsyndromic	BE, DNA
6855	PCSK1 Sequence Analysis	Obesity, Monogenic Nonsyndromic	BE, DNA
6860	POMC Sequence Analysis	Obesity, Monogenic Nonsyndromic	BE, DNA
6083	X-Linked, GPR143 Comprehensive (Seq & Del/Dup Analysis)	Oculocutaneous Albinism	BE, DNA
3529	Type 3, OPA3 Comprehensive (Seq & Del/Dup Analysis)	Optic Atrophy Type 3	BE, DNA
3144	OTC Comprehensive (Seq & Del/Dup Analysis)	Ornithine Transcarbamylase (OTC) Deficiency	BE, DNA
2574	AMER1 Comprehensive (Seq & Del/Dup Analysis)	Osteopathia Striata with Cranial Sclerosis	BE, DNA
2624	TCIRG1 Comprehensive (Seq & Del/Dup Analysis)	Osteopetrosis	BE, DNA
2604	CA2 Comprehensive (Seq & Del/Dup Analysis)	Osteopetrosis with Renal Tubular Acidosis	BE, DNA
6885	PCDH19 Sequence Analysis	PCDH19-Related X Linked Female-Limited Epilepsy w/MR	BE, DNA
3169	PDHA1 Comprehensive (Seq & Del/Dup Analysis)	PDH Complex Deficiency	BE, DNA
3899	PDHB Comprehensive (Seq & Del/Dup Analysis)	PDH Complex Deficiency	BE, DNA
3924	PDHX Comprehensive (Seq & Del/Dup Analysis)	PDH Complex Deficiency	BE, DNA
3894	PDP1 Comprehensive (Seq & Del/Dup Analysis)	PDH Complex Deficiency	BE, DNA

* Refer to Sample Specifications Table (page 11)



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MOLECULAR DIAGNOSTIC TESTING REQUISITION

Patient Last Na	me Patient First Name	MI Date of Birth (MM / DD / YYYY)	Genetic Sex
		Bate of Birth (MAY 257 TTT)	Genetic Sex
	AGNOSTIC TESTS		
SINGLE GENE			
TEST CODE	TEST NAME	DISORDER	SAMPLE TYPE *
6550	GJC2 Sequence Analysis	Pelizaeus-Merzbacher-Like Disease	BE, DNA
5365	PGM3 Sequence Analysis	PGM3-Related Disorders	BE, DNA
3139	PAH Comprehensive (Seq & Del/Dup Analysis)	Phenylalanine Hydroxylase Deficiency (PKU)	BE, DNA
6149	PLP1 Comprehensive (Seq & Del/Dup Analysis)	PLP1-Related Disorders	BE, DNA
3729	RARS2 Comprehensive (Seq & Del/Dup Analysis)	Pontocerebellar Hypoplasia Type 6	BE, DNA
6050	Prader-Willi Syndrome Methylation Analysis	Prader-Willi Syndrome	BE, DNA
7105	MAGEL2 Sequence Analysis	Prader-Willi-like Syndrome; Intellectual Disability; Autism	BE, DNA
3622	Propionic Acidemia Comprehensive Panel (Seq & Del/Dup Analysis) (PCCA & PCCB)	Propionic Acidemia	BE, DNA
6048	Prothrombin Mutation Panel (F2)	Prothrombin	BE, DNA
6790	PTEN Comprehensive (Seq & Del/Dup Analysis)	PTEN-Related Disorders	BE, DNA
5025	PNP Sequence Analysis	Purine Nucleoside Phosphorylase Deficiency	BE, DNA
2444	CTSK Comprehensive (Seq & Del/Dup Analysis)	Pycnodysostosis	BE, DNA
6950	ALDH7A1 Sequence Analysis	Pyridoxine-Dependent Seizures	BE, DNA
3919	DLAT Comprehensive (Seq & Del/Dup Analysis)	Pyruvate Dehydrogenase E2 Deficiency	BE, DNA
3754	PC Comprehensive (Seq & Del/Dup Analysis)	Pyruvate Carboxylase Deficiency	BE, DNA
5300	RAG2 Sequence Analysis	RAG2-Related Disorders	BE, DNA
6736	MECP2 Comprehensive (Seq & Del/Dup Analysis)	Rett Syndrome (MECP2-Related Disorders)	BE, DNA
6635	FOXG1 Sequence Analysis	Rett Syndrome, Congenital Variant	BE, DNA
6565	VDR Sequence Analysis	Rickets-Alopecia Syndrome	BE, DNA
6758	CREBBP Comprehensive (Seq & Del/Dup Analysis)	Rubinstein-Taybi Syndrome	BE, DNA
3929	ACADS Comprehensive (Seq & Del/Dup Analysis)	SCAD Deficiency	BE, DNA
6285	COL10A1 Sequence Analysis	Schmid Metaphyseal Chondrodysplasia (SMCD)	BE, DNA
6745	DHCR7 Sequence Analysis	Smith-Lemli-Opitz Syndrome	BE, DNA
6760	RAI1 Sequence Analysis	Smith-Magenis Syndrome	BE, DNA
6059	SMN1/SMN2 Copy Number Analysis	Spinal Muscular Atrophy (SMA) Diagnostic Test	BE, DNA
2899	PRKCG Comprehensive (Seq & Del/Dup Analysis)	Spinocerebellar Ataxia 14 (SCA14)	BE, DNA
6060	SRY Molecular Analysis	SRY-Related Phenotypes	BE, DNA
5024	ALDH5A1 Comprehensive (Seq & Del/Dup Analysis)	Succinic Semialdehyde Dehydrogenase Deficiency	BE, DNA
2510	TMLHE Sequence Analysis	TMLHE Deficiency	BE, DNA
2513	TMLHE Exon 2 Deletion Analysis	TMLHE Deficiency	BE, DNA
3969	TCN2 Comprehensive (Seq & Del/Dup Analysis)	Transcobalamin II Deficiency	BE, DNA
3624	Trifunctional Protein Deficiency Comprehensive Panel (Seq & Del/Dup Analysis) (HADHA and HADHB)	Trifunctional Protein Deficiency	BE, DNA

