

**Panorama Funded by MOHLTC**

Must include **MOHLTC CHECKLIST**, page 2 of this document  
Microdeletions are **NOT** funded - private pay

<b>Ordering Physician Billing #:</b>				<b>LifeLabs</b> Demographic Label	
<b>Ordering Physician:</b>		Name			
<b>Ordering Physician Address &amp; Contact Information:</b>		Tel: _____ Fax: _____			
<b>Physician Signature:</b>		Confirmation of patient consent: I confirm that this patient has been informed about the details associated with the genetic test(s) ordered below including its risks, benefits and limitations, and has given consent to testing as may be required by applicable law.			
<b>Copy to:</b> <input type="checkbox"/> Genetic Counsellor <input type="checkbox"/> Other Healthcare Provider		Name _____ Tel: _____ Fax: _____		Demographic Label	
<b>Bill to:</b>		Bill to Type 'H' OHIP (patient ONLY pays for Microdeletions if ordered)			
<b>Patient Last Name:</b>			<b>Patient First Name:</b>	<b>Date of Birth:</b>	
				M M D D Y Y Y Y	
<b>Unit #:</b>	<b>Street:</b>	<b>City:</b>	<b>Province:</b>	<b>Postal Code:</b>	
<b>Ontario Health Card #:</b>		# # # # # # # # # # V C		<b>Patient Telephone #:</b> ( ) -	
<b>CLINICAL QUESTIONS</b> <i>all fields are required; incomplete requisitions may result in testing delays</i>	<b>Multiple gestation?</b> <input type="checkbox"/> Y <input type="checkbox"/> N ↳ If Y: Ongoing Twins? <input type="checkbox"/> OR >2? <input type="checkbox"/> Optional: <input type="checkbox"/> Monochorionic <input type="checkbox"/> Dichorionic <input type="checkbox"/> Unknown <i>Panorama does <u>not</u> accept twins conceived using a surrogate or egg donor, high order multiple gestations (&gt;2), or vanishing twins</i>		<b>Egg donor?</b> <input type="checkbox"/> Y <input type="checkbox"/> N ↳ If Y: donor's age at egg retrieval: _____		
			<b>Surrogate?</b> <input type="checkbox"/> Y <input type="checkbox"/> N <b>Vanishing twin?</b> <input type="checkbox"/> Y <input type="checkbox"/> N		
	<b>Due Date:</b>		<i>Patient <u>must</u> be at least 9 weeks gestation at the time of blood draw</i>		
			M M D D Y Y Y Y		
<b>Maternal Weight:</b>		lbs.			

TESTS REQUESTED			
<b>Singleton pregnancies ONLY</b> please select only one of the following options:			<u>LL TR</u>
<input type="checkbox"/>	<b>Panorama® Prenatal Test (no cost to patient)</b> Testing of chromosomes 21, 13, 18, X, Y and triploidy. (Monosomy X + triploidy not screened in dizygotic pregnancies or pregnancies conceived with an egg donor or surrogate)		5518
<input type="checkbox"/>	<b>Panorama® Prenatal Test + 22q11.2 deletion (\$195)</b> Testing of chromosomes 21, 13, 18, X, Y, triploidy, and 22q11.2 deletion. <i>Not available for dizygotic twins, egg donors or surrogates</i>		5518 & 3037
<input type="checkbox"/>	<b>Panorama® Prenatal Test + Microdeletion Extended Panel [5] (\$245)</b> Testing of chromosomes 21,13,18,X,Y, triploidy, 22q deletion, Cri-du-chat,1p36 deletion, Angelman, Prader-Willi <i>Not available for twins, egg donors or surrogates.</i>		5518 & 3071
<input type="checkbox"/>	<b>YES, include the sex of the baby on the report (no cost)</b> – if the box is not ticked, the sex of the baby will not be reported		
<b>Date Blood Collected:</b>		<b>Time Blood Collected:</b>	<b>Collector Name:</b>
M M D D Y Y Y Y		H H M M	

**\*\* LIFELABS: PHOTOCOPY REQUISITION, INCLUDE 1 COPY WITH SAMPLES \*\***

Singleton pregnancies: Panorama Prenatal Test performed by LifeLabs Genetics (175 Galaxy Blvd., Suite 105, Toronto ON, M9W 0C9, Canada)  
Twins, egg donors, surrogate pregnancies: Panorama Prenatal Test performed by Natera Inc. (410 – 201 Industrial Road, San Carlos CA, 94070, USA)

**PATIENT CONSENT - MANDATORY:**

I have read and signed the Patient Consent Form, which remains with the ordering physician. I understand that 2 blood samples will be taken by LifeLabs staff. I acknowledge that my sample(s) and personal health information will be sent to LifeLabs and/or Natera for the purpose of non-invasive prenatal testing. I also understand that LifeLabs will contact me for a new blood sample if a test result cannot be provided from the original blood samples. I acknowledge that LifeLabs will send the results to my ordering physician and, if testing is performed at Natera, LifeLabs will receive results from Natera and send the results to my ordering physician. Should we be asked to disclose information about you for another reason, other than as required or permitted by law, we will contact you to obtain your consent. In the event of a high risk or no result, I acknowledge that LifeLabs may contact my healthcare provider to obtain follow-up diagnostic information to ensure quality and accuracy in reporting.

