

GLOBAL MAPS® 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) _____



GLOBAL MAPS® REQUISITION

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

INDICATION FOR TESTING (REQUIRED)

ICD10 Diagnosis Code(s):

For most accurate results, patient should not be on TPN, special diet, dietary supplements, or drug therapies. Please list all medications and/or supplements the patient has been prescribed and is currently taking:

Please provide the following clinical information regarding the patient to be tested. This information is needed to facilitate interpretation of metabolic profiling results. If the laboratory requires additional information, please indicate the healthcare provider to be contacted:

Physician Name _____ Physician Phone/Pager # _____

INDICATION CHECKLIST

INDICATION	YES*	NO	UNKNOWN
Abnormal Movements	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Ataxia	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Autism/Autistic Spectrum	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Delayed Motor Milestones	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Delayed Speech	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Developmental Regression	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Dietary Avoidances	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Dysmorphic Features	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Eye Problems	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Failure to Thrive	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Family History of Similar Disorder	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Genital Anomalies	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
GI/Liver Problems	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Hearing Loss	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Heart Problems	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Hyperextensibility	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Hypertonia/Spasticity	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Hypotonia	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Intellectual Disability	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

INDICATION	YES*	NO	UNKNOWN
Intrauterine Growth Restriction	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Joint Contractures	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Kidney Problems	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Lethargy	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Leukodystrophy	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Macrocephaly	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Microcephaly	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Obesity/Overgrowth	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Organomegaly	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Prematurity	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Seizure Disorder	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Short Stature	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Skeletal Abnormalities	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Skin Anomalies	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Structural Brain Abnormalities	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Tall Habitus	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Unusual Odor	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Vomiting	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

* If YES, please provide description below:

PREVIOUS TESTING

- Metabolic Testing
(e.g.: Newborn screening, amino acid analysis)
- Chromosomal Microarray Analysis (CMA)
- Genetic Analysis

If checked, please provide additional details about previous testing in the box below:

TESTING LOCATION

- Baylor Genetics
- Lab # _____ Family # _____
- Another laboratory (Attach a copy of the test results)

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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): _____ |

REQUIRED ITEMS CHECKLIST

- | | |
|---|---|
| <input type="checkbox"/> Indication for Study Checklist | <input type="checkbox"/> Proband Sample |
| <input type="checkbox"/> Clinical Note / Summary | <input type="checkbox"/> Requisition |

GLOBAL MAPS® TESTS

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

TEST CODE	TEST NAME	SAMPLE TYPE *
<input type="checkbox"/> 4900	Global Metabolomic Assisted Pathway Screen ¹	PE
<input type="checkbox"/> 4901	Global Metabolomic Assisted Pathway Screen	U

¹ Was plasma extracted from EDTA? Yes No

(REQUIRED When ordering Test Code 4900)

REPORTING

Turnaround time is 3 weeks after financial responsibility has been verified to receive the focused report. Once the focused report is received, the expanded report can be ordered (no additional charge). A requisition for ordering the expanded report is available on our website. Please allow 2 weeks for the expanded report. For more details regarding the reporting system, please visit BMGL.com or call 800-411-GENE.

SAMPLE SPECIFICATIONS TABLE

ABBREVIATION	SAMPLE NAME	RECOMMENDED AMOUNT		SHIPPING INSTRUCTIONS	SPECIAL NOTES
		(2 YRS - ADULT)	(NEWBORN - 2YRS)		
PE	Plasma (from EDTA)	1 - 2 cc	1 - 2 cc	Ship frozen sample in insulated container, with 3 -5 lbs dry ice, by overnight courier.	Draw blood in an EDTA (purple top) tube(s) and separate as soon as possible, freezing immediately. Send 1 -2 cc of plasma. Store the specimen frozen at -20°C. Specimen may be stored frozen up to 7 days.
U	Urine	3 - 5 cc	2 - 4 cc	Ship frozen sample in insulated container, with 3 -5 lbs dry ice, by overnight courier.	Collect random urine. Do not add preservatives. Store the specimen frozen at 20°C.

ADDITIONAL STUDIES - RESEARCH

After your results are finalized and reported there may be research studies that you may be eligible for and may be of interest to you. Please read the following statement and select the appropriate box. If the "YES"/contact option is chosen, please complete the additional information requested. Please note that if neither box is selected, the lab will default to the "NO" contact option.

YES Baylor Genetics may share my contact information with researchers who have an Institutional Review Board (IRB) approved research study for which I may be eligible for participation. There is no obligation to participate if contacted. No information, other than the contact information below, will be provided to the researcher.

INITIAL _____

Printed Name _____ Signature _____ Date (MM / DD / YYYY) _____

Relationship to Patient _____ Patient Name _____ Preferred Method of Contact:

- Email Mail Phone

Phone # _____ Alternative Phone # _____ Email _____

Address _____ City _____ State _____ Zip _____

NO I DO NOT wish to be contacted regarding participation in research studies.

*Refer to Sample Specifications Table above

INFORMED CONSENT FOR GLOBAL MAPS® TESTING

Patient Last Name

Patient First Name

MI

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

Biological Sex

INFORMED CONSENT FOR GENETIC TESTING

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. In order to ensure that you have understood the purpose and significance of genetic testing, we have provided information about the testing process and potential results below.

