

**IriSight™ Comprehensive Analysis - Prenatal**  
 Test Requisition Form  
 Page 1 of 2
Patient Name  
(fetus of)Patient Date  
of BirthAffix barcode label of fetal  
sample here

- ☐ Pregnancy and parent demographics    ☐ Signed informed consents    ☐ **Completed TRF and all clinical notes faxed to 508-302-8022**  
☐ ICD-10 codes    ☐ Ultrasound report(s)    ☐ Clinical & genetic counseling notes with pedigree (please include all family history of known chronic and inherited disease and copies of genetic test results if available)  
☐ Healthcare provider signature

**Test Order**

- ☐ IriSight™ CNV Analysis (RPG002)    ☐ Duo (For MCC)    ☐ Trio  
☐ Opt-In Variants of Uncertain Significance (for trios with clinical indication only)    *No selection will default to an opt-out option. Variants of uncertain significance (VUS) may be reported if correlated with the provided clinical symptoms of the fetus, the pregnancy and/or the family history. Additionally, VUS will only be reported for trios. For more information see test information.*  
☐ Opt-In Fluorescent *In-Situ* Hybridization (FISH)    *No selection will default to an opt-out option. Rapid FISH includes analysis for the common aneuploidies (13, 18, 21, X and Y). For more information see test information.*  
☐ If the IriSight™ CNV Analysis does not yield in a diagnostic result reflex to IriSight™ Comprehensive Analysis - Prenatal (RPG001)  
*\*Reflex to RPG001 might not be covered by insurance.*

- ☐ IriSight™ Comprehensive Analysis - Prenatal (RPG001)    ☐ Duo (For MCC)    ☐ Trio  
☐ Opt-In ACMG Secondary Findings    *No selection will default to an opt-out option and findings in this category will not be returned. For more information see test information.*  
☐ Opt-In Variants of Uncertain Significance (for trios with clinical indication only)    *No selection will default to an opt-out option. Variants of uncertain significance (VUS) may be reported if correlated with the provided clinical symptoms of the fetus, the pregnancy and/or the family history. Additionally, VUS will only be reported for trios. For more information see test information.*  
☐ Opt-In Fluorescent *In-Situ* Hybridization (FISH)    *No selection will default to an opt-out option. Rapid FISH includes analysis for the common aneuploidies (13, 18, 21, X and Y). For more information see test information.*

**Ordering Healthcare Provider**

First Name	Last Name	Title	NPI #
Facility Name			Phone
Facility Address			Fax
City	State	Zip Code	Country
			Email

**Additional Report Recipients**

Full Name	Phone	Fax	Email
Full Name	Phone	Fax	Email

**Billing Information**

ICD-10 Code(s)* <small>*ICD-10 codes must be specified here and/or in attached clinical notes</small>		Suspected Diagnosis	
<input type="radio"/> Insurance Billing			
Insurance Company		Policy #	Group #
Policy Holder First Name		Policy Holder Last Name	Policy Holder DOB
Who is the Policy Holder? <input type="radio"/> Patient <input type="radio"/> Spouse <input type="radio"/> Parent	Policy Holder's Employer		Employer's Address
<input type="radio"/> Institutional Billing		<input type="radio"/> Patient Payment <i>Who should be contacted for billing purposes?</i>	
An invoice will be sent to institution listed above. Contact us for alternate billing. Billing Contact Name: Billing Contact Email:		Payer Name: Payer Email:	Payer Phone: <i>An invoice will be sent to the patient email provided. Insurance will not be billed.</i>

**Healthcare Provider's Statement**

By my signature below, I attest that I am the referring physician, an authorized healthcare provider for the patient, or procurator thereof and this testing is medically necessary for diagnosis and/or treatment of the fetus. I attest that the guardian(s) have been appropriately consented to the test including possible results and outcomes, including variants of uncertain clinical significance (VUSs) and ACMG secondary findings, if selected, and have been given the opportunity to ask questions about the testing and/or seek genetic counseling. I agree to allow an independent genetic counselor facilitated through a third party to provide pre-test and/or post-test genetic counseling, if required by the insurer and/or referring institution. I attest that the guardian(s) have voluntarily consented to testing performed by Variantyx for diagnostic purposes through both oral and written consent.

Healthcare provider signature

Date

**IriSight™ Comprehensive  
Analysis - Prenatal  
Test Requisition Form**  
Page 2 of 2

Patient Name  
(fetus of)

Patient Date  
of Birth

Affix barcode label of fetal  
sample here

**Pregnant Patient Information**

First Name

Last Name

MI

DOB

Address

ID/MR#

City

State

Country

Zip Code

Phone

Email

Sample Type

☐ Blood ☐ Other:

Sample Collection Date

Sample will be collected

☐ In-Clinic ☐ By Variantyx ☐ Patient Given Kit

**Biological Maternal Information (if egg donor or gestational carrier)**

First Name

Last Name

MI

DOB

Sample Type

☐ Blood ☐ Other:

Sample Collection Date

Sample will be collected

☐ In-Clinic ☐ By Variantyx ☐ Patient Given Kit

**Biological Paternal Information**

First Name

Last Name

MI

DOB

Sample Type

☐ Blood ☐ Other:

Sample Collection Date

Sample will be collected

☐ In-Clinic ☐ By Variantyx ☐ Patient Given Kit

**Pregnancy History - Please attach detailed clinical notes (with pedigree if available)**

G \_\_\_\_\_ P \_\_\_\_\_ A \_\_\_\_\_

Sample collection Date\*

Sample Type

☐ Direct amniotic fluid

☐ Direct chorionic villus sampling (CVS)

☐ Fetal/cord blood

☐ Cultured cells; Source \_\_\_\_\_

☐ Genomic DNA; Source \_\_\_\_\_

Gestational Age at Collection

W \_\_\_\_\_

D \_\_\_\_\_

Expected Delivery Date

Predicted Fetal Sex

☐ Male

☐ Female

☐ Unknown

☐ Ambiguous

Multiple Pregnancy?

