

CYTOGENETICS - PRODUCTS OF CONCEPTION 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 _____ Phone _____
 Accession # _____ Hospital / Medical Record # _____
 Patient discharged from the hospital/facility: Yes No
 Biological Sex: Female Male 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 _____

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) _____

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) _____



CYTOGENETICS - PRODUCTS OF CONCEPTION REQUISITION

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

- Recurrent Pregnancy Loss (N96)
- Maternal Care for Intrauterine Death, Not Applicable or Unspecified (O36.4XX0)
- Encounter for Elective Termination of Pregnancy (Z33.2)
- Missed Abortion (O02.1)
- Blighted Ovum and Nonhydatidiform Mole (O02.0)
- Other Abnormal Products of Conception (O02.89)
- Other (Specify ICD-10 code):

GESTATIONAL INFORMATION

Date of Pregnancy Loss: _____ / _____ / _____
MM DD YYYY

Gestational Age at Loss (weeks): _____

- Type of Pregnancy:
- Singleton Pregnancy
 - Multiple Gestation

- Was an Egg Donor Used During This Pregnancy?
- Yes
 - No

CYTOGENETIC TESTS

8800 | CHROMOSOMES (TISSUE)

SAMPLE TYPE

- Fresh Tissue
- POC
- Villi

DATE OF COLLECTION

_____ / _____ / _____
MM DD YYYY

8639 | CMA - POC

SAMPLE TYPE

- Fresh Tissue
- POC
- Villi

DATE OF COLLECTION

_____ / _____ / _____
MM DD YYYY

- Frozen Tissue

- Autopsy Material

FFPE Slides #: _____

FFPE Blocks #: _____

- Maternal Cell Contamination Studies
(RECOMMENDED For Test Code 8639)

Mother Last Name _____ Mother First Name _____ MI _____

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

- Blood Sample (5 cc in EDTA Tube) collected from Biological Mother

NOTE: Fresh tissue samples should be submitted in sterile media whenever possible (see BMGL.com for specific handling instructions).

INFORMED CONSENT FOR CYTOGENETICS - PRODUCTS OF CONCEPTION TESTING

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

INFORMED CONSENT FOR GENETIC TESTING

BACKGROUND

You are considering the genetic test called Chromosomal Microarray Analysis (abbreviated CMA) for your current pregnancy. The purpose of this document is to provide information about the test so that you can decide whether it is right for you. This information is meant to be used in addition to your discussion with your physician or a genetic counselor. If you decide to have the CMA test, you will be asked to sign at the bottom of this document, indicating that you understand the information provided and wish to have testing. You will be given a copy of this document for your records.

Having the proper amount of genetic material (DNA) in each cell of the body is important for normal growth and development. The DNA is located along the 23 pairs of chromosomes (46 total) in each cell. A basic karyotype analysis can only detect the number of chromosomes in each cell and large structural changes in these chromosomes. CMA is an advanced method of looking at the structure and number of the chromosomes in our body because CMA is able to detect the large changes identified by karyotype, as well as detecting smaller regions of any missing or extra copies (copy number variant, or CNV). These smaller CNVs can also cause abnormal development.

In addition, the CMA test can detect an abnormal inheritance pattern of the chromosomes called uniparental disomy (UPD). The CMA test can also detect regions of genetic similarity, called absence of heterozygosity (AOH).

TEST REPORTING

There are several categories of results that may be reported: these include 1) No clinically significant CNV detected (normal result); 2) Clinically significant CNV detected, known to be associated with a genetic condition; 3) CNV detected in the fetus but also detected in a parent. Based on our experience thus far, this has been seen in about 10% of cases. It is generally of low concern, but should be discussed with a genetic counselor; and 4) Variation of uncertain significance detected in the fetus, but not present in either parent. This is relatively rare (seen thus far in about 1-2% of cases) and requires detailed discussion with a physician or genetic counselor.

In addition, regions of genetic similarity (AOH) may be reported if the CMA results indicate the possibility of uniparental disomy (UPD) or consanguinity. If a clinically significant abnormality has been detected, your physician or genetic counselor will discuss the information with you. A clinical geneticist (a specialist in the medical impact of genetic information) may also be consulted.

INFORMATION AND CONSENT FOR TESTING

- (1) While the CMA test is very accurate, it is possible that your fetus could have one of the medical conditions included in the CMA test that is not detected or that your fetus could have a medical condition, which cannot be detected by the CMA. This is possible because many genetic syndromes have more than one cause.
- (2) Due to the fact that many different regions of the chromosomes and many different conditions are being analyzed, there is a risk that you will learn genetic information about yourself, your fetus, or your family that is not directly related to the reason for monitoring your pregnancy. This information might relate to diseases with symptoms that may develop in the future in your fetus or possibly yourself or other family members. Gains and Losses associated with adult-onset dementia disorders will NOT be reported. See below for options regarding receipt of certain categories of results in the report.
- (3) As with any genetic test, results may be unclear and additional studies may be recommended in order to give you the most accurate information about what the lab finding may mean for the health of your fetus.
- (4) It is possible that additional information may come to light during these studies, such as family relationships not being as expected. Because interpretation of CMA results may involve study of the biological parents to determine significance of the findings, the interpretation may not be accurate if specimens from the biological parents are not available for comparative study. If pregnancy was achieved through use of an egg or sperm donor, it is important to inform your physician/genetic counselor so they can work with you and the laboratory to assure the most accurate analysis possible. Your doctor or genetic counselor may be able to coordinate obtaining samples from an egg or sperm donor if necessary.
- (5) The CMA test will be performed using materials and protocols developed at the BGL and validated by the laboratory. This laboratory is certified by standards set by the Clinical Laboratory Improvement Acts (CLIA) and the College of American Pathologists.

Due to the nature of the methodology of this testing we are unable to guarantee that all CNVs in each option will be detected by the CMA. The below options apply to the reporting of fetal data.

There are some findings that may be identified by CMA that do not directly impact the health of your pregnancy, but are considered medically actionable because they have clear medical significance to the health of the fetus later in life. Parental studies will NOT automatically be run for this category of reporting since it will not aid in the interpretation of the fetal result. The American College of Medical Genetics (ACMG) has published guidelines for the reporting of these types of medically actionable or incidental findings (PMID: 23788249, 27854360). These guidelines include a list of genes, which may be updated periodically. In accordance with an update to this policy statement (PMID: 25356965), there is the option to receive pathogenic variant information if identified in one of the listed genes. Please note that if one of these is part of a larger copy number event that meets criteria for reporting then the below option will NOT apply and the data will be reported.

