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INSTRUCTIONS: Please review all of the below sections carefully. Your test ordering option will be indicated based on your selection made in the "Test Catalog" section. Please include as much supporting documentation and information as you can, and fill out all fields for which you are able to do so. If you have any questions or concerns, please reach out to our office for clarification.

COMPREHENSIVE NEUROLOGY TESTING REQUISITION FORM

ORDERING DETAILS

1. ELIGIBILITY CRITERIA

This test is available to individuals 18 or older who are suspected of or at risk of having a Comprehensive Neurology disorder based on one or more of the following (*please select all that apply*):

REQUIRED:

To avoid processing delays, you must select the appropriate eligibility criteria for the patient from the provided list of options below.

- o Amyotrophic Lateral Sclerosis
- o Ataxia
- o Autism Spectrum Disorder
- o Cerebral Cavernous Malformation
- o Charcot-Marie-Tooth Neuropathy
- o Coenzyme Q10 Deficiency
- o Cognitive Impairment
- o Collagen Type VI-Related Disorders
- o Epilepsy
- o Muscular Dystrophy/Myopathy
- o Congenital Myasthenic Syndrome(s)
- o Creatine Metabolism Deficiency
- o Dementia
- o Dystonia
- o Emery-Dreifuss Muscular Dystrophy
- o Epileptic Encephalopathy
- o Holoprosencepahly
- o Idiopathic Generalized and Focal Epilepsy
- o Leukodystrophy/Leukoencephalopathy
- o LGMD/Congenital Muscular Dystrophy

- o Lissencephaly
- o Macrocephaly/Overgrowth Syndrome
- o Metabolic Epilepsy
- o Metabolic Myopathy/Rhabdomyolysis
- o Microcephaly/Pontocerebellar Hypoplasia
- o Migraines
- o NCL/Progressive Myoclonic Epilepsy
- o Nemaline Myopathy
- o Neuro-Ophthalmology
- o Neuronal Migration Disorder
- o Parkinson Disease
- o Periodic Paralysis
- o Polymicrogyria
- o Porphyria
- o Septo-Optic Dysplasia
- o Spastic Paraplegia
- o Spinal Muscular Atrophy
- o Tuberous Sclerosis
- o X-Linked Intellectual Disabilities

Does the patient have a family member with a known disease-causing variant in one of the genes included on the Amyotrophic Lateral Sclerosis Panel, the Dementia Panel, or the Parkinson Disease Panel?

If yes, please list the family member relationship to patient, gene, and variant, if known:

2. GENE-SPECIFIC FAMILY FOLLOW-UP TESTING

For relatives of testing participants who received a Pathogenic/Likely Pathogenic result, or approved VUS, who would like to receive gene-specific follow-up testing at no additional charge

Relatives do not need to meet the eligibility criteria listed above. If participating in gene-specific follow-up testing, please check the following box.

PATIENT INFORMATION				ORDERING PHYSICIAN & CLINIC INFORMATION				
First Name	me MI Last Name				Clinic/Practice Name			
Date of Birth (MM/DD/YYYY) Biological Sex MRN (medical record number, if available)			Phone		Fax			
			ucasian 🗌 Ashkenazi Jewish	Street Address		C	City	
Hispanic I Native American Pacific Islander French Canadian Sephardic Jewish Mediterranean Other:				State/Prov	ZIP/Postal	Country (if	f other than US)	
Phone Email address			Primary Clinic Conta	ict Name		Extension (if applicable)		
Street Address City			Primary Clinic Conta	ct Title				
State/Prov ZIP/Postal Country (if other than US)			Ordering Provider (Please select one physician per order.)					
					Physician Name		Physician NPI	
				□				
				□				
SPI	ECIMEN IN	FORMATIO	ON	□				
SPECIMEN INFORMATION Specimen type (please select and indicate number):			□					
Buccal Swab(s) - (OCD-100 kit)			□					
				□				
Number of Swabs Provided:				□				
Specimen collection date (MM/DD/YYYY):								

CLINICAL HISTORY (It is strongly enco					VEC	NO			
Cognitive Features Progressive cognitive decline -	YES	NO	UNKNOWN	Motor Features (continued) Progressive muscle weakness	YES	NO □			
amnestic presentation (memory loss, impairment in learning and recall)				and/or atrophy					
Progressive cognitive decline -			_	Muscle fasciculations and/or cramps					
language presentation (word-finding deficits)				Hyporeflexia and/or decreased or absent deep tendon reflexes					
Progressive cognitive decline - visuospatial presentation (spatial cognition-object agnosia, facial recognition, simultagnosia and alexia)				Parkinsonism (bradykinesia, postural instability, rigidity, facial masking, resting tremor) Tardive dyskinesia (irregular, jerky					
Progressive cognitive decline - executive dysfunction (impaired reasoning, judgment and problem solving)				movements), dystonia (patterned/ twisting movements and postures) and/or myoclonus (muscle jerks)					
Behavioral changes (disinhibition/				Dysarthria (difficulty speaking)					
impulsivity, apathy/inertia, and/or loss of sympathy/empathy)				Dysphagia (swallowing difficulties)					
Behavioral changes				Neuroimaging, biomarkers, genetic, and	/or neuropath	ophysiolo	gy findings:		
(perseverative/compulsive behaviors and/or hyperorality)				Abnormal MRI Major finding(s)?					
Psychiatric illness (psychosis, mania, hallucinations, delusions, etc.)				 Abnormal PET scan or CSF analysis Major finding(s)? 					
Other:				△ Abnormal brain pathology findings					
Motor Features				Major finding(s)?			,		
Cerebellar ataxia (gait and/or limb ataxia)				Abnormal EMG Major finding(s)?					
Oculomotor dysfunction				□ Previously tested for C9orf72 gene					
(ex: oculomotor apraxia, strabismus, and/or nystagmus)				Test result (positive, negative, inter [please include copy of previous te		ble]			
Increased muscle tone and/or increased extremity deep-tendon reflexes/ hyperreflexia (jaw jerk, Hoffman sign, positive Babinski sign, crossed adductors, extensor plantor response)				Other features Paget disease of bone Supranuclear palsy					
Spasticity				Autonomic dysfunction (ex: orthost	atic hypotensic	on, urinary i	ncontinence)		
Pseudobulbar affect (inappropriate laughing/crying/forced yawning)				Familial insomnia Other relevant clinical features:					
Noticed yawning/ OPTIONAL - REQUESTED VARIANTS FOR THIS PATIENT'S REPORT, IF KNOWN To have the presence or absence of specific variants commented on in this patient's report, provide the details requested below. <i>For gene-specific family follow-up, see the note following test selection menu.</i> Was the proband (individual with variant) tested at Elite Clinical Laboratory? If yes, list Accession ID: If no, attach a copy of the test results (required). Variant(s) Requested (e.g. GENE c.2200A>T (p.Thr734Ser) NM_00012345)									
				IT INFORMATION					
INSURANCE BILLING To avoid delays, Policy Holder Name To avoid delays,	please include	Patient	Relationship to Pol	surance card, along with clinical notes, medical rec icy Holder Self			al necessity (LMN). PAY/SELF-PAY		
Primary Insurance Name	Other:			Primary Insurance Group # Have you, the patier hospital inpatient (n			ould prefer to pay dit card, please		
Secondary Insurance Name	Secondary Insurance Policy #		#	four [24] hour stay) in Secondary Insurance Group #	the last fourteen (14	4) include Credit C	e a fully filled out ard Authorization		
	Secondary insurance Name Form with this test order form.								
CLINICAL HISTORY									
FAMILY HISTORY Is there a family history of the disease(s) for which the patient is being tested? Yes No If yes, describe below and attach pedigree and/or clinical notes.									
Relative / Relationship to the patient Maternal or paternal Diagnosed	•		Age at diagnosis		condition	-	Age at diagnosis		