☐ Twin

☐ Triplet

☐ Other \_\_\_\_\_

Sperm Donation?

☐ Yes

☐ No

Egg Donation?

☐ Yes

☐ No

Abnormal Findings in Previous Pregnancies

Testing Previously Performed in Current Pregnancy

☐ FISH

☐ Karyotype

☐ Microarray

☐ Maternal Serum Screening

☐ Non-Invasive Prenatal Screening

☐ Other \_\_\_\_\_

Results:

\*Note that the collection date is not the same as the date of service for test billing purposes. For more information, please see CMS guidelines.



## IriSight™ Comprehensive Analysis - Prenatal Supplement Form

Patient Name  
(fetus of)

Patient Date  
of Birth

Affix barcode label of fetal  
sample here

### Fetus Phenotypes

Phenotype			Phenotype		
Growth	<input type="radio"/>	Cystic hygroma	Cardiac	<input type="radio"/>	Aortic valve atresia
	<input type="radio"/>	Hydrops fetalis		<input type="radio"/>	Atrial septal defect (ASD)
	<input type="radio"/>	Increased nuchal translucency		<input type="radio"/>	Atrioventricular canal defect
	<input type="radio"/>	Intrauterine growth restriction (IUGR)		<input type="radio"/>	Coarctation of aorta
	<input type="radio"/>	Large for gestational age (Macrosomia)		<input type="radio"/>	Dextrocardia
	<input type="radio"/>	Oligohydramnios		<input type="radio"/>	Ebstein anomaly
	<input type="radio"/>	Placental abnormality		<input type="radio"/>	Echogenic intracardiac focus
	<input type="radio"/>	Polyhydramnios		<input type="radio"/>	Hypoplastic left heart
	<input type="radio"/>	Single umbilical artery		<input type="radio"/>	Hypoplastic right heart
	<input type="radio"/>	Other:		<input type="radio"/>	Pericardial effusion
Neurological	<input type="radio"/>	Abnormal posterior cranial fossa morphology	<input type="radio"/>	Pulmonary valve atresia	
	<input type="radio"/>	Anencephaly	<input type="radio"/>	Tetralogy of Fallot	
	<input type="radio"/>	Aplasia/hypoplasia of the corpus callosum	<input type="radio"/>	Transposition of the great arteries	
	<input type="radio"/>	Cerebellar hypoplasia	<input type="radio"/>	Truncus arteriosus	
	<input type="radio"/>	Choroid plexus cysts	<input type="radio"/>	Ventricular septal defect (VSD)	
	<input type="radio"/>	Dandy-Walker malformation	<input type="radio"/>	Other:	
	<input type="radio"/>	Decreased fetal movement	Genitourinary	<input type="radio"/>	Ambiguous genitalia
	<input type="radio"/>	Holoprosencephaly		<input type="radio"/>	Fetal pyelectasis
	<input type="radio"/>	Hydrocephalus		<input type="radio"/>	Hydronephrosis
	<input type="radio"/>	Lissencephaly		<input type="radio"/>	Hypogonadism
<input type="radio"/>	Spina bifida	<input type="radio"/>		Hypospadias	
<input type="radio"/>	Ventriculomegaly	<input type="radio"/>		Megacystis	
<input type="radio"/>	Other:	<input type="radio"/>		Polycystic kidneys	
<input type="radio"/>		<input type="radio"/>		Renal agenesis	
<input type="radio"/>		<input type="radio"/>		Urethral obstruction	
<input type="radio"/>		<input type="radio"/>		Other:	
Craniofacial	<input type="radio"/>	Cleft lip	Musculoskeletal	<input type="radio"/>	Abnormal vertebral morphology
	<input type="radio"/>	Cleft palate		<input type="radio"/>	Abnormality of the lower limb
	<input type="radio"/>	Hypertelorism		<input type="radio"/>	Abnormality of the upper limb
	<input type="radio"/>	Hypotelorism		<input type="radio"/>	Arthrogryposis multiplex congenita
	<input type="radio"/>	Macrocephaly		<input type="radio"/>	Contractures
	<input type="radio"/>	Microcephaly		<input type="radio"/>	Clubfoot
	<input type="radio"/>	Micrognathia		<input type="radio"/>	Polydactyly
<input type="radio"/>	Pierre-Robin sequence	<input type="radio"/>		Rocker bottom foot	
<input type="radio"/>	Other:	<input type="radio"/>		Scoliosis	
<input type="radio"/>		<input type="radio"/>		Short long bone	
Pulmonary	<input type="radio"/>	Abnormality of the thoracic cavity	<input type="radio"/>	Skeletal dysplasia	
	<input type="radio"/>	Congenital cystic adenomatoid malformation of lung (CCAM)	<input type="radio"/>	Syndactyly	
	<input type="radio"/>	Congenital diaphragmatic hernia	<input type="radio"/>	Other:	
	<input type="radio"/>	Diaphragmatic eventration			
	<input type="radio"/>	Pleural effusion			
<input type="radio"/>	Other:				
Gastrointestinal	<input type="radio"/>	Abnormal stomach morphology			
	<input type="radio"/>	Choanal atresia			
	<input type="radio"/>	Duodenal atresia			
	<input type="radio"/>	Echogenic fetal bowel			
	<input type="radio"/>	Gastroschisis			
	<input type="radio"/>	Omphalocele			
	<input type="radio"/>	Tracheoesophageal fistula			
<input type="radio"/>	Other:				
Other					

