

Patient and Specimen Information Form

5424 Glenridge Drive NE | Atlanta, GA 30342 USA | phone: 844.664.8378 | fax: 678.225.0212 | mnglabs.com

| | Patient and | l Specin | nen Infor | mation | | | |
|--|--|-----------------------|------------------------------|--------------------------------|---|--|--|
| Patient Last Name | | | | Patient First Name | | | |
| Patient ID # | | | | Date of Birth [MM/DD/YYYY] | | | |
| Diagnosis/ICD-10 | | | Collection Date [MM/DD/YYYY] | | | | |
| Gender ☐ Male ☐ Female | Specimen Type Whole Blood Buccal Swab | □ CSF □ Urine □ Fibro | e oblasts | ☐ Plasma/ Serum ☐ Muscle | □ DNA Tissue: | | |
| Please complete and include clinical information form, or attach clinical notes | | | | | | | |
| Referring Physician Information | | | | | | | |
| Physician Name | | | | | NPI # or equivalent (Required) | | |
| Facility / Organization | Signature | | | | | | |
| Facility Address City, State, Zip Code Same billing | as | | | | | | |
| Report Delivery □ Fax | ☐ Email | | | | Phone | | |
| Billing Information (REQUIRED) | | | | | | | |
| Self-Pay? ☐ Yes If yes, | | | <u> </u> | - | ust be received in full prior to testing. | | |
| Facility | • | | | | | | |
| Billing Address | | | | Name | | | |
| | | | | | | | |
| City, State, Zip Code | | | | | | | |
| Phone | Fax | | Email | | | | |
| | | | | | | | |
| Authorized | | Res | ults Authorized | | | | |
| Recipient Name | | | Recipient Name | | | | |
| Facility Phone | | | Facility | | Phone | | |
| ☐ Fax | | | ☐ Fax | | | | |
| □ Email | | | ☐ Email | | | | |
| Testing Checklist | | | | | | | |
| All of the following are encouraged to be included with test orders (please check the following): | | | | | | | |
| ☐ All specimens that will be analyzed must be received - please note if samples will ship separately ☐ Clinical Information Form completed ☐ Informed Consent for Genetic Testing completed and signed | | | | | | | |