Is/was this patient affected or symptomatic? ^{$†$} \Box Yes \Box No Provide details in the required clinical history questions (if applicable).	[†] Symptomatic means this patient has features or signs known or suspected to be related to the genetic testing being ordered and could include findings on physical examination, laboratory tests, and/or imaging.
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PERSONAL HISTORY

COMPREHENSIVE NEUROLOGY TEST CATALOG

Custom Neurology NGS Testing (Select the genes below) or Comprehensive Neurology NGS Testing Panel (Test All Genes)

□ADNP □AFF2 □ALDH7A1 □ANG □APTX □ARX □ASPA □ASXL1 □ATN1 □ATP1A2 □ATP7B □ATXN1	□ATXN10 □ATXN2 □ATXN3 □ATXN7 □ATXN8OS □BCL11A □BSCL2 □C12orf4 □CACNA1A □CACNA1C □CC2D1A □CCL5	CHD2 CNOT3 CNTN6 COL4A1 COL4A3BP CSNK2A1 CSTB CTNND2 DHCR7 DPYD EGR2 CEHMT1	DEN2 DEZH2 DFBX011 DFMR1 DFOXG1 DFTSJ1 DFTSJ1 DFXN DGABRG2 DGAMT DGARS DGATM	□GBA □GCH1 □GRIN2A □GRN □HEXA □HFE □HSPB1 □HTT □IKBKAP □KCNQ2 □KDM5C □L1CAM	DLRRK2 MAPT MBOAT7 MECP2 MED12 MTHFR MTM1 NDP NDUFA1 NLGN3 NLGN4X NOTCH3	 NNSD1 NTRK1 NTRK2 PABPN1 PCDH19 PDGFB PDHA1 PIK3CA PINK1 PMP22 PNKD POLG 	DPPP2R2B DPRRT2 DPSEN1 DPTEN REEP1 DSCN1A DSCN1B DSCN2A SCN8A DSCO2 DSGCE DSLC16A2	SLC2A1 SLC6A8 SLC9A6 SMN1 SMN2 SOD1 SPG11 STXBP1 STXBP1 TARDBP TBP TCF4	□TH □THAP1 □TOR1A □TSC1 □TSC2 □TTR □UBA1 □ZEB2 □ZNF41
INDICATI	ON (S) FOR	TESTING		ICD-1	0 Codes				
Inflammatory of ICD Category - 1 G00.0 Herr G00.1 Pnete G00.3 Stap G00.8 Oth G00.9 Bact ICD Category - 1 G01 G01 Men G02 Men ICD Category - 1 G03 G03.0 Non G03.1 Chrc G03.2 Beni G03.9 Men G03.9 Men G03.9 Men G04.01 Post G04.02 Post G04.03 Max G04.10 Post G04.31 Post G04.32 Post G04.33 Uth G04.34 Post G04.39 Oth G04.39 Oth G04.39 Oth G04.48 Oth G05.4 Myce ICD Category - 1 G05.3 G05.4 Myce	liseases of the ce Bacterial mening tomococcal mening phylococcal mening obylococcal mening er bacterial mening traial meningitis in bacteria Meningitis in bacteria Meningitis in bacteria Meningitis in other inf Meningitis in other inf Meningitis due to other ingitis, unspecifie Encephalitis, unspecifie Encephalitis, unspecified Encephalitis and ence distinfectious acute of stinfectious acute of the disseminated en infectious acute of stinfectious acute of stinfectious acute of stinfectious acute of the disseminated en- infectious acute of the disseminated en- infectious acute of stinfectious acute of the disseminated en- infectious acute of the disseminated en- infectious acute of the disseminated en- tifications acute of the disseminated en- ter acute necrotizing en acute necrotizing en acute necrotizing the disseminated en- tifications acute of the disseminated en- tification acute of the disseminated en- tification acute of the disseminated en- tificat	ntral nervous syste itis, not elsewhere is gitis gitis gitis nspecified terial diseases classified infec/parastc dise fectious and parasiti o ther and unspec- tits ingitis [Mollaret] er specified causes d elitis and encephal incephalitis and encephal incephalitis and encephal incephalitis and encephal inte disseminated encephal inte disseminated encephal its and encephalomyelitis phalomyelitis, unsp 5 & encephalomyeli tis and encephalomyelitis phalomyelitis, unsp 6 & encephalomyeli tis and encephalomyelitis phalomyelitis, unsp 6 & encephalomyeli tis and encephalomyelitis 7 maspinal abscess 7 maspinal phlebiti pinal abscess of cen 7 maspinal phlebiti 7 maspinal phlebiti pinal abscess of cen 7 maspinal phlebiti 7 maspinal phlebiti	classified (G00) ssified elsewhere elsewhere ases class d elsewhere c diseases classifie c diseases (G03) omyelitis (G04) ephalomyelitis, un halitis and encephal cephalitis, myelitis gomyelitis, not els agic encephalopat rrhagic encephalopat rrhagic encephalopat rrhagic encephalopat cephalopat cephal	(G01) r (G02) d elsewhere) specified alomyelitis encephalomyelitis ewhere classified thy spathy liswhr (G05) classified elsewhere G06) elswhr (G07) ies nlebitis (G08) r ease	ICD Catego G13.0 G13.2 G13.2 G13.4 ICD Catego G14 Extrapyran ICD Catego G20 ICD Catego G21.10 G21.2 G21.3 G21.3 G21.4 G21.3 G21.4 G21.3 G21.4 G21.3 G21.4 G23.0 G23.1 G23.2 G23.3 G23.4 G24.3 G24.1 G24.2 G24.3 G24.4 G24.3 G24.4 G24.3 G25.0 G25.1 G25.2 G25.3 G25.4 G25.5 G25.4 G25.70 G25.71 G25.72 G25.74 G25.75 G25.70	Paraneoplastic neuro Other systemic atrop in neoplastic disease Systemic atrophy pri Systemic atrophy pri Systemic atrophy pri Systemic atrophy pri Parkinson's disease Parkinson's disease Postencephalitic par Vascular parkinsonis Other secondary parkinsori Other degenerative Secondary parkinsori Other degenerative Striatonigral degene Other specified degene Other specified degene Other drug induced Genetic torsion dystri Idiopathic orofacial of Blepharospasm 8 Other dy Dystonia, unspecifier ry - Other extrapyra Essential tremor Drug-induced tremo Other chorea Drug induced tremo Other chorea Drug induced tremo Other chorea Drug induced tremo Other chorea Drug induced tremo Other tics of organic Drug induced tremo Other specified extra Other specified extra Other specified extra	marily affecting the ce marily affecting centra sewhere ome (G14) t disorders (G20-G26) ease (G20) insonism (G21) ic syndrome parkinsonism secondary parkinsonis nism due to other exter kinsonism mkinsonism nism, unspecified titve diseases of basa lisease clear ophthalmoplegia ration merative diseases of basa clear ophthalmoplegia ration e of basal ganglia, unsp ute dyskinesia dystonia dystonia dystonia stonia al dystonia stonia d dystonia d fistonia d fistonia d d midal and movement a origin ment disorder, unspeci sia movement disorders me tttacks upyramidal and movem	pathy central nervous system al nervous system in c il anarvous system in c il ganglia (G23) [Steele-Richardson-Ol: asal ganglia pecified ified	m in myxedema ther
G11.19 Other early-onset cerebellar ataxia G11.2 Late-onset cerebellar ataxia					G25.9	Extrapyramidal and	novement disorder, ur and movement disor	nspecified	elswhr (G26)
G11.3 Cere	bellar ataxia with	defective DNA repa		equisition Form P		.,	novement disorders ir	n diseases classified e	sewhere
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G11.4 Hereditary spastic paraplegia	Other degenerative disease
G11.8 Other hereditary ataxias	ICD Category - Alzheimer's G30.0 Alzheimer's disea
G11.9 Hereditary ataxia, unspecified ICD Category - Spinal muscular atrophy and related syndromes (G12)	G30.1 Alzheimer's disea
G12.0 Infantile spinal muscular atrophy, type I [Werdnig-Hoffman]	G30.8 Other Alzheimer
G12.1 Other inherited spinal muscular atrophy	G30.9 Alzheimer's disea
G12.20 Motor neuron disease, unspecified G12.21 Amyotrophic lateral sclerosis	ICD Category - Oth degener G31.01 Pick's disease
G12.22 Progressive bulbar palsy	G31.09 Other frontotem
G12.23 Primary lateral sclerosis	G31.1 Senile degenerat
 G12.24 Familial motor neuron disease G12.25 Progressive spinal muscle atrophy 	G31.2 Degeneration of G31.81 Alpers disease
