

PHONE 1.800.411.4363 FAX 1.800.434.9850

CONNECT





# **POSTNATAL CMA / CYTOGENETICS REQUISITION**

PATIENT INFORMATION (COMPLETE	E ONE FORM FOR EACH PERSON TESTED)			
				/
Patient Last Name	Patient First Name		MI	Date of Birth (MM / DD / YYYY)
Address	City	State Patient discharged from	Zip Genetic Sex:	Phone
Accession #	Hospital / Medical Record #	the hospital/facility: Yes No	Female Gender identity (if different	) Male ( ) Unknown from above):
REPORTING RECIPIENTS				
Ordering Physician		Institution Name		
Email (Required for International Clien	its)	Phone	Fax	
ADDITIONAL RECIPIENTS				
Name		Email	Fax	
Name		Email	Fax	
PAYMENT (FILL OUT ONE OF THE O	PTIONS BELOW)			
SELF PAYMENT				
Pay With Sample	Bill To Patient			
O INSTITUTIONAL BILLING				
O INSTITUTIONAL BILLING				
Institution Name	Institution Code Inst	itution Contact Name In	stitution Phone	Institution Contact Email
O INSURANCE				
Do Not Perform Test Until Pa	atient is Aware of Out-Of-Pocket Costs (exclud	es prenatal testing)		
REQUIRED ITEMS 1. Copy of	f the Front/Back of Insurance Card(s) 2. ICD10 D	Diagnosis Code(s) 3. Name of Ordering	g Physician 4. Insured Si	gnature of Authorization
	/ /			1 1
Name of Insured	/ / / / / /	: Name of Insured	Insu	/ // red Date of Birth (MM / DD / YYYY)
Name of modera	insured bate of Birth (init) / BB / TTTT	i Name of moured	11130	rea bate of birth (Min 7 bb 7 1111)
Patient's Relationship to Insured	Phone of Insured	Patient's Relationship to	Insured Pho	ne of Insured
Address of Insured		Address of Insured		
City	State Zip	City	Stat	e Zip
Primary Insurance Co. Name	Primary Insurance Co. Phone	Secondary Insurance Co.	Name Seco	ondary Insurance Co. Phone
Primary Member Policy #	Primary Member Group #	Secondary Member Polic	y# Sec	ondary Member Group #
understand that I am responsible for a reasons including, but not limited to, r	Baylor Genetics to provide my insurance c any co-pay, co-insurance, and unmet deductib non-covered and non-authorized services. I u in payment for this test. Please note that Med	ole that the insurance policy dictates, Inderstand that I am responsible for	as well as any amounts sending Baylor Genetics	not paid by my insurance carrier for
				//
Patient's Printed Name	Patient's S	Signature		Date (MM / DD / YYYY)
STATEMENT OF MEDICAL NECESSI	TY (REQUIRED)			
patient's medical management and tr	ne risk assessment, diagnosis, or detection o eatment decisions. The person listed as the C to the patient and they have consented to ger	Ordering Physician is authorized by la		
				//
Physician's Printed Name	Physician	s Signature		Date (MM / DD / YYYY)



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# **POSTNATAL CMA / CYTOGENETICS REQUISITION**