Next Generation Sequencing Test Request Form

| 5424 Glenridge Drive NE Atlanta, GA 30342 USA phone | e: 678.225.0222 fax: 678.225.0212 mnglabs.labcorp.com | | | | | |
|---|---|--|--|--|--|--|
| Patient Name | DOB | | | | | |
| Epilepsy | | | | | | |
| ☐ (NGS412) Myoclonic Epilepsy | | | | | | |
| Movement Disorders | | | | | | |
| Ataxia / Episodic Ataxia | | | | | | |
| ☐ (NGS324) Ataxia/Episodic Ataxia Disorders + mtDNA | | | | | | |
| Dystonias | | | | | | |
| ☐ (NGS358) Comprehensive Dystonia + mtDNA | ☐ (NGS360) Basal Ganglia Calcification Dystonia | | | | | |
| ☐ (NGS357) Parkinsons Disease/Parkinsonism | (NOSOSO) Basar Carigita Calomoation Bystonia | | | | | |
| | | | | | | |
| Neuromuscular Neuromuscular | | | | | | |
| Muscular Dystrophy / Myopathy | | | | | | |
| ☐ (NGS330) Comprehensive Muscular Dystrophy/Myopathy + mtDNA | (NGS413) Congenital Myopathies | | | | | |
| ☐ (NGS331) Congenital Myasthenic Syndromes☐ (NGS332) Hypokalemic & Hyperkalemic Periodic Paralysis | ☐ (NGS424) Duchenne/Becker Muscular Dystrophy ☐ (NGS333) Malignant Hyperthermia | | | | | |
| Neuropathies | _ (), | | | | | |
| ☐ (NGS445) Comprehensive Neuropathies | ☐ (NGS323) Amyotrophic Lateral Sclerosis | | | | | |
| ☐ (NGS346) Hereditary Sensory & Autonomic Neuropathy | (NGS405) Amyotrophic Lateral Scierosis + C9orf72 Repeat Expansion | | | | | |
| ☐ (NGS400) Pain Syndromes | ☐ (NGS337) Spastic Paraplegia + mtDNA | | | | | |
| ☐ (NGS345) Charcot-Marie-Tooth Disease + mtDNA | | | | | | |
| | | | | | | |
| Neurob | ehavioral | | | | | |
| Intellectual Disability / Autism | | | | | | |
| ☐ (NGS325) Comprehensive Intellectual Disability/Autism + mtDNA | (NGS398) Macrocephaly & Overgrowth Syndrome | | | | | |
| (NGS349) Nonsyndromic Intellectual Disability | ☐ (NGS425) Microcephaly | | | | | |
| ☐ (NGS350) Syndromic Intellectual Disability | | | | | | |
| Neurodegeneration | | | | | | |
| ☐ (NGS376) Comprehensive Dementia | (NGS356) Alzheimer Disease/Frontotemporal Dementia | | | | | |
| Brain Malformation Disorders | ☐ (NGS380) Amyloid Related Disorders | | | | | |
| ☐ (NGS372) Comprehensive Leukodystrophy/ Leukoencephalopathy + | ☐ (NGS387) Comprehensive Neuronal Migration Disorders + mtDNA | | | | | |
| mtDNA | (NOODOT) COMPTENENSIVE NEGROTIAN MIGRATION DISORDERS - MILESTAT | | | | | |
| | | | | | | |
| Neurometabolic Neurometabolic | | | | | | |
| (NGS301) Comprehensive Cellular Energetics Defects + mtDNA | ☐ (NGS313) Lysosomal Disease | | | | | |
| ☐ (NGS302) Carbohydrate Metabolism Deficiency + mtDNA | (Nocoro) Eysosomai biscase | | | | | |
| Other Inherited Disorders | | | | | | |
| Vision and Ophthalmoplegia | Other | | | | | |
| ☐ (NG464) Comprehensive Vision Loss & Eye Disorders + mtDNA | ☐ (NGS429) Familial Hemiplegic Migraine + mtDNA | | | | | |
| ☐ (NGS352) Comprehensive Ophthalmoplegia Syndromes + mtDNA | ☐ (NGS430) Stroke + mtDNA | | | | | |
| Connective Tissue Disorders | (NGS379) Polycystic Kidney Disease | | | | | |
| (NGS377) Ehlers Danlos, Ehlers Danlos-like Syndromes, and | (NGS392) Bartter/Gitelman Syndromes | | | | | |
| Aneurysm Syndromes Multi-Sensory Disorders | ☐ (NGS335) Neurofibromatosis ☐ (NGS428) Tuberous Sclerosis | | | | | |
| ☐ (NGS402) Usher Syndrome | (NGS319) Fever Syndromes | | | | | |
| ☐ (NGS460) Alport Syndrome | | | | | | |



Clinical Information Form

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Patient Name _ DOB Clinical (Check All That Apply) ☐ Retinitis Pigmentosa Hearing **Neuronal Migration** ☐ Optic Atrophy Eye ☐ Sensorineural ☐ Stickler ☐ Usher ☐ Joubert ☐ Other ☐ Stroke ☐ Other ☐ Intellectual Disability (ID) ☐ Syndromic ID Cognitive/Neurobehavioral ■ Nonsyndromic ID ☐ Autism Dementia ☐ Ataxia ☐ Episodic Ataxia ☐ Dystonia ☐ Chorea/Athetosis ☐ Parkinson Disease ☐ L-Dopa Response **Movement Disorders Connective Tissue & Bone** Epilepsy ☐ Myoclonic ☐ Other **Spasticity** ☐ Absence ☐ Tonic Clonic ☐ Spastic Paraplegia ☐ Other ☐ Ehlers Danlos ☐ Marfan ■ Aneurysms ☐ Epileptic Encephalopathy ☐ Spastic Quadriplegia ☐ Other **Nerve/Anterior Horn Cell** Neuromuscular Distal ☐ Proximal ■ Muscle Atrophy ☐ Contractures ☐ Neurofibromas ☐ Charcot-Marie-Tooth ☐ Sensory Rhabdomyolysis ■ Malignant Hyperthermia ☐ Arthrogryposis ☐ Autonomic ☐ Pain ☐ Motor ☐ Nerve Conduction ☐ Periodic Paralysis ☐ Statin Use ■ Myasthenia ☐ Other **Arrhythmias Congenital Heart Defects** Cardiomyopathy **Endocrine** ☐ Ventricular Tachycardia
☐ Brugada ☐ Heterotaxy ☐ Dilated ☐ Hypertrophic ☐ Other ☐ Hypothyroidism ☐ Long or Short QT ☐ Conduction Defect ■ Noncompaction ☐ Other ☐ Diabetes Mellitus Imaging (Check All That Apply) Brain MRI **EEG (Describe Findings) EMG/NVC** (Describe Findings) Leigh Disease ☐ Basal Ganglia Calcification ☐ Stroke ☐ Cerebellar Atrophy ☐ Abnormal Myelin (describe) Laboratory Metabolic (Describe Findings) **Genetic (Describe Findings)** ☐ Chromosomal Microarray ■ Deletion/Insertion Testing ☐ Other (comment) **CPK** Maximum ___ Minimum _ **Family History** Ethnicity (please check) ☐ Caucasian ☐ Sephardic Jewish ☐ African American (or Black) ☐ Asian Hispanic ☐ Ashkenazi Jewish ☐ Native American (or American Indian) Other: **Affected Maternal Lineage Affected Paternal Lineage Siblings** Relationship to Proband Relationship to Proband Number (specify gender) Symptoms Symptoms Healthy/Affected **Additional Comments**