G12.29 Other motor neuron disease	G31.82 Leigh's disease
G12.8 Other spinal muscular atrophies and related syndromes	G31.83 Dementia with Le
G12.9 Spinal muscular atrophy, unspecified ICD Category - Oth degeneratv disord of nervous sys in dis classd elswhr (G32)	G31.84 Mild cognitive im G31.85 Corticobasal deg
G32.0 Subacute combined degeneration of spinal cord in diseases	G31.89 Other specified d
classified elsewhere	G31.9 Degenerative dis
 G32.81 Cerebellar ataxia in diseases classified elsewhere G32.89 Other specified degenerative disorders of nervous system in diseases 	G40.804 Other G40.811 Lennox-Gastaut s
classified elsewhere	G40.812Lennox-Gastaut s
Demyelinating diseases of the central nervous system (G35-G37)	G40.813 Lennox-Gastaut sy
ICD Category - Multiple sclerosis (G35) G35 Multiple sclerosis	G40.814Lennox-Gastaut sy G40.821Epileptic spasms,
ICD Category - Other acute disseminated demyelination (G36)	G40.822 Epileptic spasms,
G36.0 Neuromyelitis optica [Devic]	G40.823 Epileptic spasms, G40.824 Epileptic spasms,
 G36.1 Acute and subacute hemorrhagic leukoencephalitis [Hurst] G36.8 Other specified acute disseminated demyelination 	G40.833 Dravet syndrome,
G36.9 Acute disseminated demyelination, unspecified	G40.834 Dravet syndrome,
ICD Category - Other demyelinating diseases of central nervous system (G37)	G40.89 Other seizures G40.901 Epilepsy, unspecif
G37.0 Diffuse sclerosis of central nervous system	G40.909 Epilepsy, unspecif
G37.1 Central demyelination of corpus callosum G37.2 Central pontine myelinolysis	G40.911 Epilepsy, unspecif
G37.3 Acute transverse myelitis in demyelinating disease of central nervous system	G40.919Epilepsy, unspecif
G37.4 Subacute necrotizing myelitis of central nervous system	ICD Category - Migraine (G4 G43.001 Migraine without
 G37.5 Concentric sclerosis [Balo] of central nervous system G37.8 Other specified demyelinating diseases of central nervous system 	G43.009 Migraine without
G37.9 Demyelinating disease of central nervous system, unspecified	G43.011 Migraine without
Episodic and paroxysmal disorders (G40-G47)	G43.019 Migraine without G43.101 Migraine with aur
ICD Category - Epilepsy and recurrent seizures (G40) G40.001 Localization-related (focal) (partial) idiopathic epilepsy and epileptic	G43.109 Migraine with aur
syndromes with seizures of localized onset, not intractable with status epilepticus	G43.111 Migraine with aur
G40.009 Localization-related (focal) (partial) idiopathic epilepsy and epileptic	G43.119 Migraine with aur G43.401 Hemiplegic migra
syndromes with seizures of localized onset, not intractable, without status epilepticus G40.011 Localization-related (focal) (partial) idiopathic epilepsy and epileptic	G43.409 Hemiplegic migra
syndromes with seizures of localized onset, intractable, with status epilepticus	G43.411 Hemiplegic migra
G40.019 Localization-related (focal) (partial) idiopathic epilepsy and epileptic syndromes with seizures of localized onset, intractable, without status epilepticus	G43.419Hemiplegic migra
G40.101 Localization-related (focal) (partial) symptomatic epilepsy and epileptic	with status migrainosus
syndromes with simple partial seizures, not intractable, with status epilepticus	G43.509 Persistent migrain
G40.109 Localization-related (focal) (partial) symptomatic epilepsy and epileptic syndromes with simple partial seizures, not intractable, without status epilepticus	without status migrainosus
G40.111 Localization-related (focal) (partial) symptomatic epilepsy and epileptic	G43.511 Persistent migrain status migrainosus
syndromes with simple partial seizures, intractable, with status epilepticus $\Box_{\rm res}$ contractable in the second seco	G43.519Persistent migrain
G40.119 Localization-related (focal) (partial) symptomatic epilepsy and epileptic syndromes with simple partial seizures, intractable, without status epilepticus	status migrainosus
G40.201 Localization-related (focal) (partial) symptomatic epilepsy and epileptic	G43.601 Persistent migrain with status migrainosus
syndromes with complex partial seizures, not intractable, with status epilepticus G40.209 Localization-related (focal) (partial) symptomatic epilepsy and epileptic	G43.609 Persistent migrain
syndromes with complex partial seizures, not intractable, without status epilepticus	without status migrainosus
G40.211 Localization-related (focal) (partial) symptomatic epilepsy and epileptic	G43.611 Persistent migrain with status migrainosus
syndromes with complex partial seizures, intractable, with status epilepticus G40.219 Localization-related (focal) (partial) symptomatic epilepsy and epileptic	G43.619Persistent migrain
syndromes with complex partial seizures, intractable, without status epilepticus	without status migrainosus
G40.301 Generalized idiopathic epilepsy and epileptic syndromes, not intractable,	G43.701 Chronic migraine G43.709 Chronic migraine w
with status epilepticus G40.309 Generalized idiopathic epilepsy and epileptic syndromes, not intractable,	G43.711 Chronic migraine
without status epilepticus	G43.719Chronic migraine
G40.311 Generalized idiopathic epilepsy and epileptic syndromes, intractable,	G43.A0 Cyclical vomiting, G43.A1 Cyclical vomiting,
with status epilepticus Generalized idiopathic epilepsy and epileptic syndromes, intractable,	G43.B0 Ophthalmoplegic
without status epilepticus	G43.B1 Ophthalmoplegic
G40.A01 Absence epileptic syndrome, not intractable, with status epilepticus	G43.C0 Periodic headache
 G40.A09 Absence epileptic syndrome, not intractable, without status epilepticus G40.A11 Absence epileptic syndrome, intractable, with status epilepticus 	G43.D0 Abdominal migrai
G40.A19 Absence epileptic syndrome, intractable, with status epilepticus	G43.D1 Abdominal migrai
G40.B01 Juvenile myoclonic epilepsy, not intractable, with status epilepticus	G43.801 Other migraine, no G43.809 Other migraine, no
G40.809 Juvenile myoclonic epilepsy, not intractable, without status epilepticus	G43.809 Other migraine, int
G40.B11 Juvenile myoclonic epilepsy, intractable, with status epilepticus G40.B19 Juvenile myoclonic epilepsy, intractable, without status epilepticus	G43.819 Other migraine, int
G40.401 Other generalized epilepsy and epileptic syndromes, not intractable,	G43.821 Menstrual migrain
with status epilepticus \Box 640.409 . Other generalized epilepsy and epileptic syndromes not intractable	G43.829 Menstrual migrain G43.831 Menstrual migrain
G40.409 Other generalized epilepsy and epileptic syndromes, not intractable, without status epilepticus	G43.839 Menstrual migrain
G40.411 Other generalized epilepsy and epileptic syndromes, intractable, with	G43.901 Migraine, unspecif
status epilepticus G40.419 Other generalized epilepsy and epileptic syndromes, intractable,	G43.909 Migraine, unspecif G43.911 Migraine, unspecif
without status epilepticus	G43.919 Migraine, unspecif