* Refer to Sample Specifications Table (page 11)



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MOLECULAR DIAGNOSTIC TESTING REQUISITION

					/	/	
Patient Last Na	me	Patient First Name		MI	Date of Birth (MM / DD /	YYYY) Go	enetic Sex
MOLECULAR D	AGNOSTIC TESTS						
SINGLE GENE	ANALYSIS						
TEST CODE	TEST NAME			DISORDER			SAMPLE TYPE *
3634	HADHB Comprehensive (Seq & Del/Dup Analysis) (H	ADHB)	Trifunctional Pr	rotein Deficiency		BE, DNA
5005	TSHR Sequence Analysis	i		TSHR-Related	Disorders		BE, DNA
3449	Type I, FAH Comprehensi	ve (Seq & Del/Dup Analysis	s)	Tyrosinemia			BE, DNA
2084	Type II, TAT Comprehensi	ve (Seq & Del/Dup Analysis	5)	Tyrosinemia			BE, DNA
6650	USH2A Sequence Analys	is		Usher Syndron	ne 2A		BE, DNA
6660	CLRN1 Sequence Analysi	is		Usher Syndron	ne 3A		BE, DNA
3359	ACADVL Comprehensive	(Seq & Del/Dup Analysis)		VLCAD Deficier	ісу		BE, DNA
2554	ATP7B Comprehensive (Seq & Del/Dup Analysis)		Wilson Disease			BE, DNA
6430	LIPA Sequence Analysis			Wolman Diseas	se		BE, DNA
Test Code 1 code. These performed	should only be used wh	equesting sequencing of en Baylor Genetics has ccuss prior to sending a	already identifie sample. A positiv	d the sequence ve control may	r which the Baylor Genetic e change in the proband/o be required in some case paylorgenetics.com.	original patient. If pr	oband testing was
Name of First F	Patient Studied	Relati	onship to Patient Stu	ıdied E	Baylor Genetics Lab #	Family #	
This Family Me	mber is Currently:						
○ A	SYMPTOMATIC If S	SYMPTOMATIC, please provi	de details. Please at	tach additional p	pages, if needed.		
○ s	YMPTOMATIC				Inc	cludo a podiargo chowir	ng familial relationships.
c	opy of Original Results Attac	hed (REQUIRED)			me	ituue a peurgree snown.	ig rannuat retationships.
CUSTOM FAM	LY SEQUENCING TEST						
TEST CODE	TEST NAME		GENE NAME	(REQUIRED)	MUTATION/UNC	LASSIFIED VARIANT (F	REQUIRED)
1580	Custom Family Member Se	equence Analysis - Gene 1					
SAMPLE SPECI	FICATIONS TABLE						