Informed Consent for Genetic Testing

In compliance with New York State Civil Law: Section 79-L

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|----|--|--|--|--|--|
| | Patient Name | DOB | | | |
| | Please provide a copy of <u>completed</u> consent | with sample and requisition. Failure to do so may delay testing. | | | |
| W | hen signed and dated, this written consent is written au | thorization to participate in genetic testing. | | | |
| 1. | | e recommended testing: (name o | | | |
| | test or MNG test code), which is performed to help di | agnose(insert disease description) | | | |
| | | ng, including the description of the purpose, methodology, and disorders is ither been reviewed with me by my physician or I have read the documentation | | | |
| 2. | disease tested for. A negative result may/may not rule | result is an indication that the individual has a genetic cause for the specific out a genetic disorder depending on clinical history and quality/type of specimes dependent testing, consult a personal physician or pursue genetic counseling. | | | |
| 3. | Level of Certainty: Is test-specific and determined by the methods employed, patient's clinical history and sometimes by the natu of the patient's condition at time of sampling. There is always a small possibility of error or failure in sample analysis; this is true wi complex testing in any laboratory. Inclusion of clinical data, such as medical history, family history, images as they relate to the diseas or disorder, will decrease the level of uncertainty in an interpretation and are encouraged to be included when submitting samples f analysis. MNG Laboratories will keep personal information private in accordance with HIPAA laws. I consent to the retention of these documents by MNG Laboratories in their database. Patient (or parent/guardian) Initials: | | | | |
| 4. | Disclosing Test Results: The following categories of persons or organizations that test results may be released to include, but are not limited to: hospitals or laboratories involved in the patient's care, referring physician(s) and primary care providers, other physician groups (consultants, surgeons), insurance companies (as provided by the patient or referring physician for payment purposes and other professionals involved in patient care that assist MNG Laboratories in carrying out treatment, payment, and operational activities. Results are kept confidential. Medical Neurogenetics complies with security and privacy statutes of the federal Healt Information Portability and Accountability Act (HIPAA). If a patient chooses to specifically declare where results may be released (other than the referring institution and ordering physician), please provide these <i>in writing</i> to the Compliance Officer, MNG Laboratories (quickresponse@mnglabs.com). | | | | |
| 5. | No clinical tests other than those authorized shall be my referring physician or other authorized healthcare MNG Laboratories will retain the samples for longer p assurance processes. I consent to have my specimens retained afte | not return any remaining sample to individuals or physicians unless requested be performed on the sample. A request for additional testing must be made by a professional and there will be an additional charge. If agreed by the patient periods for use in an anonymous fashion for research/development or for quality are completion of initial testing (this consent may be withdrawn at any paining sample). Patient (or parent/guardian) Initials: | | | |
| 6. | Testing for Genetic Conditions can be Complex: If understand what the risks and benefits are to having to understand that a biologic specimen will be obtained. | warranted, obtain professional genetic counseling prior to giving consent to full; the testing completed. I hereby consent to participate in testing described above I from me and/or members of my family. I understand that this biologic speciment if I, or members of my family, are affected or are carriers of a particular disease. | | | |
| | Signature of Patient | Date | | | |
| | Authorized Signature (Parent/Guardian) | Relationship | | | |
| | Name of Patient (please print clearly) | Name of Ordering MD (please print clearly) | | | |
| | Referring Facility (please print clearly) | Signature of Ordering MD | | | |
| | | | | | |

Important: One signature from patient (or parent/guardian), authorized person, or physician is required to complete this form. New York requires signatures from patient (or parent/guardian) OR ordering physician to complete this form.

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