es of the nervous system (G30-G32) disease (G30) ase with early onset ase with late onset 's disease ase, unspecified rative diseases of nervous system, NEC (G31) poral dementia tion of brain, not elsewhere classified nervous system due to alcohol ewy bodies npairment, so stated eneration legenerative diseases of nervous system sease of nervous system, unspecified epilepsy, intractable, without status epilepticus syndrome, not intractable, with status epilepticus yndrome, not intractable, without status epilepticus yndrome, intractable, with status epilepticus yndrome, intractable, without status epilepticus not intractable, with status epilepticus not intractable, without status epilepticus intractable, with status epilepticus intractable, without status epilepticus intractable, with status epilepticus intractable, without status epilepticus fied, not intractable, with status epilepticus fied, not intractable, without status epilepticus fied, intractable, with status epilepticus fied, intractable, without status epilepticus 3) aura, not intractable, with status migrainosus aura, not intractable, without status migrainosus aura, intractable, with status migrainosus aura, intractable, without status migrainosus ra, not intractable, with status migrainosus ra, not intractable, without status migrainosus ra, intractable, with status migrainosus ra, intractable, without status migrainosus aine, not intractable, with status migrainosus aine, not intractable, without status migrainosus aine, intractable, with status migrainosus aine, intractable, without status migrainosus ne aura without cerebral infarction, not intractable, ne aura without cerebral infarction, not intractable, ne aura without cerebral infarction, intractable, with ne aura without cerebral infarction, intractable, without he aura with cerebral infarction, not intractable, e aura with cerebral infarction, not intractable, ne aura with cerebral infarction, intractable, ne aura with cerebral infarction, intractable, without aura, not intractable, with status migrainosus vithout aura, not intractable, without status migrainosus without aura, intractable, with status migrainosus without aura, intractable, without status migrainosus , in migraine, not intractable , in migraine, intractable migraine, not intractable migraine, intractable e syndromes in child or adult, not intractable e syndromes in child or adult, intractable