**Patient Sign Here:** \_\_\_\_\_ **Date:** \_\_\_\_\_

M M D D Y Y Y Y

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**CHECKLIST**  
 must be submitted with requisition

<b>Patient Name:</b> (Last, First)	<b>Patient Healthcard #:</b>												
The Provincial Council for Maternal and Child Health (PCMCH) has recommended specific indications for NIPT funding. <b>Please complete either Category I or II and attach to page 1 of the Panorama Funded by MOHLTC requisition.</b> Please confirm that your patient meets the following indications by checking the appropriate boxes. <b>Ordering physician on page 1 must match physician information/signature on page 2 CHECKLIST</b>													
<b>CATEGORY I: For investigation of trisomy 21, 18 or 13 ONLY.</b> <input type="checkbox"/> <b>Singleton gestation</b> (NIPT in the context of twin pregnancies requires consultation with a geneticist or maternal fetal medicine specialist (see Section B)) with appropriate pre-test counselling including a discussion of the limitations of the test. <b>And any one of the following:</b> <input type="checkbox"/> A maternal multiple marker screening test (eg. FTS/IPS/Quad etc.) positive for aneuploidy. <input type="checkbox"/> Women of advanced maternal age, defined as $\geq 40$ years of age at expected time of delivery. <input type="checkbox"/> Fetal nuchal translucency (NT) $\geq 3.5$ mm <input type="checkbox"/> Pregnancy history of aneuploidy / previous child with aneuploidy.													
<b>Physician signature</b>	<b>Date</b>												
<b>CSN#</b> (Billing #)													
<b>CATEGORY II:</b> There are several situations where additional specialist consultation is necessary to determine whether NIPT is warranted and to provide appropriate pre and post- test counselling. <b>NIPT funding for the following criteria must be submitted by a genetics or maternal fetal medicine (MFM) specialist.</b>  <b>Risk indicators:</b> <b>A/</b> <input type="checkbox"/> Fetal congenital anomalies identified on ultrasound, which are suggestive of trisomy 21, 18 or 13. Specify: _____  <b>OR:</b> <b>B/</b> <input type="checkbox"/> Risk of aneuploidy for trisomy 21, 18 or 13 > than that of a positive maternal multiple marker screen. <ul style="list-style-type: none"> <li>o Women less than 40 years of age at expected date of delivery must have at least one other risk factor noted.</li> <li>o The risk of aneuploidy can be calculated by including any combination of risk indicators including soft markers, biochemistry, maternal age, etc.</li> </ul> Please indicate all risk factors present <input type="checkbox"/> Twin pregnancy <input type="checkbox"/> Soft markers (check all that apply): <table border="1" style="margin-left: 20px; border-collapse: collapse; width: 60%;"> <tr><td style="padding: 2px;">Absent nasal bone</td><td style="padding: 2px;">Increased nuchal fold / edema</td></tr> <tr><td style="padding: 2px;">Choroid plexus cysts</td><td style="padding: 2px;">Increased nuchal translucency</td></tr> <tr><td style="padding: 2px;">Clinodactyly</td><td style="padding: 2px;">Intracardiac echogenic focus / foci</td></tr> <tr><td style="padding: 2px;">Cystic hygroma</td><td style="padding: 2px;">Short femur</td></tr> <tr><td style="padding: 2px;">Hyperechogenic bowel</td><td style="padding: 2px;">Short humerus</td></tr> <tr><td style="padding: 2px;">Hypoplastic nasal bone</td><td style="padding: 2px;">Ventriculomegaly</td></tr> </table> <input type="checkbox"/> Maternal age: _____ <input type="checkbox"/> Other, specify: _____		Absent nasal bone	Increased nuchal fold / edema	Choroid plexus cysts	Increased nuchal translucency	Clinodactyly	Intracardiac echogenic focus / foci	Cystic hygroma	Short femur	Hyperechogenic bowel	Short humerus	Hypoplastic nasal bone	Ventriculomegaly
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<b>OR:</b> <b>C/</b> <input type="checkbox"/> NIPT for sex chromosome determination (at least one of the following): <ul style="list-style-type: none"> <li><input type="checkbox"/> risk of a sex-limited disorder</li> <li><input type="checkbox"/> ultrasound findings suggestive of either a sex chromosome aneuploidy</li> <li><input type="checkbox"/> ultrasound findings suggestive of a disorder of sex determination (DSD)</li> </ul>													
<b>Genetics or MFM specialist's name</b> (Please print)													
<b>Physician Signature</b>	<b>Date</b>												
<b>CSN#</b> (Billing #)	<b>Genetics or MFM Centre</b>												