

# Molecular Genetics Laboratory

## Cancer and Endocrine NGS Requisition

Scanning Label or Accession # *(lab only)*

For detailed testing information, refer to **APL Genetics & Genomics Webpage** (<http://ahsweb.ca/lab/if-lab-genetics-and-genomics>) and **APL Test Directory** (<http://ahsweb.ca/lab/apl-td-lab-test-directory>)

<b>Patient</b>	PHN		Expiry: _____		Date of Birth <i>(dd-Mon-yyyy)</i>	
	Legal Last Name			Legal First Name		Middle Name
	Alternate Identifier		Preferred Name		<input type="checkbox"/> Male <input type="checkbox"/> Female	Phone
					<input type="checkbox"/> Non-binary <input type="checkbox"/> Prefer not to disclose	
Address			City/Town		Prov	Postal Code
<b>Provider(s)</b>	Authorizing Provider Name <i>(last, first, middle)</i>			Copy to Name <i>(last, first, middle)</i>		Copy to Name <i>(last, first, middle)</i>
	Address			Phone	Address	Address
	CC Provider ID	CC Submitter ID	Legacy ID	Phone	Phone	
	Clinic Name			Clinic Name	Clinic Name	
<b>Collection</b>		Date <i>(dd-Mon-yyyy)</i>	Time <i>(24 hr)</i>	Location	Collector ID	
Genetic Counsellor/Clinic Contact Name <i>(last, first)</i>					Phone	
<b>Specimen</b>						
<input type="checkbox"/> Whole Blood in EDTA tube		<input type="checkbox"/> Extracted DNA		<input type="checkbox"/> Fluid, amniotic*		
<input type="checkbox"/> Tissue, chorionic villi*		<input type="checkbox"/> Cord blood*		<input type="checkbox"/> Other <i>(specify):</i> _____		
*If specimen type is prenatal or cord blood, maternal specimen must be collected for maternal cell contamination studies						
<b>Health Care Provider Important Information</b>						
1. All sections of the requisition must be completed.						
2. By providing this requisition to the patient/family, the health care provider confirms that they have reviewed the pre-test counselling information (available on the Genetics & Genomics website) with the patient/family, and the patient/family consents to testing.						
3. Direct patient to take requisition to a local blood collection location to have blood specimen drawn.						
<b>Billing Information:</b> Must be completed if the patient does not have a valid Alberta Personal Health Number						
Genetic testing is not covered by inter-provincial billing agreements. Alberta Precision Laboratories (APL) will bill a provincial medical services plan provided there is a letter of pre-approval received with the requisition or Institutional Billing information provided below. By completing the Institutional Billing section, the health care provider confirms they have obtained any necessary pre-approval. For patient pay, contact the testing laboratory.						
<b>Institutional Billing Information <i>(if pre-approval letter not attached)</i></b>						
Address						
Contact Name <i>(last, first)</i>						
Phone			Fax			
<b>MGL Use Only</b>						
Patient Number		Family Number		Rec'd	Quantity	

## Molecular Genetics Laboratory Cancer and Endocrine NGS Requisition

Last Name (Legal)	First Name (Legal)
PHN	

### Section I - Reason for Testing (Select one only)

**Confirmation of Diagnosis.**

Patient has signs or symptoms of the disease / disorder.

**Presymptomatic or Predictive Testing.**

Patient does not presently have symptoms; positive family history

**Carrier Testing.**

No symptoms; at risk of being a carrier of a recessive disorder

**Required for Family Study.**

**Prenatal Testing**

**Other** \_\_\_\_\_

### Section II - Family History of Indicated Disease

Unknown family history

No known family history

Possible family history

#### Documented family history

Clinical Diagnosis ONLY

Molecular Diagnosis (provide a copy of the familial variant report and complete information in Section V)

**Is RUSH testing needed?**  Yes (provide details below)

Results will alter the **immediate** management and/or treatment of this patient (specify): \_\_\_\_\_

Results will impact an ongoing pregnancy (provide EDD, and procedure date if applicable): \_\_\_\_\_

If RUSH testing is required, please provide a target date (yyyy-Mon-dd) (required): \_\_\_\_\_

Note: TAT is a minimum of 4 weeks.

