Laboratory of Personalized Genomic Medicine **Department of Pathology and Cell Biology** 630 W. 168th Street, P&S 17th Floor, Room 401 New York, New York 10032 Tel: 212-305-9706 Fax: 212-342-0420





College of Physicians and Surgeons

Precision Genomics Laboratory

701 West 168th Street, HHSC 1401A New York, New York 10032 Tel: 212-305-6094 Fax: 212-305-6687

Molecular Genetic Pathology Testing Requisition Form Must be filled out completely. Informed consent MUST be signed by the Patient, Parent/Legal Guardian or Legal Next of Kin.

PATIENT INFORMATION:			ORDERING PHYS	SICIAN INFORMATION:		
LAST NAME: FIRST	NAME:	M.I.:	LAST NAME:	FIRST NA	AME:	M.I.:
DATE OF BIRTH: MRN:	Gender:		INSTITUTION:		NPI #:	
Address:	☐ MALE	□ Female	Address:			
hobitos.						
City, State & ZIP:			CITY, STATE & ZIP:			
Home Phone: Work	PHONE:		TELEPHONE NUMBER:		FAX NUMBER:	
INSURANCE INFORMATION:			EMAIL ADDRESS:			
NAME OF POLICY HOLDER:	DATE OF BIRTH:		SIGNATURE:		DATE:	
Relationship to Patient: Self Parent	SPOUSE CHI	ILD	GENETIC COUNSELOR NA	AME:	EMAIL ADDRESS:	
NAME & ADDRESS OF INSURANCE COMPANY:						
			INSTITUTIONAL BILLI	NG: Do you have a Pathol	ogy Billing Accou	int?
POLICY NUMBER:	GROUP NUMBER:		□ Yes P.0.#			
	NAME OF POLICY HOLDER:			logy-billing@columbia.edu		
SECONDARY INSURANCE CARRIER:	NAME OF POLICY HOLDER:			CARE PRACTITIONER:		
POLICY NUMBER:	GROUP NUMBER:		performing genetic	predisposition testing an	nd maintained in	the patient's medical
				the appropriate disease/s he patient/legal guardian		
MEDICARE PATIENTS ONLY: Check here to confi	irm that an Advance	Beneficiary		ure. If the patient consen		
Notice (ABN) was signed by the Patient:				han 60 days, please inclu have obtained a signed		
		athology		accordance with New		
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Molecular Genetic Pathology Patient Medical Questionnaire

To support the laboratory analysis, please provide a detailed clinical picture of the patient's clinical syndrome, the clinical differential diagnosis, and a pedigree. All of these will help in the interpretation of the pathogenicity of the variants that will be found.

PATIENT INFORMATION								
LAST NAME:	FIRST NAME:		DATE OF BIRTH:		TODAY'S DATE:			
Gender: Male Female	ETHNIC BACKGROUND:	🗆 AFRICAN AMERICAN	🗆 Ashkenazi	Jewish 🛛 Asia	N 🗆 CAUG	CASIAN	□ HISPANIC	
		□ NATIVE AMERICAN	□ OTHER JEW	ISH 🗆 ОТН	ER			
REASON FOR TESTING / CLINICAL INFORMA	ATION							
CLINICAL DIAGNOSIS:						AGE OF ONS	ET:	
POSITIVE FAMILY HISTORY: YES NO	\square N/A IF yes, explain:			Deceased: 🛛 Ye	s 🗆 No	AUTOPSY	: □YES □	No
OTHER SIGNIFICANT OBSERVATIONS:								