ABBREVIATION	SAMPLE NAME	RECOMMEN	DED AMOUNT	SHIPPING INSTRUCTIONS	SPECIAL NOTES	
ADDICEVIATION	JAMI LE NAME	(2 YRS - ADULT) (NEWBORN - 2YRS)		Shirr ind instructions	SI EGIAL NOTES	
BE	Blood in EDTA tube (purple-top)	3 - 5 cc	2 - 3 cc			
BUC	Buccal Swab	See Special Notes	See Special Notes	Ship at room temperature in an insulated container by	Collected with ORAcollect•Dx (OCD-100) self-collection kit (provided by Baylor Genetics with instructions). It is highly recommended that the sample be collected by a healthcare professional. Buccal swab is an accepted sample type for FMR1 CGG Repeat Expansion Analysis (test code 6573) only.	
СВ	Cord Blood	N/A	1 - 2 cc	overnight courier. Do not heat or freeze.	Ensure properly labeled. Also send 3 cc of maternal blood in properly labeled EDTA tube for MCC studies at no charge as needed.	
DNA	DNA, Extracted	10 -15 ug	10 -15 ug		Minimal concentration of 50ng/uL; A260/A280 of ~1.7	
SA	Saliva	See Special Notes	See Special Notes		Collected with Oragene DNA Self-Collection Kit.	



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INFORMED CONSENT FOR MOLECULAR DIAGNOSTIC TESTING

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

The testing ordered by your healthcare provider can determine if you or your child have a variant associated with a genetic disease. "Your child" can also mean your unborn child, for the purposes of this consent.

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.

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 cannot be used for whole exome sequencing (WES), whole genome sequencing (WGS), or Huntington's disease testing. These tests have specific consents that 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 CONFIDENTIALITY AND SPECIMEN RETENTION

• 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.



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INFORMED CONSENT FOR MOLECULAR DIAGNOSTIC TESTING											
				/ /							
Pat	ient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex						
PAT	PATIENT CONFIDENTIALITY AND SPECIMEN RETENTION (CONT.) ····································										
	 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. 										
9	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.										
1	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.										
	•	aboratory in accordance with the labor	,	•	10.1.0						
i	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.										
9	submission serves to contribute	nderstand and agree that variants ider knowledge to the medical community. ase and further that the contents of thi	. I understand that	limited clinical information is also rec	uired for the submission						
		identifies the underlying genetic caus e management or treatment of disease		n your family, this information may not	help in predicting the						
FINANCIAL AGREEMENT AND GUARANTEE											
bill to r of a dir- par hea to e Bay	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 healthcare provider and billed to me by Baylor Genetics.										
l ur	nderstand that a completed Adv	ance Beneficiary Notice (ABN) is requir	ed for Medicare pa	atients if the service is deemed not me	edically necessary.						
RE	CONTACT FOR RESEARCH CONSE	ENT									
cor	ntact patients or their provider(search involving the sample(s) a	earch relating to health, disease preve s) directly as part of this research. I agr nd/or information associated with this e information on research at Baylor Ger	ree to allow Baylor testing. I understa	Genetics to contact me or my provide and that patients generally receive no	r(s) about possible						
lf I	wish to opt out of being reconta	cted for research purposes by Baylor (Genetics, I underst	and that I may check the box below:							
□F	Please do not contact me regard	ing any research that uses information	n obtained from thi	s testing.							
	any research I may be contacte l be made via secure email if po	ed about, I prefer contact through the fossible):	ollowing methods	(please check all that apply – if no cho	ices are selected, contact						
	Email □ Phone □ Mail										



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INFORMED CONSENT FOR MOLECULAR DIAGNOSTIC TESTING

			/ /							
Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / Y	YYYY) Gen	etic Sex					
PATIENT AUTHORIZATION										
By signing this statement of consent, I acknowledge that I have read, understand, and hereby grant my informed consent for genetic testing. I have received appropriate explanations from my healthcare provider about the planned genetic test(s) and possible results. I have been informed by my healthcare provider about the availability and importance of genetic counseling and have been provided with written information identifying a genetic counselor or medical geneticist who can provide such counseling services. All my questions have been answered and I have had the necessary time to make an informed decision about the genetic test(s). I hereby give permission to Baylor Genetics to conduct genetic testing as recommended by my physician.										
Patient's Printed Name		Patient's Signature		/ 	/) / YYYY)					
Patient's Parent / Personal Representative*	Name I	Patient's Parent / Personal Repre	sentative Signature	/ 	/) / YYYY)					
				/	/					
Relationship of Personal Representative to	the Patient (Ordering Provider's Signature		Date (MM / DE) / YYYY)					

^{*}If you are signing as a person with legal authority to act on behalf of the patient, you may be required to provide evidence of your authority.