

Exome with CNV Evaluation Clinical History Form

Client account number:	Client name:		
Primary patient name:			
Primary patient DOB:		Patient phone number:	
If ordering trio or duo testing, please provide the	e names and DOB of add	itional family members:	
Name:			DOB:
Name:		'	
Proband information			
Reason(s) for referral for exome sequencing, ie,	what was the initial pre	senting symptom?	
		Age of onset of init	ial presenting symptom:
Ethnicity (check all that apply):			
☐ African or descended from the African continu ☐ Central or South American ☐ French Canad ☐ Sephardic Jewish ☐ Western/Northern Eur	lian □ Hispanic or Lat	ino □ Middle Eastern □ Pacific Islar	nder or Native Hawaiian
Is there a family history of a similar or related di	isorder? □Yes □Nc		
Consanguinity (related by blood, eg, parents rela	ated by blood)? □ Yes	□ No □ Unsure If yes, please specif	:y:
History of a bone marrow transplant: ☐ Yes* *If yes, please contact 1.866.GENE.INFO to spea		Specialist before sending any samples.	
Previous genetic testing			
Chromosome analysis ☐ No results/not perfo	rmed 🗆 Yes/results: .		
Chromosomal microarray analysis □ No resul	ts/not performed 🛚 Ye	s/results:	
Other molecular studies, including prenatal test	ting:		
Reporting of Secondary Findings (Physician Si	ignature Required)		
Secondary Findings Report: I have provided general explained the potential benefits and limitations patient's consent regarding the reporting of secondary findings recommedically actionable secondary findings recommedically	netic counseling to the p s of receiving secondary condary findings as indic	findings, have answered the patient's qua ated below. Check the appropriate option	estions, and have obtained the n below to order reporting of
☐ YES , Please report pathogenic variants in ger	nes determined to be me	dically actionable by the ACMG policy sta	atement
□ NO, Please DO NOT report pathogenic variant	s in genes determined to	be medically actionable by the ACMG pe	olicy statement
SIGNATURE REQUIRED			
Medical Professional's Signature:			Date
Patient Authorization to Use De-Identified Sp	ecimen (Patient Signati	ure Required)	
Test results are confidential and will only be repapplicable state and federal law.			presentative or consistent with
To promote medical understanding and develop in a de-identified way (without identifying the te or other personal identifying information will no	est subject) for research,	educational studies, commercial purpos	ses and/or publication. Your name
You are not required to consent to any of these u affect processing or testing of the specimen, the indicate your choice regarding the use of the de	test results or the service	es provided by Quest Diagnostics in conn	ection with this testing. Please
☐ YES, I consent to the use of my de-identified s	specimen as described	above.	
\square $\mathbf{N0}, I$ do not consent to the use my de-identifi	ed specimen as describ	ed above.	
Signature of Patient, Parent or Legally Authorized Repr	resentative		Date
Print Name			Relationship