GENERAL SYMPTOMS / SIGNS	CUTANEOUS (CONTINUED):		NEOPLASIA (CONTINUED):
BEHAVIORAL/ PSYCHIATRIC	CUTANEOUS (CONTINUED): Papules	HEMATOLOGY	Head and Neck
Attention Deficit Hyperactivity Disorder	Papules Psoriasis	Anemia	
□ Anxiety	□ Psoriasis □ Striae	Clotting Disorder	□ Hematopoietic/ Lymphatic □ Intrathoracic
□ Autism	□ Striae □ Urticaria	Hemophilia	Intrathoracic Neurofibroma
Behavioral/Psychiatric Abnormality		Neutropenia	
□ Dementia	LI ENDOCRINE/ EXOCRINE:	□ Thrombocytopenia	Optic Gliomas
Pervasive Developmental Delay	□ Adrenal	IMMUNE SYSTEM	□ Skin, Soft Tissue, Bone and Joints
	□ Bone	Autoimmune Disorders	LI NEUROLOGICAL
□ Aortic Atresia	Diabetes Mellitus	□ Immunodeficiency	Abnormal Gyri (Lissencephaly)
□ ASD	□ Hypothalamus		Agenesis of the Corpus Callosum
AV Canal Defect	Pancreas	METABOLIC/ LABORATORY ABNORMALITIES	□ Asthenia
Arrial Septal Defect	Parathyroid	Biliary Tract	Cerebellar Hypoplasia
Coarctation of the Aorta	Pituitary	Blood Gases	Cerebellar Signs
□ Congestive Heart Failure	Reproductive	Electrolytes	Cerebral Blindness
Dextrocardia and Situs Inversus	□ Thyroid	□ Gastrointestinal	Cerebral Palsy
Dextrocardia and Situs Inversus Double Outlet Right Ventricle	GASTROINTESTINAL	□ Glycoproteins	Dandy Walker (posterior fossa abnormality)
Ebstein's Anomaly	Absent Stomach	□ Kidney	□ Encephalopathy
Echogenic Intracardiac Focus	Absent Stomach Echogenic Focus	□ Lipids	□ Headache
Hypoplastic Left Heart	□ Gastrointestinal Pseudo Obstruction	□ Liver	□ Holoproscencephaly
Pulmonary Valve Atresia	Gastroschisis	Mitochondrial	□ Hypertonia
		□ Pancreas	□ Hypotonia
Tetralogy of Fallot Tetralogy of Fallot	Meconium Ileus/ Anal Atresia Ourschale sele	MUSCULOSKELETAL	□ Macrocephaly
□ Transposition of the Great Vessels	Omphalocele		Microcephaly – List HC if known:
Truncus Arteriosus	□ Tracheoesophageal Fistula □ Pyloric Stenosis	□ Clenched Hands	□ Neuropathy
Ventricular Septal Defect		□ Club Foot (bilateral)	□ Neural Tube Defect
COGNITIVE/DEVELOPMENTAL	Genitourinary	Contractures (arthrogryposis)	□ Seizures
□ Learning Disability	Ambiguous Genitalia	□ Cramps After Exercise	□ Stroke
Developmental Delay	□ Cryptorchidism	Diaphragmatic Hernia	Structural Brain Anomaly
Gross Motor Delay	□ Hydronephrosis	Exercise Intolerance	Ventriculomegaly/Hydrocephaly
□ Fine Motor Delay	Hypoparathyroidism	□ Ioint Laxity	PRENATAL/PERINATAL HISTORY
□ Speech Delay	□ Hypospadias	\Box Limb Anomaly	2 Vessel Cord
□ Intellectual Disability/MR	□ Hypothyroidism	□ Linio Anomary □ Mesomelia	Decreased Fetal Movement
5,	□ Kidney Malformation	□ Micromelia	
CRANIOFACIAL	Megacystis (incl. posterior valves)	□ Myoclonus	Echogenic Focus
Cleft Lip +/- Cleft Palate	□ Myoglobinuria		□ Floppy Baby
Dysmorphic Facial Features	Polycystic Kidneys	 Ophthalmoplegia Scoliosis 	□ Increased Nuchal Translucency
Ear Malformation	Renal Agenesis	\square Polydactyly	
Hyper/Hypotelorism	Urethra/Ureter Obstruction	Proximal Limb Weakness	□ Non-Immune Hydrops Fetalis
□ Macrocephaly	GROWTH	\Box Skeletal Dysplasia	Oligohydraminos
□ Microcephaly	□ Failure to Thrive	Syndactyly	Perinatal Insult Delabadamentia
CUTANEOUS		□ Vertebral Anomaly	Polyhydramnios
□ Acne	□ Short Stature		Prematurity
\Box Alopecia	_	LI NEOPLASIA	PULMONARY
□ Axillary/Inguinal Freckling	HEARING/VISION	Cancer Syndromes	□ CCAM/Small Thoracic Cavity
□ Blisters	Abnormality of Vision	Malignancies	Diaphragmatic Hernia
□ Café-Au-Lait Spots	Abnormality of Eye Movement	□ Breast	□ Eventration of Diaphragm
□ Congenital Nevus	□ Hearing Loss	□ CNS	□ Pleural Effusion
Ectodermal Dysplasia	Migraine Headache	□ Endocrine	□ Pulmonary Sequestration
	Optic Atrophy	GI	□ Respiratory Insufficiency
□ Loose Skin	□ Ptosis	🗆 GU	· · · · · · · · · · · · · · · · · · ·
L LOUSE SKIII	□ Retinopathy	□ Hamartoma	

OTHER SYMPTOMS List and describe	PREVIOUS TESTING & STUDIES	PREVIOUS TESTING & STUDIES List and describe				
COLLECTION REQUIREMENTS Samples not to exceed 3 tubes, regardless of testing						
CYSTIC FIBROSIS SCREENING (1) 2mL Lavender-top EDTA Tube LG11 TEST (1) 2mL Lavender-top EDTA Tube PRADER-WILLI/ANGELMAN DNA ANALYSIS (2) 2mL Lavender-top EDTA Tube SANGER SEQUENCING FOR TARGETED GENE (1) 2mL Lavender-top EDTA Tube THROMBOPHILIA RISK PANEL (1) 2mL Lavender-top EDTA Tube WARFARIN SENSITIVITY TEST (1) 2mL Lavender-top EDTA Tube	SOMA CYTOSCAN HD, TARGETED & WHOLE GENOME PERIPHERAL BLOOD	WHOLE EXOME SEQUENCING WHOLE BLOOD IN EDTA				