The purpose of genetic testing is to identify the cause of a suspected disease in you or your family. The testing analyzes your genetic material (DNA) for an abnormal change (variant) that could explain the disease you or members of your family are experiencing. Genetic testing can be a diagnostic test, which is used to identify or rule out a specific genetic condition. Genetic screening tests are used to assess the chance for a person to develop or have a child with a genetic condition. Genetic screening tests are not typically diagnostic, and results may require additional testing.

The purpose of this test is to see if you or your child may have a genetic variant or chromosome rearrangement. This may cause a genetic disorder or may determine the chance that you or your child will develop or pass on a genetic disorder in the future. "Your child" can also mean your unborn child, for the purposes of this consent.

In a genetic test, depending on the case, you can be tested for:

- A single gene/variant responsible for a specific, suspected genetic disease.
- Multiple genes in parallel.

The sample/specimen that is needed to perform the genetic test is stated in the test order form and is typically blood or purified DNA, but may also be tissue, saliva or buccal swab.

RESULTS

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

- **Positive:** Positive or "abnormal" results mean there is a change in the genetic material found that is related to your/your child's medical issues or that you/your child are at an increased risk of developing the disorder in the future. It is possible to test positive for more than one genetic variant.
- **Negative:** Negative or "normal" results mean no relevant genetic change related to your/your child's medical issues was detected. This does not mean there is no genetic change, but it may mean that the type of testing performed could not detect it.
- **Results of Unclear Significance:** Testing can detect change(s) in DNA which we do not yet fully understand. These alterations are also referred to as variants of uncertain significance (VUS). Additional studies may be recommended if a VUS is identified in a gene that may be associated with your/your child's medical concerns.
- **Secondary / Incidental Findings:** Testing can sometimes detect a change in a person's DNA unrelated to the reason for testing. If this change has medical or reproductive significance, it is called a secondary or incidental finding.

CONSIDERATIONS AND LIMITATIONS

- Results may indicate affected status, increased risk to someday be affected with, and/or reproductive risk for a genetic disorder. It is important to understand that genetic tests, even if negative, are not exhaustive. It is not possible to exclude risks for all possible genetic diseases for yourself and your family members.
- A positive test result is an indication that the individual(s) being tested may be predisposed to or have the specific disease or condition which prompted testing. You might consider additional independent testing, consult a personal physician, or pursue genetic counseling.
- It is possible that the knowledge of the test results may result in psychological stress for you and your family. It is always recommended to discuss the results with your healthcare provider or genetic counselor.
- If several family members are tested, the correct interpretation of the results depends on the provided relationships between 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 discrepancy is identified, it may be necessary to report this to the physician who ordered the testing.
- Genetic testing is highly accurate. Rarely, inaccurate results may occur for various reasons. These reasons include, but are not limited to, mislabeled samples, inaccurate reporting of clinical/medical information, or rare technical errors.
- If you sign this consent form, but you no longer wish to have your sample(s) tested, you can contact your physician to cancel the test. If testing is complete, but you have not received your results yet, you can inform your physician that you no longer wish to receive the results. If you withdraw consent for testing after 5pm CST the next business day following sample receipt by the laboratory, you will be charged for the full cost of the test.

PATIENT CONFIDENTIALITY AND SPECIMEN RETENTION

- Results will only be released to a licensed healthcare provider, to those allowed access to test results by law, and to those authorized in writing.
- In rare cases, persons with genetic diagnoses have experienced problems with insurance coverage and employment. The U.S. Federal Government has enacted several laws that prohibit discrimination based on genetic test results by health insurance companies and employers. In addition, these laws prohibit unauthorized disclosure of this information. For more information, you can visit www.genome.gov/10002077.
- Samples will be retained in the laboratory in accordance with the laboratory retention policy.
- After testing is complete, the de-identified submitted specimen may be used for test development and improvement, internal validation, quality assurance, and training purposes. DNA specimens are not returned to individuals or to referring health care providers unless specific prior arrangements have been made.



INFORMED CONSENT FOR GLOBAL MAPS® TESTING

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

INFORMED CONSENT FOR GENETIC TESTING

PATIENT CONFIDENTIALITY AND SPECIMEN RETENTION (CONT.)

- Samples from residents of New York State will not be included in the de-identified research studies described in this authorization and will not be retained for more than 60 days after test completion, unless specifically authorized by your selection. No tests other than those authorized shall be performed on the biological sample.
- Information including results, indications for testing and clinical status obtained from this testing may be shared with healthcare providers, scientists and healthcare databases or used in scientific publications or presentations, but the personal identifying information of all persons studied will not be revealed in such data sharing or publications/presentations.

RESEARCH & RECONTACT CONSENT

For more information on research at Baylor Genetics, please visit baylorgenetics.com. Please read the below statements carefully and check the appropriate box.

Note: If left blank, consent is interpreted as "NO."

- I agree to use of my de-identified specimen for research to improve genetic testing for all patients and contribute to scientific research.
 - I am a New York State Resident, and I give Baylor Genetics permission to store my specimen in accordance to the laboratory retention policy for internal quality assurance and possible research studies.
- In addition to agreeing above, I agree to be contacted by Baylor Genetics regarding research opportunities.

PATIENT AUTHORIZATION

By signing this statement of consent, I acknowledge that I have read and understand the informed consent for genetic testing. I have received appropriate explanations from my physician regarding the purpose, scope, type and significance of the planned genetic testing and achievable results. All my questions have been answered and I have had the necessary time to make an informed decision about the genetic test.

I give permission to Baylor Genetics to conduct genetic testing as recommended by my physician.

Patient Signature Date (DD/MM/YYYY)

Printed Name