



THE HOSPITAL FOR  
SICK CHILDREN  
Paediatric  
Laboratory Medicine

555 University Avenue  
Room 3416, Roy C. Hill Wing  
Toronto, ON, M5G 1X8, Canada  
Tel: 416-813-7200 x1  
Fax: 416-813-7732  
(CLIA # 99D1014032)

Patient Name:  
Preferred Name (if different):  
Date of Birth (DD/MM/YYYY):  
Legal Sex:  Male  Female  Non-binary/U/X  
Sex Assigned at Birth (if different):  Male  Female  Unassigned  
Gender Identity:  Male  Female  Non-binary/U/X  
MRN:  
Parent's Name:  
Address:  
For Canada Only  
Provincial Health Card #:  
Issuing Province:

Version:

## Genome Diagnostics

[www.sickkids.ca/en/care-services/for-health-care-providers/lab-testing-services](http://www.sickkids.ca/en/care-services/for-health-care-providers/lab-testing-services)

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### Referring Physician (required):

Name: \_\_\_\_\_  
Facility/Ward/Clinic (required): \_\_\_\_\_  
Address: \_\_\_\_\_  
Phone: \_\_\_\_\_ Fax: \_\_\_\_\_  
Email address: \_\_\_\_\_  
Signature: \_\_\_\_\_

### Reason for Testing (required):

- Diagnosis  Carrier testing  
 Familial mutation/variant Analysis  Prenatal testing  
 Bank DNA only  Variant re-assessment  
 Parental sample  
 Other (Specify): \_\_\_\_\_

### If expedited testing is requested, indicate reason:

- Pregnancy (Gestational age (weeks)) \_\_\_\_\_  
 Other (Specify): \_\_\_\_\_

### Copy Report To Another Healthcare Provider (all information is required):

Name: \_\_\_\_\_  
Address: \_\_\_\_\_  
Phone: \_\_\_\_\_ Fax: \_\_\_\_\_

### Familial Mutation / Targeted Variant Analysis:

**\*If proband testing was performed elsewhere, a copy of the original report (all pages) is required. Send a positive control sample if available.**

Gene & NM #: \_\_\_\_\_  
Mutation/variant(s): \_\_\_\_\_  
SickKids Laboratory/Order number: \_\_\_\_\_  
SickKids Pedigree/Family number: \_\_\_\_\_  
Name of proband: \_\_\_\_\_  
Relationship to proband: \_\_\_\_\_  
Name(s) & DOB of other submitted family members: \_\_\_\_\_

### Sample Information (required):

Date obtained (DD/MM/YYYY): \_\_\_\_\_ - \_\_\_\_\_ - \_\_\_\_\_ Referring laboratory reference #: \_\_\_\_\_

- Blood in EDTA (purple top tube): min. 4 mL (0.5-3 mL for newborns)  
 DNA: min. 10 ug in low TE buffer (Source: \_\_\_\_\_)  
\*Unable to perform MLPA analysis on externally extracted DNA (contact lab)  
 Direct CVS: min. 10 mg direct villi  
 Cultured villi: 1-2 confluent T25 flasks  
 Cultured amniocytes: 1-2 confluent T25 flasks  
 Tissue (Source: \_\_\_\_\_)  
 Other (Specify: \_\_\_\_\_)

### Closed consent:

- (If checked, all remaining DNA will be discarded upon notification by the ordering physician that all DNA testing has been completed)

### Laboratory Use:

Date (DD/MM/YYYY) | Time Received:

\_\_\_\_\_ | \_\_\_\_\_ h

Lab/Order #: \_\_\_\_\_

Specimen type, amt & # of tubes: \_\_\_\_\_

Comments:

Pedigree/Family No./Patient/Order No. \_\_\_\_\_ / \_\_\_\_\_

### Clinical Diagnostics and Family History (required):

Draw or attach a pedigree and provide any relevant information below, including clinical and family history details, as this is important for accurate interpretation of results.

Ethnicity: \_\_\_\_\_

### Ordering Checklist:

- Specimen tube labeled with at least two identifiers  
 Completed test requisition form  
**Clinical information must be provided for all tests. Pages 4-5 must be completed for all tests. Testing will not proceed until these are provided.**  
 Proband's report and positive control (familial/targeted variant testing only)  
 Completed billing form (page 6, if applicable)

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### LIST OF TESTS AVAILABLE BY DISEASE

For prenatal testing and cases where a familial mutation/variant is known, include information on page 1.