Clinical de	tails			
			ale Other (please specify):	
Common di	agnoses (please provide m us genitalia Autism sp	ore information about these common c ectrum disorder Congenital heart d	%tile Height: diagnoses using the check boxes furthe efect □ Developmental delay □ Dysi iital anomalies □ Seizures □ Structu	r below): morphic features
Exon	ne with CNV	Evaluation Clinic	cal History Form	
		Endocrine Adrenal gland abnormality Adrenal insufficiency Cushing syndrome Diabetes insipidus Diabetes mellitus Growth hormone deficiency Hirsutism Immunologic abnormality Specify: Obesity Pancreatic insufficiency Parathyroid dysfunction	Hair & skin (continued) Hair Alopecia Brittle Coarse Hypopigmentation Hemangioma Hyperextensible skin Hyperpigmented macule Hypopigmented macule Hypopigmented macule Hypertrichosis Infections	☐ Generalized myoclonic ☐ Generalized tonic ☐ Generalized tonic-clonic ☐ Infantile spasms ☐ Spasticity Pre/perinatal history ☐ Conceived via artificial reproductive technology ☐ Congenital diaphragmatic hernia ☐ Cystic hygroma Pre/perinatal history (continued)
☐ Patent fo☐ Teratolog☐ Thromboo☐ Thromboo☐ Tortuosity☐ Truncus a☐ Ventricula	ot dilation ia ia ital defect aortic valve ropathy on of aorta ormality ve prolapse iotus arteriosus ramen ovale y of Fallot cytopenia sis	□ Thyroid dysfunction Eye defects & vision □ Amblyopia □ Aniridia □ Anophthalmia □ Blue sclerae □ Cataracts □ Congenital □ Postnatal □ Cherry red spot □ Coloboma □ Corneal abnormality □ Ectopia lentis □ Microphthalmia □ Nystagmus □ Ptosis □ Retinitis pigmentosa □ Strabismus	□ Lipoma □ Nail abnormality □ Neurofibroma □ Rash Metabolic □ Acidosis □ Lactic □ Metabolic □ CSF lactate level (abnormal) □ Dicarboxylic aciduria □ Hyperammonemia □ Hyperglycemia □ Hypoglycemia □ Hypoglycemia □ Hypopammonemia □ Hypoammonemia □ Hypoammonemia □ Ketosis □ Organic aciduria □ Phosphokinase (abnormal)	□ Encephalocele □ Increased nuchal translucency □ Intrauterine growth restriction □ Oligohydramnios □ Polyhydramnios □ Omphalocele □ Prematurity □ Teratogen exposure Specify: □ Respiratory □ Apnea □ Asthma □ Bronchiectasis □ Hyperventilation □ Hypoventilation □ Pneumothorax □ Recurrent infections
Craniofacia Bifid uvul Cleft lip Cleft pala Craniosyr Epicanth Hyperteld Hyperteld Macrocep Microcep Microgna Nose abn Palpebra Philtrum Teeth abr	ate nostosis al folds orism orism ohaly haly thia ormality l fissures abnormality normality	□ Visual impairment □ Blind □ Cortical □ Myopia Gastrointestinal □ Anal malformation □ Constipation (chronic) □ Crohn's disease □ Diarrhea (chronic) □ Esophageal atresia □ Gastroparesis □ Hepatic failure □ Hepatomegaly □ Hirschsprung disease □ Inflammatory bowel disease	□ Plasma carnitine (abnormal) □ Serum creatine (abnormal) □ Serum pyruvate (abnormal) Musculoskeletal □ Arthrogryposis □ Camptodactyly □ Contractures □ Fractures □ Hemihypertrophy □ Hyperlordosis □ Hypermobility □ Hypermobility □ Hypertonia □ Hypotonia □ Kyphosis □ Muscle atrophy □ Muscular dystrophy	□ Respiratory failure □ Respiratory insufficiency Structural brain abnormalities □ Aplasia/hypoplasia of the cerebellar vermis □ Aplasia/hypoplasia of the cerebellum □ Basal ganglia abnormality □ Brain atrophy □ Brainstem abnormality □ Cerebral dysmyelination □ Cerebral hypomyelination □ Cerebral white matter abnormality □ Corpus callosum abnormality □ Corpus callosum abnormality □ Cortical dysplasia □ Encephalocele □ Holoprosencephaly
□ Developm □ Developm □ Intellectu □ Mild □ Modera □ Severe □ Profou	D Dectrum disorder nental delay nental regression nal disability ate nd	□ Intestinal pseudo-obstruction □ Pancreatitis □ Pyloric stenosis □ Splenomegaly □ Vomiting (episodic/cyclic) □ Tracheoesophageal fistula Genitourinary □ Ambiguous genitalia □ Cryptorchidism □ Hypogonadism □ Hypospadias □ Kidney abnormality □ Agenesis	Myopathy Myotonia Oligodactyly Overgrowth Polydactyly Rib defects Scoliosis Short stature Skeletal dysplasia Spina bifida Syndactyly Talipes equinovarus Tall stature Vertebral anomalies	Hydrocephalus Leukoencephalopathy Leukodystrophy Lissencephaly Neuronal migration abnormality Pachgyria Polymicrogyria Ventriculomegaly Other Allergies (severe) Fever (episodic) Failure to thrive
Ear & heari Deafness Acquir Conger Bilater Unilate Condur Senson Low-set & Pinna abr	ed nital al eral ctive rineural ears normality Jlar pit	□ Horseshoe □ Partially duplicated □ Polycystic □ Ovarian streak □ Polycystic ovarian syndrome □ Testicular abnormality □ Ureter abnormality □ Urethra abnormality □ Hair & skin □ Albinism □ Blistering □ Cafe-au-lait spots	Neurologic Areflexia Ataxia Chorea Dystonia Epileptic encephalopathy Neuropathy Seizures Absence Atonic Febrile Generalized clonic	Heterotaxy Lethargy Organomegaly Pain (chronic) Attach any imaging or laboratory results. Questions? To speak with a Genomic Science Specialist, please call: 1.866.GENE.INFO Please fax or email the form to 1.949.668.7818 or Preauthorization_ neurology@QuestDiagnostics.com