



Benefits of providing COMPLETE and LEGIBLE information:

- Reduces turnaround time when processing patient samples
Promotes patient safety through reduced transcription errors

RUH SCH SPH Other

DEPARTMENT OF PATHOLOGY AND LABORATORY MEDICINE

GENERAL REQUISITION – CYTOGENETICS AND MOLECULAR GENETICS TESTING

Page 1 of 2

Form with sections: PATIENT INFORMATION, PHYSICIAN INFORMATION, SPECIMEN REQUIREMENTS, TEST PRIORITY, TEST REQUESTED, and For Laboratory Use Only. Includes various checkboxes, text boxes, and callouts.

Patient identification MUST include:
First and last name
PHN or other unique identifier
Date of birth
This information may be on a label or in writing.

Please provide physician's FULL name

Cytogenetics specimen information

Molecular specimen information

Indicate priority

Indicate which test is being ordered

THIS IS A TWO-PAGE FORM. COMPLETE IN FULL TO AVOID DELAY IN TESTING.

GENERAL REQUISITION – CYTOGENETICS AND MOLECULAR GENETICS TESTING

Page 2 of 2

Genomics Laboratory
 Royal University Hospital
 Room 5617, 5th Floor, 1955 Building
 103 Hospital Drive, Saskatoon, SK S7N 0W8
 Phone: 306-655-1706 Fax: 306-655-6462
 Email: cytogenetics@saskatoonhealthregion.ca

CLINICAL INDICATIONS (Please complete to ensure accurate and timely reporting)		
Prenatal: <input type="checkbox"/> Gestational age: _____ (low) <input type="checkbox"/> Twin pregnancy: <input type="checkbox"/> Twin A <input type="checkbox"/> Twin B <input type="checkbox"/> Positive MSS (include report) <input type="checkbox"/> Maternal age greater than 40 <input type="checkbox"/> Ultrasound anomalies (specify below) <input type="checkbox"/> Fetal growth restriction <input type="checkbox"/> Previous pregnancy with chromosomal abnormality (specify below) <input type="checkbox"/> Family history of chromosome abnormality (specify below) <input type="checkbox"/> Family history of genetic disease (must be referred through Medical Genetics) <input type="checkbox"/> Other: _____	<div style="border: 1px solid purple; border-radius: 15px; background-color: #e6e6fa; padding: 10px; margin-bottom: 10px;"> Provide clinical information: <ul style="list-style-type: none"> Why is the test being requested? What is the diagnosis? Check off indication or choose Other and write information </div> <input type="checkbox"/> Failure to thrive <input type="checkbox"/> Developmental delay <input type="checkbox"/> Sex chromosome abnormality <input type="checkbox"/> Short stature <input type="checkbox"/> Excessive growth <input type="checkbox"/> Infertility <input type="checkbox"/> Multiple miscarriages (greater than/equal to 3) <input type="checkbox"/> Family history of chromosome abnormality <input type="checkbox"/> Other: _____	Oncology: <input type="checkbox"/> New diagnosis <input type="checkbox"/> Follow up <input type="checkbox"/> Relapse <input type="checkbox"/> Post-treatment <input type="checkbox"/> Post-transplant <input type="checkbox"/> Potential donor <input type="checkbox"/> AML <input type="checkbox"/> APL <input type="checkbox"/> ALL <input type="checkbox"/> MDS <input type="checkbox"/> CML <input type="checkbox"/> CLL <input type="checkbox"/> MM/PCD <input type="checkbox"/> Lymphoma <input type="checkbox"/> MPD
FAMILY HISTORY, PEDIGREE, AND ADDITIONAL INFORMATION		
PRE-TEST CONSENT FOR GENETIC TESTING		
<p>The following points are to be discussed with the patient/guardian by the health-care provider prior to order genetic testing:</p> <ol style="list-style-type: none"> 1. Specimen analysis will be limited to related clinical condition(s) as indicated above. 2. For genetic testing, patient sample(s) will be collected in order to extract DNA or otherwise process for genetic testing and any remaining specimen will be stored by the Genomics Lab. If there is enough sample left when testing is complete, it may be used anonymously (identifying information will be removed) for research and quality assurance purposes. 3. Correlation with clinical information may be required for accurate interpretation. Correct interpretation of results can also depend on accurate family relationships being provided. 4. The genetic test(s) may not detect all genetic abnormalities associated with this condition. The accuracy of the genetic test requested as well as the potential benefits and limitations have been reviewed with the patient. 5. DNA analysis may reveal a change in a gene that has unknown clinical significance. It may also reveal unexpected information about family relationships (e.g., non-paternity). 6. The results of this testing are entirely confidential but may be used without identifying information for the interpretation of genetic testing for family members. 7. Genetic test results can be complex and may have important implications. When the genetic testing is complete, results will be discussed with the patient by the health-care provider. 8. Genetic counseling is available if required by the patient/guardian. <p>I have reviewed the points above, as well as the risks and benefits of genetic testing, and I have answered all the patient's/guardian's questions. I consent to the indicated testing(s).</p>		
Ordering Physician: _____ Printed Name	Ordering physician name and signature are REQUIRED for the patient/guardian. _____ Signature (Required)	
Date: _____		

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