					/	/	
Patient Last Nan	ne	Patient First Name		MI	Date of Birtl	(MM / DD / YYYY	Genetic Sex
ETHNICITY							
African Ame Ashkenazi J East Asian ( Finnish French Cana	lewish China, Japan, Korea)	Native America	(Saudi Arabia, Qata n	ar, Iraq, Turkey) andinavian, UK, Gern	Sou Sou Sou	ıth Asian (India, P ıtheast Asian (Vie	opines, Micronesia, Malaysia, Indonesia) akistan) tnam, Cambodia, Thailand) Caucasian (Spain, Italy, Greece)
INDICATION FO	R TESTING (REQUIRED)						
CMA OPTIONS	•••••	•••••	•••••••	(	CHROMOSOME/FISH	OPTIONS	
Autism Spectrum Failure to Thrive  Developmental Delay Multiple Congenital Anomalies  Dysmorphic Features Seizure Disorder  Other (Specify):			<ul><li>☐ Autosomal Trisomies</li><li>☐ Ambiguous Genitalia</li><li>☐ Fetal Demise</li><li>☐ Other (Specify):</li></ul>			☐ Infertility ☐ Sex Chromosome Abnormalities ☐ Multiple Miscarriages	
ICD10 Diagnosis	Code(s):						
SAMPLE INFOR	RMATION						
Date of Collectio	/ n (MM / DD / YYYY)	s	AMPLE TYPE ···  Blood in EDTA  Skin Biopsy		Buccal Swab Extracted DNA fro	m	Skin Fibroblast
	L MICROARRAY ANALYSIS						
Products of Cond	ception (POC) and fetal tissue	tests should be re	quested using the	"Cytogenetics - Prod	ucts of Conception Red	quisition", which c	an be found at baylorgenetics.com.
TEST CODE						SPECIFY REGION OF INTEREST	
	Chromosomal Microarray Analysis (CMA) - HR + SNP Screen (Comprehensive)  Chromosomal Microarray Analysis (CMA) - HR + SNP Screen BE + BH, SF, SB, BUC only or DNA						
8655	BE + BH, SF, SB, BUC only or DNA  BE + BH, SF, SB, BUC only or DNA						
For Chromosomal I	Microarray Analysis tests, the sam	ple types BE+BH are	preferred. BUC and DI	NA are also acceptable s	sample types.		
PARENTAL STU	IDIES RECOMMENDED IN C	HILD'S CMA REP	ORT (ATTACH CO	PY)			
Mother	First, MI, Last	Date of Bi	// rth (MM/DD/YYYY)		ASYMPTOMATIC	SYMPTOMA	ATIC (Specify:)
Father	,,		, ,	(	ASYMPTOMATIC	SVMPTOMA	ATIC (Specify:)
	First, MI, Last	Date of Bi	// rth (MM/DD/YYYY)			O 31111 10111	
	FICATIONS TABLE						
ABBREVIATION	SAMPLE NAME	RECOMMEND (2 YRS - ADULT)	DED AMOUNT (NEWBORN - 2 YRS)	SHIPPING INSTRUCTIONS		SPECIAL NOTES	
BE	Blood in EDTA tube (purple-top)	3 - 5 cc	2 - 3 cc	Ship at room temperature overnight courier. Do not h	in an insulated container by		
ВН	Blood in Sodium Heparin tube (green top)	3 - 5 cc	1 - 2 cc	,	in an insulated container by		
BUC	Buccal Swab	See "Special Notes"	See "Special Notes"	Baylor Genetics with instructions). We high collected by a healthcare professional.  Baylor Genetics with instructions). We high collected by a healthcare professional.  Buccal swab is an accepted sample type fo		accepted sample type for Chromosomal Microarray s 8665 or 8655) and FMR1 CGG Repeat Expansion	
CB         Cord Blood         N/A         1 - 2 cc			Ship at room temperature overnight courier. Do not h	in an insulated container by leat or freeze.	Ensure properly labeled. Also send 3 cc of maternal blood in p labeled EDTA tube for MCC studies at no charge as needed.		
DNA	DNA, Extracted	10 - 15 ug	10 - 15 ug	Ship at room temperature overnight courier. Do not h	in an insulated container by leat or freeze.	Minimal concentrat	tion of 50ng/uL; A260/A280 1.75-2.0
SA	Saliva	See "Special Notes"	See "Special Notes"	Ship at room temperature overnight courier. Do not h	in an insulated container by leat or freeze.	Collected with Orag Genetics with instr	gene DNA Self-Collection Kit (provided by Baylor uctions).



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## POSTNATAL CMA / CYTOGENETICS PENIISITION