- ne, not intractable
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- ied, not intractable, without status migrainosus
 - ied, intractable, with status migrainosus
- G43.919 Migraine, unspecified, intractable, without status migrainosus

G40.42 Cyclin-Dependent Kinase-Like 5 Deficiency Disorder G40.501 Epileptic seizures related to external causes, not intractable, with status G40.509 Epileptic seizures related to external causes, not intractable G40.801 Other epilepsy, not intractable, with status epilepticus G40.802 Other epilepsy, not intractable, with status epilepticus G40.803 Other epilepsy, intractable, with status epilepticus	ICD Category - Other headache syndromes (G44) G44.001 Cluster headache syndrome, unspecified, intractable
 G40.501 Epileptic seizures related to external causes, not intractable, with status G40.509 Epileptic seizures related to external causes, not intractable G40.801 Other epilepsy, not intractable, with status epilepticus G40.802 Other epilepsy, not intractable, without status epilepticus 	
 G40.509 Epileptic seizures related to external causes, not intractable G40.801 Other epilepsy, not intractable, with status epilepticus G40.802 Other epilepsy, not intractable, without status epilepticus 	
 G40.801 Other epilepsy, not intractable, with status epilepticus G40.802 Other epilepsy, not intractable, without status epilepticus 	
G40.802 Other epilepsy, not intractable, without status epilepticus	G44.009 Cluster headache syndrome, unspecified, not intractable
	G44.011 Episodic cluster headache, intractable
\Box (40.902) Other enderse intractable with status endertises	G44.019 Episodic cluster headache, not intractable
\Box G40.005 Uther epilepsy, intractable, with status epilepticus	G44.021 Chronic cluster headache, intractable
G44.039 Episodic paroxysmal hemicrania, not intractable	G44.029 Chronic cluster headache, not intractable
G44.041 Chronic paroxysmal hemicrania, intractable	
	G44.031 Episodic paroxysmal hemicrania, intractable
G44.049 Chronic paroxysmal hemicrania, not intractable	G47.35 Congenital central alveolar hypoventilation syndrome
G44.051 Short lasting unilateral neuralgiform headache with conjunctival	G47.36 Sleep related hypoventilation in conditions classified elsewhere
injection and tearing (SUNCT), intractable	G47.37 Central sleep apnea in conditions classified elsewhere
G44.059 lasting unilateral neuralgiform headache with conjunctival injection	G47.39 Other sleep apnea
5 5 , , ,	
and tearing (SUNCT), not intractable	G47.411 Narcolepsy with cataplexy
G44.091 Other trigeminal autonomic cephalgias (TAC), intractable	G47.419 Narcolepsy without cataplexy
G44.099 Other trigeminal autonomic cephalgias (TAC), not intractable	G47.421 Narcolepsy in conditions classified elsewhere with cataplexy
G44.1 Vascular headache, not elsewhere classified	G47.429 Narcolepsy in conditions classified elsewhere without cataplexy
G44.201 Tension-type headache, unspecified, intractable	G47.50 Parasomnia, unspecified
G44.209Tension-type headache, unspecified, not intractable	G47.51 Confusional arousals
G44.211 Episodic tension-type headache, intractable	G47.52 REM sleep behavior disorder
G44.219 Episodic tension-type headache, not intractable	G47.53 Recurrent isolated sleep paralysis
G44.221 Chronic tension-type headache, intractable	G47.54 Parasomnia in conditions classified elsewhere
G44.229 Chronic tension-type headache, not intractable	G47.59 Other parasomnia
G44.301 Post-traumatic headache, unspecified, intractable	G47.61 Periodic limb movement disorder
G44.309 Post-traumatic headache, unspecified, not intractable	G47.62 Sleep related leg cramps
G44.311 Acute post-traumatic headache, intractable	G47.63 Sleep related bruxism
G44.319 Acute post-traumatic headache, not intractable	G47.69 Other sleep related movement disorders
G44.321 Chronic post-traumatic headache, intractable	G47.8 Other sleep related movement disorders
G44.329 Chronic post-traumatic headache, not intractable	G47.9 Sleep disorder, unspecified
G44.40 Drug-induced headache, not elsewhere classified, not intractable	Nerve, nerve root and plexus disorders (G50-G59)
G44.41 Drug-induced headache, not elsewhere classified, intractable	ICD Category - Disorders of trigeminal nerve (G50)
G44.51 Hemicrania continua	G50.0 Trigeminal neuralgia
G44.52 New daily persistent headache (NDPH)	
	G50.1 Atypical facial pain
G44.53 Primary thunderclap headache	G50.8 Other disorders of trigeminal nerve
G44.59 Other complicated headache syndrome	G50.9 Disorder of trigeminal nerve, unspecified
G44.81 Hypnic headache	ICD Category - Facial nerve disorders (G51)
G44.82 Headache associated with sexual activity	
G44.83 Primary cough headache	G51.0 Bell's palsy
	G51.1 Geniculate ganglionitis
G44.84 Primary exertional headache	G51.2 Melkersson's syndrome
G44.85 Primary stabbing headache	G51.31 Clonic hemifacial spasm, right
G44.86 Cervicogenic headache	G51.32 Clonic hemifacial spasm, left
G44.89 Other headache syndrome	
·	G51.33 Clonic hemifacial spasm, bilateral
ICD Category - Transient cerebral ischemic attacks and related syndromes (G45)	G51.39 Clonic hemifacial spasm, unspecified
G45.0 Vertebro-basilar artery syndrome	G51.4 Facial myokymia
G45.1 Carotid artery syndrome (hemispheric)	G51.8 Other disorders of facial nerve
G45.2 Multiple and bilateral precerebral artery syndromes	G51.9 Disorder of facial nerve, unspecified
G45.3 Amaurosis fugax	
	ICD Category - Disorders of other cranial nerves (G52)
