

POSTNATAL CMA / CYTOGENETICS REQUISITION
PATIENT INFORMATION (COMPLETE ONE FORM FOR EACH PERSON TESTED)

Patient Last Name	Patient First Name	MI	Date of Birth (MM / DD / YYYY)
Address	City	State	Zip
Accession #	Hospital / Medical Record #	Patient discharged from the hospital/facility: <input type="radio"/> Yes <input type="radio"/> No	
		Genetic Sex: <input type="radio"/> Female <input type="radio"/> Male <input type="radio"/> Unknown Gender identity (if different from above): _____	

REPORTING RECIPIENTS

Ordering Physician	Institution Name
Email (Required for International Clients)	Phone
	Fax

ADDITIONAL RECIPIENTS

Name	Email	Fax
Name	Email	Fax

PAYMENT (FILL OUT ONE OF THE OPTIONS BELOW)

SELF PAYMENT
 Pay With Sample Bill To Patient

INSTITUTIONAL BILLING

Institution Name	Institution Code	Institution Contact Name	Institution Phone	Institution Contact Email
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INSURANCE
 Do Not Perform Test Until Patient is Aware of Out-Of-Pocket Costs (excludes prenatal testing)

REQUIRED ITEMS 1. Copy of the Front/Back of Insurance Card(s) 2. ICD10 Diagnosis Code(s) 3. Name of Ordering Physician 4. Insured Signature of Authorization

Name of Insured	Insured Date of Birth (MM / DD / YYYY)	Name of Insured	Insured Date of Birth (MM / DD / YYYY)
Patient's Relationship to Insured	Phone of Insured	Patient's Relationship to Insured	Phone of Insured
Address of Insured		Address of Insured	
City	State	Zip	
City	State	Zip	
Primary Insurance Co. Name	Primary Insurance Co. Phone	Secondary Insurance Co. Name	Secondary Insurance Co. Phone
Primary Member Policy #	Primary Member Group #	Secondary Member Policy #	Secondary Member Group #

By signing below, I hereby authorize Baylor Genetics to provide my insurance carrier any information necessary, including test results, for processing my insurance claim. I understand that I am responsible for any co-pay, co-insurance, and unmet deductible that the insurance policy dictates, as well as any amounts not paid by my insurance carrier for reasons including, but not limited to, non-covered and non-authorized services. I understand that I am responsible for sending Baylor Genetics any and all payments that I receive directly from my insurance company in payment for this test. Please note that Medicare does not cover routine screening tests.

Patient's Printed Name	Patient's Signature	Date (MM / DD / YYYY)
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STATEMENT OF MEDICAL NECESSITY (REQUIRED)

This test is medically necessary for the risk assessment, diagnosis, or detection of a disease, illness, impairment, symptom, syndrome, or disorder. The results will determine my patient's medical management and treatment decisions. The person listed as the Ordering Physician is authorized by law to order the test(s) requested herein. I confirm that I have provided genetic testing information to the patient and they have consented to genetic testing.

Physician's Printed Name	Physician's Signature	Date (MM / DD / YYYY)
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POSTNATAL CMA / CYTOGENETICS REQUISITION

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

ETHNICITY

- | | | |
|--|---|---|
| <input type="radio"/> African American | <input type="radio"/> Hispanic American | <input type="radio"/> Pacific Islander (Philippines, Micronesia, Malaysia, Indonesia) |
| <input type="radio"/> Ashkenazi Jewish | <input type="radio"/> Mennonite | <input type="radio"/> South Asian (India, Pakistan) |
| <input type="radio"/> East Asian (China, Japan, Korea) | <input type="radio"/> Middle Eastern (Saudi Arabia, Qatar, Iraq, Turkey) | <input type="radio"/> Southeast Asian (Vietnam, Cambodia, Thailand) |
| <input type="radio"/> Finnish | <input type="radio"/> Native American | <input type="radio"/> Southern European Caucasian (Spain, Italy, Greece) |
| <input type="radio"/> French Canadian | <input type="radio"/> Northern European Caucasian (Scandinavian, UK, Germany) | <input type="radio"/> Other (Specify): _____ |

INDICATION FOR TESTING (REQUIRED)
CMA OPTIONS

- | | |
|---|--|
| <input type="checkbox"/> Autism Spectrum | <input type="checkbox"/> Failure to Thrive |
| <input type="checkbox"/> Developmental Delay | <input type="checkbox"/> Multiple Congenital Anomalies |
| <input type="checkbox"/> Dysmorphic Features | <input type="checkbox"/> Seizure Disorder |
| <input type="checkbox"/> Other (Specify): _____ | |