POSTNAT	AL CMA / CYTOGENETICS REQUISITION								
			//						
Patient Last Na	me Patient First Name	MI	Date of Birth (MM / DD / YYYY)	Genetic Sex					
CYTOGENETIC TESTS									
Products of Conception (POC) and solid tissue tests should be requested using the Cytogenetics - Products of Conception Requisition, which can be found at baylorgenetics.com									
TEST CODE	ST CODE TEST NAME								
8600	Chromoso	ome Analysis		BH					
8480	FISH for SRY - Related Phenotype	es (Metaphase & Interph	nase Cells) **	ВН					
** Testing on m	etaphase cells requires cell culturing.								
NOTE: The follo	wing tests (8425 and 8426) REQUIRE selecting an accompanying te	st (8665, 8655, or 8600	)						
TEST CODE	TEST	NAME		SAMPLE TYPE*					
8425	Rapid FISH - AneuVysion (+13/+1	18/+21/X/Y) (Interphase	e cells ONLY)	ВН					
8426	Rapid FISH - Sex Chromosome	s (X/SRY) (Interphase c	ells ONLY)	ВН					
		+							
TEST CODE	TEST	NAME		SAMPLE TYPE*					
8665	Chromosomal Microarray Analysis (Ci	BE + BH, SF, SB, BUC only or DNA							
8655	Chromosomal Microarray	BE + BH BUC or DNA							
8600	Chromosome Analysis BH								
CMA + FMR1									
	buccal swab sample is needed if test codes 8665 and 6573 are ord								
TEST CODE	TEST	NAME		SAMPLE TYPE*					
8665	Chromosomal Microarray Analysis (CMA) - HR + SNP Screen (Comprehensive)  BE + BH SF, SB, BUC only or								
6573	FMR1 CGG Repeat Expansion Analysis			BE, BUC, SAL, or DNA					
If negative, reflex to:									
TEST CODE	TEST NAME								
1500	Proband Whole Exome Sequencing								
1600	Trio Whole Exome Sequencing								
1602	Additional Affected Sibling for Trio*								
2055 Comprehensive mtDNA Analysis by Massively Parallel Sequencing (MitoNGS <sup>SM</sup> )									
* The Sibling Trio should be ordered along with, or after a completed Trio (#1600) for the same biological family.  Note: Please include the WES Advantage requisition and consents.									
FISH STUDIES	FISH STUDIES								

Products of Conception (POC) and fetal tissue tests should be requested using the "Cytogenetics - Products of Conception Requisition", which can be found at baylorgenetics.com/requisitions/

TEST	T CODE	TEST NAME	SAMPLE TYPE	TES	T CODE	TEST NAME	SAMPLE TYPE
	8462	Charcot-Marie-Tooth Neuropathy Type 1A	ВН		8474	Neurofibromatosis Type I	ВН
	8440	DiGeorge/Velocardiofacial Syndrome (22q and 10p) Panel	ВН		8480	SRY-Related Phenotypes	ВН
	8486	DiGeorge/Velocardiofacial Syndrome Type I (22q)	ВН		8485	X-Linked Ichthyosis	ВН
	8465	DiGeorge/Velocardiofacial Syndrome Type II (10p)	ВН		8490	Chromosome X and Y Centromere Analysis	ВН
	8467	Hereditary Neuropathy w/ Liability to Pressure Palsies	ВН		*8405	Custom Familial FISH Studies	ВН

 $<sup>^*</sup>$ Note: Please include the previous report and note the region of interest. Contact the lab to confirm appropriate probe coverage is available.

<sup>\*</sup> Refer to Sample Specifications Table (page 2)



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#### INFORMED CONSENT FOR POSTNATAL CMA / CYTOGENETICS TESTING

Patient Last Name	Patient First Name		/ Date of Birth (MM / I	/ DD / YYYY)	Genetic Sex
TEST INFORMATION					
	ith information regarding genetic testing, ding the reason for this testing, we have p				
or their family. DNA is the genetic ma our body. Each person has a unique s	etermine if a genetic disease may be pres sterial that we receive from our parents. G set of DNA and most of the differences in o alled variants) that might cause disease, m	enes are made of ur DNA do not im	f DNA and are the inpact our health. G	instructions for ma enetic testing analy	intaining the health of zes DNA to find any
The testing ordered by your healthca mean your unborn child, for the purp	re provider can determine if you or your coses of this consent.	hild have a variar	nt associated with	a genetic disease.	"Your child" can also
Depending on why genetic testing is	needed, you might be tested for:				
<ul> <li>A known variant that has already be</li> <li>A single gene or variant that cause</li> <li>Multiple genes at the same time. T</li> <li>Multiple types of testing that each</li> </ul>	es a specific, suspected disease. hese genes might cause similar diseases	or might cause d	iseases that are ui	nrelated to each otl	ner.

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).

RESULTS .....