### Section III - Patient Clinical Information

Sex at birth  Female  Male  Unknown

Date of last chemotherapy (if applicable) → Date (yyyy-Mon-dd) \_\_\_\_\_

Has this patient received a blood product in the preceding three months?

Yes indicate blood product: \_\_\_\_\_

Has the patient had a bone marrow transplant?  Yes (**Blood is an incompatible specimen type.**)

Please provide any relevant information regarding your patient's clinical presentation (ex. tumour site, age at diagnosis, multiple primary tumour, pathology, hormone receptors):

If applicable: IHC result (required for Lynch testing): \_\_\_\_\_

Has the patient had BRCA 1/2 testing of tumour tissue? \_\_\_\_\_

### Section IV - Pedigree (Provide any relevant family history details, with family member names, ages, and diagnoses included as applicable. If more space is required, attach a separate sheet.)

Patient Ethnicity / Ancestry: \_\_\_\_\_

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**Section V - Specific Variant Testing**

Complete this section only if you are requesting testing for a variant previously identified in the family. Specific variant testing is available for all genes available on the panels listed below.

Inherited Cancer and Endocrine Gene Panels, Specific Variant

Gene	Mutation / Variant

What is the phenotype / presentation in the proband? \_\_\_\_\_

Other family members previously tested in MGL  No  Yes ▼

INDEX patient name	MGL Reference Number
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Which laboratory performed the proband testing?  Calgary  Edmonton  Other *(specify)* \_\_\_\_\_

Testing a positive control is recommended if the proband testing was performed at another lab.  
**A clear copy of the test report on a family member is required if the testing was performed at another laboratory**

**Section VI - Cancer and Endocrine NGS Panel Requests**

**Breast and Ovarian Cancer Panels**

- Breast Ovarian Cancer Panel, Core
- Breast Ovarian Cancer Panel, Extended
- Ovarian Cancer Panel

**Endocrine Disorder Panels**

- Endocrine Neoplasia Panel
- Paraganglioma/Pheochromocytoma Predisposition Panel
- Renal Cancer Panel

**Gastrointestinal Cancer Panels**

- Gastrointestinal/Polyposis Panel
- Gastrointestinal Stromal Tumor Panel
- Lynch Syndrome Panel
- Pancreatic Cancer Panel

**Hematological Panels**

- Familial Acute Myeloid Leukemia Panel
- Fanconi Anemia and DNA Repair Disorders Panel

**Overgrowth Panels**

- Hereditary Multiple Osteochondromatosis Panel
- Overgrowth Panel

**Pediatric Panels**

- Pediatric Cancer Panel
- Schwannomatosis Panel
- Tuberous Sclerosis

**Skin Panels**

- Skin Cancer Panel
- Xeroderma Pigmentosum Panel

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**Section VII - Single Gene Test Request**

- CASR-Related Disorder
- Inherited Cancer and Endocrine Custom Gene Panel
- Inherited Cancer and Endocrine Single Gene (complete only if panel is not appropriate for patient)

Requesting test for \_\_\_\_\_ *(indicate the gene)* **and** the presenting phenotype: \_\_\_\_\_

Requesting test for \_\_\_\_\_ *(indicate the gene)* **and** the presenting phenotype: \_\_\_\_\_

Neurofibromaosis Type 1 ***Please indicate if your patient has the following***

- Greater than 6 café-au-lait macules greater than 5 mm, postpubertal greater than 15 mm
- Greater than 2 neurofibromas or 1 plexiform neurofibroma
- Axillary or inguinal freckling
- Optic glioma
- Greater than 2 Lisch nodules
- A distinctive osseous lesion
- A first degree relative with NF1 per the above criteria

If the patient does not fulfill NIH diagnostic criteria for NF1, please provide reason for testing as a comment:

**Section VIII - Variant Reinterpretation**

Complete this section only if you are requesting reinterpretation of a variant previously identified in the family

Gene	Mutation / Variant
What is the clinical phenotype / presentation in the family?	

**A clear copy of the test report is required if the testing was preformed at another laboratory**

**Section IX - Additional Comments**