CHROMOSOME/FISH OPTIONS

- | | |
|---|---|
| <input type="checkbox"/> Autosomal Trisomies | <input type="checkbox"/> Infertility |
| <input type="checkbox"/> Ambiguous Genitalia | <input type="checkbox"/> Sex Chromosome Abnormalities |
| <input type="checkbox"/> Fetal Demise | <input type="checkbox"/> Multiple Miscarriages |
| <input type="checkbox"/> Other (Specify): _____ | |

ICD10 Diagnosis Code(s): _____

SAMPLE INFORMATION

 _____ / _____ / _____
 Date of Collection (MM / DD / YYYY)

SAMPLE TYPE

- | | | | |
|---------------------------------------|------------------------------------|--|------------------------------|
| <input type="radio"/> Blood in EDTA | <input type="radio"/> Buccal Swab | <input type="radio"/> Blood in Sodium Heparin | <input type="radio"/> Saliva |
| <input type="radio"/> Skin Fibroblast | <input type="radio"/> Skin Biopsy* | <input type="radio"/> Extracted DNA from _____ | |

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.

CHROMOSOMAL MICROARRAY ANALYSIS (CMA) TESTS

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.

TEST CODE	TEST NAME	SAMPLE TYPE*	SPECIFY GENE OF INTEREST	SPECIFY REGION OF INTEREST
<input type="checkbox"/> 8665	Chromosomal Microarray Analysis (CMA) - HR + SNP Screen (Comprehensive)	BE + BH, CB, SF, SB, BUC only or DNA		
<input type="checkbox"/> 8655	Chromosomal Microarray Analysis (CMA) - HR (Basic)	BE + BH, SF, SB, BUC only or DNA		

For Chromosomal Microarray Analysis tests, the sample types BE+BH are preferred. BUC and DNA are also acceptable sample types.

PARENTAL STUDIES RECOMMENDED IN CHILD'S CMA REPORT (ATTACH COPY)

- | | | |
|---|------------------------------------|--|
| <input type="checkbox"/> Mother _____ / _____ / _____
First, MI, Last Date of Birth (MM/DD/YYYY) | <input type="radio"/> ASYMPTOMATIC | <input type="radio"/> SYMPTOMATIC (Specify): _____ |
| <input type="checkbox"/> Father _____ / _____ / _____
First, MI, Last Date of Birth (MM/DD/YYYY) | <input type="radio"/> ASYMPTOMATIC | <input type="radio"/> SYMPTOMATIC (Specify): _____ |

SAMPLE SPECIFICATIONS TABLE

ABBREVIATION	SAMPLE NAME	RECOMMENDED AMOUNT		SHIPPING INSTRUCTIONS	SPECIAL NOTES
		(2 YRS - ADULT)	(NEWBORN - 2 YRS)		
BE	Blood in EDTA tube (purple-top)	3 - 5 cc	2 - 3 cc	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze.	
BH	Blood in Sodium Heparin tube (green top)	3 - 5 cc	1 - 2 cc	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze.	
BUC	Buccal Swab	See "Special Notes"	See "Special Notes"	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze.	Collect with ORACollect•Dx (OCD-100) self-collection kit (provided by Baylor Genetics with instructions). We highly recommend the sample be collected by a healthcare professional. Buccal swab is an accepted sample type for Chromosomal Microarray Analysis (test codes 8665 or 8655) and FMR1 CGG Repeat Expansion Analysis (test code 6573).
CB	Cord Blood	N/A	1 - 2 cc	Ship at room temperature in an insulated container by 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	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze.	Minimal concentration of 50ng/uL; A260/A280 1.75-2.0
SAL	Saliva	See "Special Notes"	See "Special Notes"	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze.	Collected with Oragene DNA Self-Collection Kit (provided by Baylor Genetics with instructions).
SF	Cultured Skin Fibroblast	2 T25 flasks	2 T25 flasks	Ship at room temperature in an insulated container by overnight courier. Do not heat or freeze.	Send 2 T25 flasks at 80-100% confluence.
SB	Skin Biopsy	5mm^3	5mm^3	Ship at ambient temperature (18-25°C/64-77°F). Protect paraffin tissue from excessive heat. Ship in cooled container during summer months. Sample must arrive within 72 hrs.	Collect 5 cubic millimeters of skin from a central location (e.g., buttock or upper thigh) rather than from a distal location (e.g., foot) to enhance cell viability. Place sample in a separate sterile container with RPMI media. In the absence of RPMI media, place sample in a sterile container with a small amount of sterile saline. Unacceptable Conditions: Specimens placed in formalin or other fixatives.

* 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.