#### 22q11 Deletion Syndrome

- 22q11 deletion/duplication analysis (*external DNA not accepted*)

#### Angelman Syndrome

- Methylation and deletion/duplication analysis (*external DNA not accepted*)  
 UPD15 analysis (*please submit parental samples*)

#### Ashkenazi Jewish Carrier Screening

- Recurrent mutation analysis (7 diseases):  
Bloom syndrome, Canavan disease, Familial Dysautonomia, Fanconi  
Anemia Group C, Mucopolidosis Type IV, Niemann-Pick disease, Tay-Sachs  
disease

#### ETHNICITY (required):

- Ashkenazic  Sephardic  French Canadian  Cajun  
 Non-Jewish  Other \_\_\_\_\_

#### Autoinflammatory Disease \*

**Clinical information must be provided on pages 4 and 5**

- Autoinflammatory Diseases (AID) NGS panel  
 Recurrent Fever Syndrome (RFS) NGS panel  
 Hemophagocytic Lymphohistiocytosis (HLH) NGS panel  
 Aicardi-Goutieres Syndrome (AGS) NGS panel  
 Deletion/duplication analysis

#### Becker Muscular Dystrophy

- DMD deletion/duplication analysis (*external DNA not accepted*)  
 DMD sequence analysis

#### Beckwith-Wiedemann Syndrome

- IC1 and IC2 methylation† and 11p15 deletion/duplication analysis  
(*external DNA not accepted*)  
 UPD11 analysis  
 CDKN1C sequence analysis  
† No methylation analysis on CVS samples

#### Bone Marrow Transplantation

- Post-transplant monitoring

#### Caffey Disease

- COL1A1 recurrent mutation analysis

#### Cancer Related Tests

##### Li-Fraumeni Syndrome

- TP53 sequence analysis  
 TP53 deletion/duplication analysis (*external DNA not accepted*)

##### Rhabdoid Tumour Predisposition Syndrome

- SMARCB1 sequence analysis  
 SMARCB1 deletion/duplication analysis (*external DNA not accepted*)

#### Charge Syndrome

- CHD7 sequence analysis  
 CHD7 deletion/duplication analysis (*external DNA not accepted*)

#### Cherubism

- SH3BP2 recurrent mutation analysis  
 SH3BP2 sequence analysis

#### Congenital Muscular Dystrophies

- Sequence analysis panel:  
FKTN (FCMD), FKR, POMGnT1, POMT1, POMT2

#### Connective Tissue Disease \*

**Clinical information must be provided on pages 4 and 5**

**If more than one panel is requested, rationale must be provided on page 5.**

- Ehlers Danlos Syndrome NGS panel  
 Osteogenesis Imperfecta NGS panel  
 Osteopetrosis and Disorders of Increased Bone Density NGS panel  
 Bone Involvement NGS panel  
 Deletion/duplication analysis

#### Craniosynostosis

- Apert Syndrome (FGFR2 recurrent mutations analysis)  
 Crouzon Syndrome (FGFR2, FGFR3 recurrent mutation analysis)  
 Pfeiffer Syndrome (FGFR1, FGFR2, FGFR3 recurrent mutation analysis)  
 Saethre-Chatzen Syndrome (TWIST1 sequence analysis and FGFR3  
recurrent mutation analysis)  
 Non-Syndromic Craniosynostosis (FGFR3 recurrent mutation analysis)  
 TWIST1 deletion/duplication analysis (*external DNA not accepted*)

#### Cystic Fibrosis and/or CFTR-Related Disorders \*\*

**Indication (provide additional clinical details on page 1 and/or pages 4-5):**

- Fetal echogenic bowel (*ensure parental samples are linked to each other on  
both requisitions with at least two identifiers*)  
 Clinical diagnosis of cystic fibrosis  
 CFTR-related disorders  
 Male factor infertility:  oligo/azoospermia  C(B)AVD  
 Family history of cystic fibrosis  
 Positive newborn screen (*ensure familial samples are linked to each other  
on all requisitions with at least two identifiers; send NSO report*)

**Tests (indication specific):**

- CFTR recurrent mutation analysis  
 CFTR sequence analysis  
 CFTR deletion/duplication analysis (*external DNA not accepted*)

#### Dopamine Beta-Hydroxylase Deficiency

- DBH Sanger sequence analysis

#### Duchenne Muscular Dystrophy

- DMD deletion/duplication analysis (*external DNA not accepted*)  
 DMD sequence analysis  
 DMD mRNA analysis (*contact the laboratory before ordering*)

