Centre universitaire de santé McGill University Health Centre	PATIENT STAMP OR LABEL HERE
MOLECULAR GENETICS REQUISITION	
Core Molecular Diagnostic Laboratory (CMDL)	Birth date (YYYY-MM-DD):* / /
1001 Decarie boul., E05.5051, Montreal, QC, H4A 3J1	Father's name:
Email: <u>molecular.genetics@muhc.mcgill.ca</u> Tel: 514-934-1934 x23383 / x23298	Mother's name:
Fax: 514-938-7405	Medical Record # (MRN):*
Referring Physician:*	RAMQ #:* For babies, please provide mother's RAMQ #
Name (Last, First):*	Sex:* 🔲 Male 🔲 Female 🔲 Unknown
License #:* Institution:*	Test Requested:*
Address:*	(Write below OR check boxes on page 2 if more than one test is requested)
E-mail address:*	Reason for Testing:*
Tel:* Fax:*	Diagnostic testing (affected case)
(Fax # to send results)	Carrier testing (unaffected case – for recessive conditions) Predictive testing (unaffected case – for dominant conditions)
Genetic counsellor/Nurse:	Prenatal testing (maternal sample required)
Tel: Fax:	Drug response (<i>pharmacogenetics</i>)
I acknowledge that the patient/guardian is aware of the benefits, limitations and risks associated with the requested test(s) and that I have obtained	□ Other – Specify:
informed consent to perform genetic testing for this patient. I authorize the laboratory to fax results to the number provided above.	Reason for expedited testing (if applicable):
aboratory to fax results to the number provided above.	Pregnancy (Gestational age: weeks on / /)
Signature:* Date:* / /	Other reason – Specify:
Sample Information:*	Familial Variant Testing: Please attach a copy of the proband's report. If the familial variant was not
Collection Date – Time:* / / at h min Collected by (Last, First):*	previously tested at the CMDL, please provide a sample from a family member known to be positive for this variant (i.e. a positive familial control).
□ 2 x 4 mL blood in EDTA tubes (purple top tube) – 2 mL for newborns	Gene (HGNC symbol):
DNA: min 10 µg – Source:	Variant(s) (HGVS format):
Amniotic fluid: min 10 mL	CMDL Family #:
Cultured amniocytes: 2 x T25 flasks (confluent)	Name of proband:
Direct CVS: min 10 mg direct villi	Relationship to proband:
Cultured CVS: 2 x T25 flasks (confluent)	Pedigree / Clinical Information:
Tissue – Specify:	Please draw or attach pedigree and provide relevant clinical information.
Other – Specify:	
Please check our test directory for test-specific requirements: https://muhc.ca/health-professionnals-and-teaching/muhc-clinical-laboratories	
CMDL - Laboratory use only:	
Date – Time received:	
h min SAMPLE LABEL(S) HERE	
Sample type and # of tubes:	Ethnicity:*
	Ordering Checklist:*
	 Specimen tubes labelled with at least two identifiers* Completed test requisition (this form)*
Patient #:	
Patient #: Family #:	Completed testing eligibility criteria form (if applicable)

CMDL - Laboratory use only:	PATIENT STAMP OR LABEL HERE
Date – Time received:	Name (Last, First);*
1 1	
h min SAMPLE LABEL(S) HERE	Birth date (YYYY-MM-DD):* / /
	Father's name:
Sample type and # of tubes:	Mother's name:
	Medical Record # (MRN):*
Patient #:	RAMQ #:*
Family #:	For babies, please provide mother's RAMQ # Sex:* Male Female Unknown
LIST OF TESTS & DISEASES	
Please use this page ONLY if more than one test is requested.	
For prenatal testing and analysis of familial variants, please provide information on page 1 (Familial Variant Testing section).	
Alpha-1-antitrypsin deficiency	Hexosaminidase A deficiency (Tay-Sachs disease)
SERPINA1 genotyping	HEXA sequencing
Amyotrophic lateral sclerosis (ALS)	Hidrotic ectodermal dysplasia (Clouston syndrome)
ALS panel (SOD1, FUS, TARDBP, C9ORF72, ATXN2)	□ <i>GJB</i> 6 sequencing
C9ORF72 repeat expansion	
ATXN2 repeat expansion	Huntington disease
Angelman / Prader-Willi syndrome: 🗌 Angelman 🔲 Prader-Willi	HTT repeat expansion
Methylation and deletion/duplication analysis	Male infertility
Uniparental disomy of chromosome 15 (parental samples required)	Y-chromosome microdeletion analysis
Ashkanari Jawish Carrier Sereening	MCAD definioner
Ashkenazi Jewish Carrier Screening ASPA, ELP1, HEXA sequencing (Canavan disease, familial	MCAD deficiency ACADM sequencing
dysautonomia, Tay-Sachs disease)	
Canavan diasaas	Methylmalonic acidemia
Canavan disease	MMACHC sequencing
	Nonsyndromic hearing loss
Cancer predisposition syndrome	DFNB1 panel (GJB2, GJB6 sequencing)
 Hereditary cancer predisposition syndrome panel Cancer predisposition single-gene sequencing – Specify: 	
	PAH deficiency: PKU Hyperphenylalaninemia
(Medical Genetics only)	PAH sequencing
Cystic fibrosis and CFTR-related conditions	Pharmacogenetics
CFTR sequencing	DPYD genotyping
	Destinated ensurelaidy
Familial dysautonomia	Postnatal aneuploidy QF-PCR (chromosomes 13, 18, 21, X, Y)
HBB-related hemoglobinopathies	For MUHC (Glen) only
Phenotype: HbS HbC HbE Other:	Bank DNA (Medical Genetics only)
β-thalassemia: 🗌 Major 🗌 Intermedia 🗌 Trait/Minor	Bank RNA (<i>Medical Genetics only</i>)
HBB sequencing	Maternal cell contamination analysis (<i>maternal sample required</i>)
Hereditary dyslipidemia	Zygosity analysis
APOE genotyping (dysbetalipoproteinemia)	Specimen matching analysis
LDLR, APOB, PCSK9 sequencing (familial hypercholesterolemia)	Other test – Specify:
	(Please contact us first for information)
Hereditary hemochromatosis	