

## Chromosomal Microarray Prenatal Patient Information

Instructions: The accurate interpretation and reporting of genetic results is contingent upon the reason for referral, clinical information provided, and family history. Supply the information requested below and send paperwork with the specimen or return by fax to

Mayo Clinic Laboratories, Attn: Cytogenetics +1-507-266-5700 or email mclglobal@mayo.		1759. Phone: 507	<b>'-266-5700</b>	/ International clients:	
Patient Information					
Patient Name (Last, First, Middle)		Birth Date (mm-dd-yyyy)		Sex ☐ Male ☐ Female	
Referring Provider Name (Last, First)		Phone		Fax*	
Genetic Counselor Name (Last, First)		Phone		Fax*	
Reason for Testing	*Fax number given mus	t be from a fax machi	ine that compl	ies with applicable HIPAA regulations	
Clinical Information Check all that app	lv.				
Primary Indication for Testing  Advanced maternal age	Cardiac  ☐ Aortic atresia		uloskeletal Acromelia	l	
<ul> <li>☐ Fetal abnormality</li> <li>☐ Abnormal maternal serum screening</li> <li>☐ Abnormal cell-free DNA screening</li> <li>(NIPT, NIPS, cfDNA); increased risk for:</li> </ul>	<ul> <li>□ Atrial septal defect</li> <li>□ Atrioventricular (AV) canal def</li> <li>□ Coarctation of the aorta</li> <li>□ Dextrocardia/situs inversus</li> <li>□ Double outlet right ventricle</li> </ul>	ect	Limb anoi	ot ctures (arthrogryposis) nomaly nelia/micromelia	
Perinatal History  2 vessel cord Donor egg or gestational carrier Hydrops Increased nuchal translucency (includes cystic hygroma) Intrauterine growth restriction (IUGR) Oligohydramnios Polyhydramnios	<ul> <li>□ Ebstein anomaly</li> <li>□ Echogenic intracardiac focus</li> <li>□ Hypoplastic left heart</li> <li>□ Hypoplastic right heart</li> <li>□ Tetralogy of Fallot</li> <li>□ Transposition of the great ves</li> <li>□ Truncus arteriosus</li> <li>□ Ventricular septal defect</li> <li>□ Other:</li> </ul>	sels Genito	Ambiguous genitalia  Hydronephrosis kidney malformation  Megacystis (including posterior valves)  Polycystic kidneys  Renal agenesis  Urethra/ureter obstruction  Other:  Family History  Parents with 2 or more miscarriages  Other relatives with previous  pregnancies with similar clinical history (explain below):		
Other:	Pulmonary  Congenital cystic adenomatoi malformation (CCAM)/small thoracic cavity Diaphragmatic hernia Pleural effusion Pulmonary sequestration Other:  Gastrointestinal Absent stomach Echogenic focus Gastroschisis Meconium ileus/anal atresia Omphalocele Tracheoesophageal fistula Other:	family			
Other:			J UUIGI		

Clinical Descriptions Include any additional relevant clinical information. List karyotype, if known, and provide a report.