POSTNATAL CMA / CYTOGENETICS REQUISITION

Patient Last Name _____ 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	TEST NAME	SAMPLE TYPE*
<input type="checkbox"/> 8600	Chromosome Analysis	BH
<input type="checkbox"/> 8480	FISH for SRY - Related Phenotypes (Metaphase & Interphase Cells) **	BH

** Testing on metaphase cells requires cell culturing.

NOTE: The following tests (8425 and 8426) REQUIRE selecting an accompanying test (8665, 8655, or 8600)

TEST CODE	TEST NAME	SAMPLE TYPE*
<input type="checkbox"/> 8425	Rapid FISH - Aneuvysion (+13/+18/+21/X/Y) (Interphase cells ONLY)	BH
<input type="checkbox"/> 8426	Rapid FISH - Sex Chromosomes (X/SRY) (Interphase cells ONLY)	BH



TEST CODE	TEST NAME	SAMPLE TYPE*
<input type="checkbox"/> 8665	Chromosomal Microarray Analysis (CMA) - HR + SNP Screen (Comprehensive)	BE + BH, SF, SB, CB, BUC only or DNA
<input type="checkbox"/> 8655	Chromosomal Microarray Analysis (CMA) - HR (Basic)	BE + BH BUC or DNA
<input type="checkbox"/> 8600	Chromosome Analysis	BH

CMA + FMR1 TESTING
NOTE: Only one buccal swab sample is needed if test codes 8665 and 6573 are ordered together.

TEST CODE	TEST NAME	SAMPLE TYPE*
<input type="checkbox"/> 8665	Chromosomal Microarray Analysis (CMA) - HR + SNP Screen (Comprehensive)	BE + BH, SF, SB, CB, BUC only or DNA
<input type="checkbox"/> 6573	FMR1 CGG Repeat Expansion Analysis	BE, BUC, SAL, or DNA

If negative, reflex to:

TEST CODE	TEST NAME
<input type="checkbox"/> 1500	Proband Whole Exome Sequencing
<input type="checkbox"/> 1600	Trio Whole Exome Sequencing
<input type="checkbox"/> 1602	Additional Affected Sibling for Trio*
<input type="checkbox"/> 2055	Comprehensive mtDNA Analysis by Massively Parallel Sequencing (MitoNGS SM)

* 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

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

*Note: Please include the previous report and note the region of interest. Contact the lab to confirm appropriate probe coverage is available.

* Refer to Sample Specifications Table (page 2)

INFORMED CONSENT FOR POSTNATAL CMA / CYTOGENETICS TESTING

Patient Last Name _____

Patient First Name _____

MI _____

Date of Birth (MM / DD / YYYY) _____ / _____ / _____

Genetic Sex _____

GENERAL GENETIC TESTING CONSENT

This consent form cannot be used for whole exome sequencing (WES), whole genome sequencing (WGS), biochemical testing, or Huntington disease testing. Consent forms for other tests are located at Baylor Genetics' website (<https://www.baylorgenetics.com/consent/>).

For the purposes of this consent, "I", "my", "you", and "your" can refer to you, your child, your unborn child, or other individual you are the legal representative of.

TEST INFORMATION

Your healthcare provider (doctor, genetic counselor, or other person with medical training) wants to order one or more tests to find a cause for your health issues. This testing can see if there is a cause for your health issues or if there is an increased chance for a health issue to happen to you or your family. Some of these tests look for changes, called variants, in a person's DNA. DNA is our genetic material. You might have testing for variants in one or more genes, specific parts of DNA that are needed for our health. Variants can also be found in other places in the genome (all of the DNA that a person has). Some tests might look for changes in proteins or analytes that cause health issues. The testing ordered will depend on your health issues as well as what is already known about you and your family's genetics. These tests may also explain health issues that your family may have. Even if this test finds the cause of your health issues, this may not help treat or manage those issues.

Before you sign this consent form, you should speak with your healthcare provider. They can help you understand this testing and what it means for your health.

TEST RESULTS

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

- **Positive:** A variant in the DNA was found that is related to your health issues or a health issue that you are at an increased risk of having in the future. These changes that cause disease are also known as pathogenic variants.
- **Negative:** No variants in the DNA were found that are related to your health issues or that would increase your risk of a health issue in the future.
- **Variant of Uncertain Clinical Significance (VUS):** A variant in the DNA was found that we do not know its effect, if any, on health. More testing may be needed for you or your family if a VUS is found that may be associated with your health issues.
- **Secondary and Incidental Findings:** Testing can sometimes find a variant in the DNA not related to the reason for testing. If this result is expected to affect your health, it is called a secondary or incidental finding.