#### Fabry Disease

- GLA sequence analysis  
 GLA deletion/duplication analysis (*external DNA not accepted*)  
 GLA mRNA analysis (*contact the laboratory before ordering*)

#### Fragile X Syndrome & FMR1-related disorders

- Fragile X syndrome  
 Fragile X-associated primary ovarian insufficiency  
 Fragile X-associated tremor ataxia syndrome (FXTAS)

#### Fragile X E Syndrome \*\*\*

- AFF2 trinucleotide repeat analysis  
(*See testing requirements*)

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### LIST OF TESTS AVAILABLE BY DISEASE

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#### Fragile X E Syndrome \*\*\*

- AFF2 trinucleotide repeat analysis (See testing requirements)

#### Hearing Loss: Non-Syndromic, Autosomal Recessive

- GJB2 sequence analysis  
 GJB6 deletion/duplication analysis (external DNA not accepted)

#### Hearing Loss: Pendred Syndrome

- SLC26A4 sequence analysis  
 SLC26A4 deletion/duplication analysis (external DNA not accepted)

#### Hereditary Hearing Loss \*

**Clinical information must be provided on pages 4 and 5**

When the Common and Non-Syndromic Hearing Loss Panel is requested, STRC dosage is tested.

- Common and Non-Syndromic Hearing Loss NGS panel  
 Usher Syndrome NGS panel  
 Stickler Syndrome NGS panel  
 Alport Syndrome, Norrie Syndrome, Treacher Collins Syndrome, Waardenburg Syndrome NGS panel  
 Deletion/duplication analysis

#### Hereditary Hemorrhagic Telangiectasia

- ACVRL1 sequence analysis  
 ENG sequence analysis  
 ACVRL1 and ENG deletion/duplication analysis (external DNA not accepted)  
 SMAD4 sequence analysis

#### Hereditary Spastic Paraplegia \*

**Clinical information must be provided on pages 4 and 5**

- Autosomal Dominant HSP NGS panel  
 Autosomal Recessive HSP NGS panel  
 X-Linked HSP NGS panel  
 Deletion/duplication analysis

#### Hunter Disease

- IDS sequence analysis  
 IDS deletion/duplication analysis (external DNA not accepted)  
 IDS mRNA analysis (contact the laboratory before ordering)

#### Identity Testing

- Zygosity studies  
 Maternal Cell Contamination Studies (maternal sample required)

#### Neurofibromatosis type 1/Legius syndrome \*

**Clinical information must be provided on pages 4 and 5**

- NF1 sequence analysis  
 NF1 deletion/duplication analysis (external DNA not accepted)  
 SPRED1 sequence analysis  
 SPRED1 deletion/duplication analysis (external DNA not accepted)

#### Neuronal Ceroid Lipofuscinoses (Batten Disease)

- PPT1 (CLN1), TPP1 (CLN2) and CLN3 recurrent mutation analysis  
 Sequence analysis panel:  
PPT1 (CLN1), TPP1 (CLN2), CLN3 CLN5, CLN6, CLN7, CLN8, CLN10

#### Noonan Syndrome and RASopathies \*

**Clinical information must be provided on pages 4 and 5**

- Noonan Syndrome and RASopathies panel  
 Deletion/duplication analysis for SPRED1 only (external DNA not accepted)

#### Prader-Willi Syndrome

- Methylation and deletion/duplication analysis (external DNA not accepted)  
 UPD15 analysis (parental samples required)

#### Renal Diseases

- Atypical Hemolytic Uremic Syndrome / Membranoproliferative Glomerulonephritis sequence analysis  
 Focal Segmental Glomerulosclerosis sequence analysis

#### Russell-Silver Syndrome

- IC1 methylation and 11p15 deletion/duplication analysis (external DNA not accepted)  
 UPD7 analysis (parental samples required)

#### Shwachman-Diamond Syndrome

- SBDS sequence analysis

#### Simpson-Golabi-Behmel Syndrome

- GPC3 sequence analysis and GPC3 and GPC4 deletion/duplication analysis (external DNA not accepted)

#### Skeletal Dysplasia

- Achondroplasia (FGFR3 recurrent mutation analysis)  
 Hypochondroplasia (FGFR3 recurrent mutation analysis)  
 Thanatophoric Dysplasia (FGFR3 recurrent mutation analysis)

#### Spinal and Bulbar Muscular Atrophy

- AR trinucleotide repeat analysis

#### Spinal Muscular Atrophy

- SMN1 and SMN2 deletion/duplication analysis (external DNA not accepted)