G45.4 Transient global amnesia	G52.0 Disorders of olfactory nerve
G45.8 Other transient cerebral ischemic attacks and related syndromes	G52.1 Disorders of glossopharyngeal nerve
G45.9 Transient cerebral ischemic attack, unspecified	G52.2 Disorders of vagus nerve
ICD Category - Vascular syndromes of brain in cerebrovascular diseases (G46)	G52.3 Disorders of hypoglossal nerve
G46.0 Middle cerebral artery syndrome	
	G52.7 Disorders of multiple cranial nerves
	G52.8 Disorders of other specified cranial nerves
G46.1 Anterior cerebral artery syndrome	
G46.2 Posterior cerebral artery syndrome	G52.9 Cranial nerve disorder, unspecified
G46.2 Posterior cerebral artery syndrome G46.3 Brain stem stroke syndrome	ICD Category - Cranial nerve disorders in diseases slassified elsewhere (G53)
 G46.2 Posterior cerebral artery syndrome G46.3 Brain stem stroke syndrome G46.4 Cerebellar stroke syndrome 	
 G46.2 Posterior cerebral artery syndrome G46.3 Brain stem stroke syndrome G46.4 Cerebellar stroke syndrome G46.5 Pure motor lacunar syndrome 	ICD Category - Cranial nerve disorders in diseases slassified elsewhere (G53)
 G46.2 Posterior cerebral artery syndrome G46.3 Brain stem stroke syndrome G46.4 Cerebellar stroke syndrome G46.5 Pure motor lacunar syndrome G46.6 Pure sensory lacunar syndrome 	ICD Category - Cranial nerve disorders in diseases slassified elsewhere (G53) G53 Cranial nerve disorders in diseases classified elsewhere ICD Category - Nerve root and plexus disorders (G54)
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 G46.2 Posterior cerebral artery syndrome G46.3 Brain stem stroke syndrome G46.4 Cerebellar stroke syndrome G46.5 Pure motor lacunar syndrome G46.6 Pure sensory lacunar syndrome G46.7 Other lacunar syndromes 	ICD Category - Cranial nerve disorders in diseases slassified elsewhere (G53) G53 Cranial nerve disorders in diseases classified elsewhere ICD Category - Nerve root and plexus disorders (G54) G54.0 Brachial plexus disorders G54.1 Lumbosacral plexus disorders
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G56.82 Other specified mononeuropathies of left upper limb	ICD Category - Other disorders of peripheral nervous system (G64)
G56.83 Other specified mononeuropathies of bilateral upper limbs	G64 Other disorders of peripheral nervous system
G56.90 Unspecified mononeuropathy of unspecified upper limb	ICD Category - Sequelae of inflammatory and toxic polyneuropathies (G65)
G56.91 Unspecified mononeuropathy of right upper limb G56.92 Unspecified mononeuropathy of left upper limb	G65.0 Sequelae of Guillain-Barré syndrome
G56.93 Unspecified mononeuropathy of bilateral upper limbs	G65.2 Sequelae of toxic polyneuropathy G5.2 Sequelae of toxic polyneuropathy
ICD Category - Mononeuropathies of lower limb (G57)	Diseases of myoneural junction and muscle (G70-G73)
G57.00 Lesion of sciatic nerve, unspecified lower limb	ICD Category G70
G57.01 Lesion of sciatic nerve, right lower limb	G70.00 Myasthenia gravis without (acute) exacerbation
G57.02 Lesion of sciatic nerve, left lower limb	G70.01 Myasthenia gravis with (acute) exacerbation
G57.03 Lesion of sciatic nerve, bilateral lower limbs	G70.1 Toxic myoneural disorders
G57.10 Meralgia paresthetica, unspecified lower limb G57.11 Meralgia paresthetica, right lower limb	 G70.2 Congenital and developmental myasthenia G70.80 Lambert-Eaton syndrome, unspecified
G57.12 Meralgia paresthetica, left lower limb	G70.81 Lambert-Eaton syndrome in disease classified elsewhere
G57.13 Meralgia paresthetica, bilateral lower limbs	G70.89 Other specified myoneural disorders
G57.20 Lesion of femoral nerve, unspecified lower limb	G70.9 Myoneural disorder, unspecified
G57.21 Lesion of femoral nerve, right lower limb	ICD Category - Myasthenia gravis and other myoneural disorders (G70)
G57.22 Lesion of femoral nerve, left lower limb G57.23 Lesion of femoral nerve, bilateral lower limbs	 G71.00 Muscular dystrophy, unspecified G71.01 Duchenne or Becker muscular dystrophy
G57.30 Lesion of lateral popliteal nerve, unspecified lower limb	G71.02 Facioscapulohumeral muscular dystrophy
G57.31 Lesion of lateral popliteal nerve, right lower limb	G71.09 Other specified muscular dystrophies
G57.32 Lesion of lateral popliteal nerve, left lower limb	G71.11 Myotonic muscular dystrophy
G57.33 Lesion of lateral popliteal nerve, bilateral lower limbs	G71.12 Myotonia congenita
G57.40 Lesion of medial popliteal nerve, unspecified lower limb G57.41 Lesion of medial popliteal nerve, right lower limb	G71.13 Myotonic chondrodystrophy G71.14 Drug induced myotonia
G57.42 Lesion of medial popliteal nerve, left lower limb	G71.19 Other specified myotonic disorders
G57.43 Lesion of medial popliteal nerve, bilateral lower limbs	G71.20 Congenital myopathy, unspecified
G57.50 Lesion of medial popliteal nerve, bilateral lower limbs	G71.21 Nemaline myopathy
G57.51 Tarsal tunnel syndrome, right lower limb	G71.220 X-linked myotubular myopathy
G57.52 Tarsal tunnel syndrome, left lower limb	G71.228 Other centronuclear myopathy G71.29 Other congenital myopathy
G57.53 Tarsal tunnel syndrome, bilateral lower limbs G57.60 Lesion of plantar nerve, unspecified lower limb	G71.3 Mitochondrial myopathy, not elsewhere classified
G57.61 Lesion of plantar nerve, right lower limb	G71.8 Other primary disorders of muscles
G57.62 Lesion of plantar nerve, left lower limb	G71.9 Primary disorder of muscle, unspecified
G57.63 Lesion of plantar nerve, bilateral lower limbs	ICD Category - Primary disorders of muscles (G71)
G57.70 Causalgia of unspecified lower limb	G72.0 Drug-induced myopathy