CONSIDERATIONS AND LIMITATIONS

- You should speak with your provider before signing this consent form to understand the risks, benefits, and alternatives to testing.
- Testing may show you have, or are at increased chance of having, a health issue. It may show that you have an increased chance of having a child with a health issue.
- Even if the variant(s) causing your health issues are found, how these issues might progress or improve with treatment might not be known. Affected family members with the same variant might not be affected like you are.
- Depending on the results of testing, more testing may be needed to understand these results. This testing might be needed for you and/or other family members.
- A negative result does not rule out the chance for health issues. Our knowledge of variants and how they cause disease may change over time as we learn more about genetics. Testing has limitations to what it can find as well.
- Certain factors may lead to incorrect results. These include mislabeled samples, incorrect information in the test order, and rare technical errors.
- More sample may be needed from you if the first sample is not sufficient to complete testing.

PATIENT CONFIDENTIALITY AND SAMPLE RETENTION

- If several family members are tested, knowing the correct biological relationships among them is important. In rare cases, testing can show that family members are not related as expected. If this is found, we may contact the provider who ordered your testing.
- If this testing is requested to be cancelled after the order and sample are sent to the laboratory, please see our Test Cancellation Policy at www.baylorgenetics.com/cancel-test/.
- Only Baylor Genetics and its contracted partners will have access to your sample for the ordered testing. Results from testing will only be released to: (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. You have the right to access your test results from Baylor Genetics by providing a written request. You also have the right to request raw data obtained from your sample by providing a written request or HIPAA Authorization Form.
- In rare cases, people with genetic diseases may have problems with health insurance and employment. The U.S. Federal Government has several laws that prohibit discrimination based on test results by health insurance companies and employers. These laws also prohibit unauthorized disclosure of this information. For more information, please visit www.genome.gov/10002077.
- Samples will be kept in the laboratory based on our retention policy. Once testing completes, de-identified sample may be used for test development, quality assurance, and training purposes. Samples are not returned to patients or providers unless requested prior to testing. You and your heirs will not receive payments, benefits, or rights to any resulting products or discoveries.
- The information from your testing may be used in scientific research, publications or presentations, but your specific identity will not be revealed. We may contact your provider to obtain more clinical information about you. Baylor Genetics also performs other types of scientific research and may contact you to see if you would like to be involved.
- Variants found may be submitted to databases. The medical community uses these databases to collect information about how variants might cause disease to improve testing and treatment for patients. An example is ClinVar, a free, public archive of reports on human genetics. Limited clinical information may need to be shared with these databases. In rare cases, this information may be enough to allow you or your family members to be identified.
- For more information on privacy practices at Baylor Genetics, please visit www.baylorgenetics.com/privacy-practices/.

INFORMED CONSENT FOR POSTNATAL CMA / CYTOGENETICS TESTING

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

USE OF DATA AND SPECIMEN FOR RESEARCH PURPOSES

Biological specimens, test results, and associated information may be used by Baylor Genetics and its research partners for anonymous or coded research purposes, including improving genetic testing, advancing knowledge of genetic conditions, and developing new technologies, including inclusion in de-identified clinical databases, only with the patient's informed consent. Patient data and specimen will not be used for anonymous or coded research, unless authorized by marking below. A patient's decision to decline participation shall not affect their ability to receive testing from Baylor Genetics.

For Oregon patients, please consult the state specific consent form found at www.baylorgenetics.com/forms.

I authorize Baylor Genetics the use of my specimen and de-identified data for research.

FOR SAMPLES FROM NEW YORK STATE RESIDENTS

Samples from New York State residents shall not be included in research without written consent. Samples will not be retained for more than sixty (60) days after receipt by Baylor Genetics, unless authorized by marking below. No tests other than those authorized shall be performed on the samples.

I authorize Baylor Genetics to retain sample(s) longer based on our retention policy for test development, quality assurance, and training purposes.

FINANCIAL AGREEMENT

By signing below, I hereby authorize Baylor Genetics to provide my insurance carrier any information necessary, including test results, for processing my insurance claim. I understand that I am responsible for any co-pay, co-insurance, and unmet deductible that the insurance policy dictates. I 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. Please note, some payers may not cover certain screening tests.

If my health insurer does not cover the test or I do not have health insurance, I have received a good faith estimate of the cost for the genetic testing ordered by my provider and agree to pay for the cost of the genetic testing billed to me by Baylor Genetics based on that good faith estimate. More information is available in Baylor Genetics' No Surprises Act and Good Faith Estimate Notice located at <https://www.baylorgenetics.com/no-surprises-act/>.

A Medicare Advance Beneficiary Notice (ABN) is required for services Medicare identifies as not medically necessary.

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 Name Patient's Signature Date Signed (MM / DD / YYYY)

 Patient's Parent / Personal Representative* Name Patient's Parent / Personal Representative Signature Date Signed (MM / DD / YYYY)

*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 or parent, you may be required to provide evidence of your authority.