#### Trismus Pseudocamptodactyly Syndrome

- MYH8 sequence analysis

#### X-Inactivation Analysis

- Other (PRIOR APPROVAL REQUIRED; CONTACT LABORATORY):

\*Next-Generation Sequencing (NGS) testing will only be initiated if the clinical information sections (pages 4-5) are completed. For more information on our Next-Generation Sequencing (NGS) panels, including the list of genes tested, visit our website: [www.sickkids.ca/en/care-services/for-health-care-providers/lab-testing-services](http://www.sickkids.ca/en/care-services/for-health-care-providers/lab-testing-services)

\*\*\* For information on the testing algorithm for Cystic Fibrosis, visit <https://www.sickkids.ca/en/care-services/for-health-care-providers/lab-tests/244-Cystic-Fibrosis/> on our website  
\*\*\* For information on the testing requirement for Fragile X E, visit the Specimen Requirements section for Fragile X E Syndrome on our website: [www.sickkids.ca/en/care-services/for-health-care-providers/lab-tests/250-FRAXE](http://www.sickkids.ca/en/care-services/for-health-care-providers/lab-tests/250-FRAXE)

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### DISEASE SPECIFIC FEATURES

#### Autoinflammatory Disorders (RFS/HLH)

- Abnormal inflammatory response
- Fevers
- Arthritis
- Pulmonary complications
- Gastrointestinal irritation
- Hepatosplenomegaly
- Lymphadenopathy
- Hemophagocytosis
- Oral ulcers
- Rash, specify: \_\_\_\_\_
- Ocular inflammation specify: \_\_\_\_\_
- Edema (periorbital, optic disk)
- Vision loss
- Other: \_\_\_\_\_

#### Hearing Loss

- Age of onset: \_\_\_\_\_
- Sensorineural hearing loss
- Conductive hearing loss
- Mixed hearing loss
- Bilateral  Unilateral
- Syndromic  Non-syndromic
- Ear anomalies  Ear tags
- Eye anomalies  Renal anomalies
- White forelock  Cardiac anomalies
- Hirschsprung disease
- Other: \_\_\_\_\_

#### Hereditary Spastic Paraplegia (HSP)

- Abnormal corpus callosum
- Cognitive impairment
- Ataxia  Spasticity
- Hyperreflexia  Seizures
- Hypertonia  Hypotonia
- Dystonia  Dysarthria
- Extensor plantar reflex
- Other: \_\_\_\_\_

The following investigations are required before molecular testing of HSP is undertaken:

- MRI – Brain and spinal cord
- Biochemical testing - Vitamin B12, vitamin E, very long chain fatty acids, lysosomal work-up, plasma amino acids and serum lipoprotein analysis (as appropriate)

#### Neurofibromatosis type 1 (NF1) / Legius Syndrome

- The patient meets the NIH criteria for a clinical diagnosis of NF1  
(**>2 of the clinical features below**).
- Café-au-lait macules  ≥6 CALS  (#: \_\_\_\_\_)
- Neurofibromas, ≥ 2 or ≥ 1 Plexiform
- Freckling, axillary or inguinal
- Optic glioma
- ≥2 Lisch nodules (iris hamartomas)
- Osseous lesion (type: \_\_\_\_\_)
- First degree relative diagnosed with NF1 by above criteria
- Other: \_\_\_\_\_
- The patient does not meet the NIH diagnostic criteria for NF1.  
Rationale for testing must be provided on page 5.

#### Connective Tissue Disorders (CTD)

##### Ehlers Danlos Syndrome (EDS)

Indicate the suspected clinical diagnosis in the patient:

- Classic  Vascular
- Kyphoscoliotic  Other: \_\_\_\_\_

Check applicable CTD features below.

##### Osteopetrosis and Disorders of Increased Bone Density

Check applicable CTD features below.

##### CTD Related Clinical Features:

- Joint hypermobility:  
Beighton score: \_\_\_\_\_
- Arterial aneurysms, dissection or rupture
- Intestinal rupture
- Molluscoid pseudotumors
- Subcutaneous spheroids
- Loose/stretchable skin
- Smooth/velvety skin
- Widened atrophic scars

##### Osteogenesis Imperfecta (OI)

If the patient does not present with one of the test indications below, rationale for testing must be provided on page 5.

- Fetal findings on anatomy ultrasound consistent with OI.
  - Fractures with minimal or no trauma in the absence of other known disorders of bone metabolism.
  - Vertebral fractures
  - Dentinogenesis imperfecta
  - Low ALP for age/gender (ALPL analysis only will be performed)
- Check applicable CTD features below.

