For PDF Fillable Requisitions, the following applies:

- 1. The form shall be completed using a Digital Health assigned computer.
- 2. Absolutely no personal health information shall be electronically saved on a computer.
- 3. The completed form shall not be shared electronically. If you reasonably believe that e-mailing the information is the only available method of communication or the only way to send the information then you must adhere to the Privacy guideline titled "E-mailing Personal Health Information".
- 4. All forms must be completed in their entirety, e.g. if a staff member has only completed half of a form they cannot save their work and then come back to complete it at a later date.
- 5. Once the personal health information has been recorded onto the form, it is to be printed immediately, deleted (not saved) from the computer, and then stored securely inside the client (paper) health record or scanned into an electronic record.
- 6. Do not print unnecessary duplicate copies of the form.
- 7. Regular audits of the Digital Health assigned computer shall be undertaken to ensure that no personal health information is being duplicated and saved.

 Please have all specimens delivered to:
 Additional requisitions / sample requirements at:

 Central Services
 https://apps.sbgh.mb.ca/labmanual/test/findTestPrepare

 MS551 820 Sherbrook Street
 Cytogenetics Laboratory Health Sciences Centre

 Winnipeg Manitoba R3A 1R9
 Phone 204-787-2489 Fax 204-787-1384

Place LIS Sticker Here

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 ¹Chromosome Analysis Specimen Collection: 2-4mL in NaHep. 	□ Ir □ Pi □ Fa	 Infection model reading reacting re						
 ²Microarray Specimen Collection: 4mL in EDTA <u>and</u> 4mL in NaHep. Infants <1 year of age: 1mL in EDTA <u>and</u> 1mL in NaHep. 		 Developmental delay/ Intellectual disability Multiple congenital anomalies (list in Additional information) Dysmorphic features (list in Additional information) 			□ Fa Pr □ 22	 Autism spectrum disorder Family studies Proband: 22q11.2 targeted array Other: 		
 FISH (Clinical Geneticists only) Specimen Collection: 2-4mL in NaHep. 		 Ambiguous genitalia Turner syndrome confirmation Sex chromosome mosaicism X-linked ichthyosis 		 Family studies Proband: Microarray follow-up ³Other: 				
 Chromosome Breakage Studies (Clinical Geneticists and Hematolog Specimen Collection: 2-6mL in NaHep Additional information: 	ists only)	taxia Telangiectasia anconi Anemia (Sar		l by noon, -	the same day	of collectio	on, Mond	ay to Thursday.)
 ¹ Physicians may order chromoson may order chromosome analysis for geneticist on-call. ² Microarray can only be ordered by a pediatric cardiologist or pediatric 	r sex chromosome diso y a Genetics Profession	rders. For all other c al or developmental	<i>linical indications re</i> pediatrician. 22q11	equire a co .2 targete	onsult with tl d array may	he clinical be ordere	-	Genomics LIS Label