- 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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INFURMED CONSENT FOR	PUSINAIAL CMA / CYTUG	ENETICS TEST						
Patient Last Name	Patient First Name	MI	// Date of Birth (MM / DD / YYYY)	Genetic Sex				
ratient Last Name	Fatient First Name	IVII	Date of Bit (II (MIM / DD / 1111)	Genetic Sex				
PATIENT CONFIDENTIALITY AND SE	PECIMEN RETENTION (CONT.)							
	e, however in rare cases, inaccurate f clinical/medical information, or rar		Reasons for this include, but are not l	imited to, mislabeled				
cancel the test. If you wish to ca	ncel testing, the laboratory must be r	notified of the cance	an contact the healthcare provider w llation request before 5 PM CST the b ntil after this time, you will be charge	ousiness day after the				
• 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.								
enacted several laws that prohi		est results by health	e coverage and employment. The U.S insurance companies and employers www.genome.gov/10002077.					
• Samples will be retained in the	aboratory in accordance with the lab	oratory retention p	olicy.					
	es. DNA specimens are not returned t		elopment and improvement, internal eferring heath care providers unless					
	York State will not be included in res ple. No tests other than those author		out your written consent and will not be med on the biological sample.	e retained for more than				
submission serves to contribute	knowledge to the medical communit	ty. I understand that	e submitted to public databases, such : limited clinical information is also re nformation may, although unlikely, in	equired for the submission				
	t identifies the underlying genetic cau se management or treatment of disea		n your family, this information may no	ot help in predicting the				
FINANCIAL AGREEMENT AND GUAR	ANTEE							
billing, I hereby authorize Baylor G to my insurance carrier which is re of appealing any denial of benefits directly to Baylor Genetics. I under part of a verification of benefits in health insurance plan. If my insura- to endorse the insurance check as	enetics to bill my health insurance pleasonably required for billing. I addit by my insurance carrier. I irrevocab stand that my out-of-pocket costs may estigation. I agree to be financially reince provider sends a payment direct appropriate and forward such checks rendered. If I do not have health insu	an on my behalf, an ionally designate B ly assign associated a be different than esponsible for all ar lly to me for unpaid to Baylor Genetics	tic testing ordered by my healthcare in durther authorize Baylor Genetics to aylor Genetics as my designated reprofunction and to the estimated amount indicated to mounts as indicated on the explanation services performed by Baylor Genetics within thirty (30) days of receipt there are for the full cost of the genetic testing the services are for the full cost of the genetic testing the services are for the full cost of the genetic testing the services are for the full cost of the genetic testing the services are for the services	o release any information esentative for purposes ect that payment be made e by Baylor Genetics as n of benefits issued by my cs on my behalf, I agree eof, as payment towards				
I understand that a completed Adv	ance Beneficiary Notice (ABN) is requ	uired for Medicare p	atients if the service is deemed not m	nedically necessary.				
RECONTACT FOR RESEARCH CONS	ENT							
contact patients or their provider(s research involving the sample(s) a	s) directly as part of this research. I a	gree to allow Baylo is testing. I underst	opment, and other scientific purposes r Genetics to contact me or my provid and that patients generally receive no t baylorgenetics.com.	er(s) about possible				
If I wish to opt out of being reconta	cted for research purposes by Baylor	r Genetics, I unders	tand that I may check the box below:					
☐ Please do not contact me regard	ing any research that uses information	on obtained from th	is testing.					
For any research I may be contactor will be made via secure email if po		following methods	(please check all that apply – if no ch	oices are selected, contact				
□Email □Phone □Mail								



BAYLOR GENETICS 2450 HOLCOMBE BLVD. GRAND BLVD. RECEIVING DOCK

PHONE 1.800.411.4363 FAX CONNECT





# HOUSTON, TX 77021-2024 1.800.434.9850

			/	/	
Patient Last Name	Patient First Name	MI	Date of Birth (MM	/ DD / YYYY)	Genetic Sex
PATIENT AUTHORIZATION					
By signing this statement of consent appropriate explanations from my hyprovider about the availability and it or medical geneticist who can provide informed decision about the genetic of the provided in th	nealthcare provider about mportance of genetic condessuch counseling servertest(s).	ut the planned genetic test(s ounseling and have been pro vices. All my questions have	) and possible results. I h vided with written inform been answered and I hav	nave been informed b nation identifying a g	oy my healthcare enetic counselor
Patient's Printed Name		Patient's Signature		Date (M	/ M / DD / YYYY)
Patient's Parent / Personal Representative	e* Name	Patient's Parent / Personal R	Representative Signature	Date (M	// M / DD / YYYY)
Relationship of Personal Representative to	o the Patient	Ordering Provider's Signatur	re	Date (M	// M / DD / YYYY)

INFORMED CONSENT FOR POSTNATAL CMA / CYTOGENETICS TESTING

<sup>\*</sup>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.