##### Bone Involvement

Check applicable CTD features below.

- Recurrent spontaneous tendon rupture
- Easy bruising
- Myopia
- Lens dislocation
- Blue/gray sclerae
- Thumb or wrist sign
- Club foot
- Scoliosis
- Marfanoid habitus
- Short stature
- Shortened long bones
- Recurrent pneumothoraces
- Joint subluxations/dislocations
- Fractures
- Bone deformity
- Wormian bones
- Increased bone mineral density
- Diaphyseal sclerosis
- Hearing loss
- Osteosclerosis
- Other: \_\_\_\_\_

##### Noonan Syndrome and RASopathies

- Increased nuchal translucency
- Developmental delay
- Characteristic facies
- Broad or webbed neck
- Heart defect (specify: \_\_\_\_\_)
- Hypertrophic cardiomyopathy
- Short stature (%ile: \_\_\_\_\_)
- Pectus deformity
- Lymphatic dysplasias
- Characteristic hematological abnormality (specify: \_\_\_\_\_)
- Other RASopathy features: (specify: \_\_\_\_\_)
- For postnatal patients: The patient must present with ≥ 2 of the above features for molecular testing to be undertaken.

### FAMILY HISTORY (Required)

Draw or attach a pedigree and provide any relevant information below, including clinical and family history details, as this is important for accurate interpretation of results.

Ethnicity: \_\_\_\_\_

## Genome Diagnostics

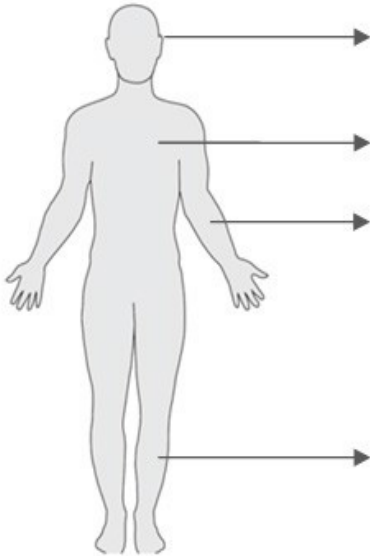
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### ADDITIONAL RELEVANT CLINICAL INFORMATION

#### Previous Genetic Testing

- No  
 Yes – Test Results: \_\_\_\_\_



### GENERAL CLINICAL FEATURES

#### Perinatal history

- Premature birth  
 IUGR  
 Oligohydramnios  Polyhydramnios  
Other: \_\_\_\_\_

#### Growth

- Failure to thrive  
 Growth retardation/short stature  
 Overgrowth  
 Macrocephaly  Microcephaly  
Other: \_\_\_\_\_

#### Physical/cognitive development

- Delayed fine motor development  
 Delayed gross motor development  
 Delayed speech and language  
 Autistic behavior  
 Intellectual disability  
 Developmental regression  
Other: \_\_\_\_\_

#### Behavioral

- Autistic features  
 Obsessive-compulsive disorder  
 Other psychiatric symptoms  
Other: \_\_\_\_\_

#### Cancer/Malignancy

- Age of onset: \_\_\_\_\_  
 Tumor type: \_\_\_\_\_  
 Location(s): \_\_\_\_\_

#### Craniofacial/Ophthalmologic

- Abnormal face shape  
 Blindness  Cataracts  
 Coloboma  Optic atrophy  
 Ophthalmoplegia  Ptosis  
 Retinitis pigmentosa  
 Oral cleft  
Other: \_\_\_\_\_

#### Brain malformations/abnormal imaging

- Abnormality of the basal ganglia  
 Agenesis of the corpus callosum  
 Brain atrophy  
 Cortical dysplasia  
 Hemimegalencephaly  
 Heterotopia  
 Holoprosencephaly  
 Hydrocephalus  
 Lissencephaly  
 Periventricular leukomalacia  
Other: \_\_\_\_\_

#### Cardiac/congenital heart malformations

- ASD  VSD  
 Coarctation of aorta  
 Hypoplastic left heart  
 Tetralogy of Fallot  
 Cardiomyopathy  
 Arrhythmia/conduction defect  
Other: \_\_\_\_\_

#### Gastrointestinal

- Gastroschisis/omphalocele  
 Gastrointestinal reflux  
 Pyloric stenosis  
 Tracheoesophageal fistula  
 Hepatic failure  
 Chronic intestinal pseudo-obstr.  
 Hirschsprung disease  
 Recurrent vomiting  
 Chronic diarrhea  
 Constipation  
Other: \_\_\_\_\_