G57.71 Causalgia of right lower limb G57.72 Causalgia of left lower limb	G72.1 Alcoholic myopathy G72.2 Myopathy due to other toxic agents
G57.73 Causalgia of bilateral lower limbs	G72.3 Periodic paralysis
G57.80 Other specified mononeuropathies of unspecified lower limb	G72.41 Inclusion body myositis [IBM]
G57.81 Other specified mononeuropathies of right lower limb	G72.49 Other inflammatory and immune myopathies, not elsewhere classified
G57.82 Other specified mononeuropathies of left lower limb	G72.81 Critical illness myopathy
G57.83 Other specified mononeuropathies of bilateral lower limbs	 G72.89 Other specified myopathies G72.9 Myopathy, unspecified
G57.90 Unspecified mononeuropathy of unspecified lower limb	
 G57.90 Unspecified mononeuropathy of unspecified lower limb G57.91 Unspecified mononeuropathy of right lower limb G57.92 Unspecified mononeuropathy of left lower limb 	ICD Category - Other and unspecified myopathies (G72) G72.0 G72.0 Drug-induced myopathy
G57.91 Unspecified mononeuropathy of right lower limb	ICD Category - Other and unspecified myopathies (G72) G72.0 Drug-induced myopathy G72.1 Alcoholic myopathy
G57.91 Unspecified mononeuropathy of right lower limb G57.92 Unspecified mononeuropathy of left lower limb G57.93 Unspecified mononeuropathy of bilateral lower limbs ICD Category - Other mononeuropathies (G58)	ICD Category - Other and unspecified myopathies (G72) G72.0 Drug-induced myopathy G72.1 Alcoholic myopathy G72.2 Myopathy due to other toxic agents
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ICD Category G82	
	ICD Category G93
G82.20 Paraplegia, unspecified	G93.0 Cerebral cysts
G82.21 Paraplegia, complete	G93.1 Anoxic brain damage, not elsewhere classified
G82.22 Paraplegia, incomplete	G93.2 Benign intracranial hypertension
G82.50 Quadriplegia, unspecified	G93.3 Postviral fatigue syndrome
G82.51 C1-C4 complete	G93.40 Encephalopathy, unspecified
G82.52 C1-C4 Incomplete	G93.41 Metabolic encephalopathy
G82.53 C5-C7 complete	
	G93.49 Other encephalopathy
G82.54 C5-C7 Incomplete	G93.5 Compression of brain
ICD Category G83	G93.6 Cerebral edema
G83.0 Diplegia of upper limbs	
	G93.7 Reye's syndrome
G83.10 Monoplegia of lower limb affecting unspecified side	G93.81 Temporal sclerosis
G83.11 Monoplegia of lower limb affecting right dominant side	G93.82 Brain death
G83.12 Monoplegia of lower limb affecting left dominant side	
_	G93.89 Other specified disorders of brain
G83.13 Monoplegia of lower limb affecting right nondominant side	G93.9 Disorder of brain, unspecified
G83.14 Monoplegia of lower limb affecting left nondominant side	ICD Category G94
G83.20 Monoplegia of upper limb affecting unspecified side	
G83.21 Monoplegia of upper limb affecting right dominant side	G94 Other disorders of brain in diseases classified elsewhere
	ICD Category G95
G83.22 Monoplegia of upper limb affecting left dominant side	
G83.23 Monoplegia of upper limb affecting right nondominant side	
G83.24 Monoplegia of upper limb affecting left nondominant side	G95.11 Acute infarction of spinal cord (embolic) (nonembolic)
	G95.19 Other vascular myelopathies
G83.30 Monoplegia, unspecified affecting unspecified side	G95.20 Unspecified cord compression
G83.31 Monoplegia, unspecified affecting right dominant side	
G83.32 Monoplegia, unspecified affecting left dominant side	G95.29 Other cord compression
G83.33 Monoplegia, unspecified affecting right nondominant side	G95.81 Conus medullaris syndrome
	G95.89 Other specified diseases of spinal cord
G83.34 Monoplegia, unspecified affecting left nondominant side	
G83.4 Cauda equina syndrome	G95.9 Disease of spinal cord, unspecified
G83.5 Locked-in state	ICD Category G96
_	G96.00 Cerebrospinal fluid leak, unspecified
G83.81 Brown-Séquard syndrome	
G83.82 Anterior cord syndrome	G96.01 Cranial cerebrospinal fluid leak, spontaneous
G83.83 Posterior cord syndrome	G96.02 Spinal cerebrospinal fluid leak, spontaneous
G83.84 Todd's paralysis (postepileptic)	G96.08 Other cranial cerebrospinal fluid leak
G83.89 Other specified paralytic syndromes	G96.09 Other spinal cerebrospinal fluid leak
G83.9 Paralytic syndrome, unspecified	G96.11 Dural tear
G89-G99	G96.12 Meningeal adhesions (cerebral) (spinal)
ICD Category G89	G96.191 Perineural cyst
G89.0 Central pain syndrome	G96.198 Other disorders of meninges, not elsewhere classified
G89.11 Acute pain due to trauma	G96.810 Intracranial hypotension, unspecified
G89.12 Acute post-thoracotomy pain	G96.811 Intracranial hypotension, spontaneous
G89.18 Other acute postprocedural pain	G96.819 Other intracranial hypotension
G89.21 Chronic pain due to trauma	G96.89 Other specified disorders of central nervous system
G89.22 Chronic post-thoracotomy pain	
	G96.9 Disorder of central nervous system, unspecified
G89.28 Other chronic postprocedural pain	ICD Category G97
G89.29 Other chronic pain	G97.0 Cerebrospinal fluid leak from spinal puncture
G89.3 Neoplasm related pain (acute) (chronic)	
	G97.1 Other reaction to spinal and lumbar puncture
G89.4 Chronic pain syndrome	G97.2 Intracranial hypotension following ventricular shunting
ICD Category G90	G97 31 Intraoperative bemorrhage and bematoma of a pervous system organ
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G90.01 Carotid sinus syncope	or structure complicating a nervous system procedure
G90.01 Carotid sinus syncope G90.09 Other idiopathic peripheral autonomic neuropathy	
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Additional ICD Codes:.....

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Physician Consent for Genetic Testing

Advanced Genomics continually updates its panels based on the most recent evidence. If an order is placed using an outdated test requisition form, educated entering and the second s

upgrade ordered tests to their current versions. Test IDs containing add-on codes will include the original panel as well as the add-on.

By signing this form, the medical professional acknowledges that the individual/family member authorized to make decisions for the individual (collectively, the "Patient") has been supplied information regarding and consented to undergo genetic testing, substantially as set forth in **Advanced Genomics** Informed Consent for Genetic T esting **Advanced Genomics**. The medical professional will retain evidence that the patient consented to genetic testing. The Patient has been informed that **tAdvanced Genomics** may notify them of clinical updates related to genetic Test results (in consultation with the ordering medical professional as indicated) and has been informed that DE identified(also referred to as pseudonymised) patient data may be used and shared with third parties in connection with the Program, for research and commercial purposes. For orders originating outside the United States, the Patient has been informed that their personal information and specimen will be transferred to and processed in the United States. The medical professional warrants that (i) he/she will not seek reimbursement for No charge test from any third, including but not limited to government healthcare programs: (ii) participation in the program will not influence the his/her medical this decisions; (iii) he/ She is not obligated to participate in or to encourage patients to participate in any clinical trial or other research program conducted by a sponsor; and (v) he/she will participate in the Program in accordance with applicable laws. The medical professional consents to the sharing of organization and clinician contact information with third parties, including commercial organizations, who may contact the medical professional directly in connection with the Program. A list of third party partners will be provided upon request. I attest that I am authorized under applicable law to order this test.

Medical professional signature (required)

L.

Patient Informed Consent for Genetic Testing

(Patient's or Patient's Legal Guardian's Full Name)

(Test Type/Name)

Date (MM/DD/YYYY)

enomics	to conduct	(genetic
			<i>,</i>

(Patient's or Patient's Legal Guardian's Full Name) (Test Type/Name) testing, as ordered by my physician/authorized healthcare provider or as ordered by my child's/dependent's physician/authorized healthcare provider, and also authorize the collection of a sample for the purpose of this testing.

By signing below, I acknowledge and/or consent to the following statements:

1. My physician or his/her designee (such as a genetic counselor) has fully explained to me the following:

_, authorize Advanced G

(A) the purpose, description, and nature of this test and its potential uses.

(B) the reliability of positive or negative results and the level of certainty that a positive result for a disease or condition(s) holds, serving as a predictor of such disease or condition(s).

(C) the effectiveness and limitations of the genetic test and the meaning of the possible genetic test results.

(D) the implications of conducting this genetic testing, including the medical risks and benefits.

(E) the description of the disease or condition(s) being tested for.

(F) the availability and importance of meeting with a genetic counselor or medical geneticist, which I have been

provided information on identifying an individual from whom I might obtain counseling, that I may seek prior to signing this consent.

(G) that a positive result is an indication I may be predisposed to or could have the specific disease or condition(s) tested for. That if I wish to discuss my results and their meaning, I should consult with my physician and/or pursue genetic counseling. That if I wish to consider further independent testing after reviewing my r esults, I should consult with my physician.

2. I understand that I will receive my test results from my physician/authorized healthcare provider unless I have explicitly directed otherwise.

3. I understand that I have a right to confidential treatment of my sample and results, and that my results will only be disclosed as I have authorized in this consent.

(A) I authorize my test results to be disclosed to the following person(s), should they be requested:

* Please list all authorized person(s) below with first name, last name, and relation to patient.

4. I understand that only my physician's offic e and **Advanced Genomics** will have access to my sample and that my sample will only be used for the purposes which I have authorized in this consent.

Patient's Statement

I, the undersigned, have been informed about the test's purpose, procedures, possible benefits, possible risks, and I have been provided a copy of this consent for my records. I have been provided the opportunity to ask questions before signing this consent, and I have been assured that I can ask further questions at any time. I voluntarily agree to this genetic testing.

Patient signature (required)