#### Genitourinary abnormalities

- Ambiguous genitalia  
 Cryptorchidism  
 Hypospadias  
 Hydronephrosis  
 Kidney malformation  
 Renal agenesis  
 Proximal renal tubulopathy  
Other: \_\_\_\_\_

#### Endocrine

- Diabetes mellitus Type 1  
 Diabetes mellitus Type 2  
 Hypothyroidism  
 Hypoparathyroidism  
 Pheochromocytoma/paragan glioma  
Other: \_\_\_\_\_

#### Neurological/Muscular

- Ataxia  Hypotonia  
 Chorea  Hypertonia  
 Dystonia  Spasticity  
 Exercise intolerance/ easy fatigue  
 Headache/migraine  
 Muscle weakness  
 Seizures (type: \_\_\_\_\_)  
 Stroke/stroke-like episodes  
Other: \_\_\_\_\_

#### Skeletal/Limb abnormalities

- Contractures  Club foot  
 Polydactyly  Syndactyly  
 Vertebral anomaly  Scoliosis  
Other: \_\_\_\_\_

#### Skin/Hair

- Abnormality of the hair pattern, quantity  
 Abnormal nail growth  
 Abnormal pigmentation  
 Café-au-lait macules  
 Neoplasms of the skin  
 Neurofibromas  
 Blistering  
 Ichthyosis  
Other: \_\_\_\_\_



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Tel: 416-813-7200 x1  
Fax: 416-813-7732  
(CLIA # 99D1014032)

Patient Name:  
Preferred Name (if different):  
Date of Birth (DD/MM/YYYY):  
Legal Sex:  Male  Female  Non-binary/U/X  
Sex Assigned at Birth (if different):  Male  Female  Unassigned  
Gender Identity:  Male  Female  Non-binary/U/X  
MRN:  
Parent's Name:  
Address:

For Canada Only  
Provincial Health Card #:  
Issuing Province:

Version:

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### BILLING FORM

**Completion of Billing Form NOT required for patients with an Ontario Health Card Number.**

The hospital, referring laboratory, referring physician, or a patient/guardian will be billed for the services rendered, upon direction from the referring physician.

- Invoices are sent upon completion of each test/service.
- Invoices are itemized and include the date of service, patient name, CPT code, test name and charge.
- Contact SickKids' Genome Diagnostics Laboratory at 416-813-7200 x1 with billing inquiries.

#### How to complete the Billing Form:

- Referring Physician completes the appropriate section below to specify billing method.
- Send requisition and completed "Billing Form" with specimen.

#### Section 1: Complete to have the Healthcare Provider billed:

Referring Laboratory's Reference #: \_\_\_\_\_

Billing address of hospital, referring laboratory, clinic, referring physician, or medical group (if different from requisition):

Name: \_\_\_\_\_

Address: \_\_\_\_\_

City: \_\_\_\_\_ Prov/State: \_\_\_\_\_

Postal/Zip Code: \_\_\_\_\_ Country: \_\_\_\_\_

Contact Name: \_\_\_\_\_ Contact Telephone #: \_\_\_\_\_

#### Section 2: Complete to have Patient/Guardian billed directly:

*If electing to have patient/guardian billed:*

- Patient/Guardian billing information below must be complete; otherwise, the healthcare provider will be billed.
- Advise the patient/guardian to expect a bill from the Genome Diagnostics laboratory.
- The patient's valid credit card information must be provided.
- Unfortunately, personal checks are not accepted as a method of payment.
- **In this case, the patient/guardian is solely responsible for the charges.**

Send bill to (check one):  Patient  Guardian

Method of Payment (check one):  American Express  MasterCard  Visa

Name as it appears on credit card: \_\_\_\_\_

Credit card #: \_\_\_\_\_

Expiry date on credit card: \_\_\_\_\_

Signature of credit card holder (Required): \_\_\_\_\_

#### Mailing Address of Patient/Guardian (if different from requisition):

Name: \_\_\_\_\_

Address: \_\_\_\_\_

\_\_\_\_\_ Apt. #: \_\_\_\_\_

City: \_\_\_\_\_ Prov/State: \_\_\_\_\_

Postal/Zip Code: \_\_\_\_\_ Country: \_\_\_\_\_

#### Additional Contact Information

Patient's phone # with area code:

\_\_\_\_\_

- or -

Guardian's phone # with area code:

\_\_\_\_\_