

SPECIMEN INFORMATION

Specimen: Blood DNA Other _____ **Date Collected:** (mm/dd/yyyy) ____ / ____ / ____
 Cord Blood* CVS* Amnio*
 DNA* derived from: _____ Space for Lab Use Only
 (Choose One) Cord Blood CVS Amnio

**Maternal specimen is required to perform Maternal Cell Contamination testing. For MCC sample, complete only the first page of the requisition form to submit with specimen. Note an additional \$950 charge will be added for duplicate analysis, MCC studies, internal control tests and rush processing.*

PATIENT INFORMATION

First name: _____ **MI:** _____ **Institution:** _____
Last name: _____ **Medical Record Number:** _____
Date of Birth: (mm/dd/yyyy) ____ / ____ / ____ **Is the patient adopted?** No Yes
Gender: Male Female Unknown/Unspecified **Is the patient deceased?** No Yes, date: _____
Is patient pregnant? No Yes **EDD:** _____ **Race and Ethnicity: Please check ALL that apply**
 Address: _____ White Ashkenazi Jewish Asian
 City: _____ State: _____ Zip Code: _____ Hispanic Black/African American
 Phone: _____ Native Hawaiian or other Pacific Islander
 Email: _____ American Indian/Native Alaskan Other _____

REFERRING PROVIDER INFORMATION

Referring Provider	Genetic Counselor / Additional Contacts
Name (First, Last): _____	Name (First, Last): _____
Phone: _____ Fax: _____	Phone: _____ Fax: _____
Email: _____	Email: _____
Institution: _____	Institution: <input type="checkbox"/> Same as Referring Provider <input type="checkbox"/> Provided below
Address: _____	_____
_____	_____
City: _____ State: _____	Place facility sticker here
Zip Code: _____ Country: _____	_____

PAYMENT INFORMATION

Please note: Payment information must be completed for testing to begin.

<input type="checkbox"/> Patient Pay (please complete section in its entirety)** <input type="checkbox"/> Check (please attach to forms)* <small>*Please make checks payable to Partners Personalized Medicine*</small> <input type="checkbox"/> Credit card (please fill out credit card information in its entirety) Card type: <input type="checkbox"/> Mastercard <input type="checkbox"/> Visa <input type="checkbox"/> AMEX Name (as it appears on card): _____ Credit card number: _____ Expiration Date: _____ 3 Digit Security Code: _____ <small>**For patient pay, please provide billing address and contact information. If same as above, please note section as such.**</small> Patient Pay Billing Address: _____ City: _____ State: _____ Zip Code: _____ Country: _____ Home: _____ Cell/Work: _____ Email: _____	<input type="checkbox"/> Referring Institution (please complete section in its entirety) <small>*For new referring facilities, please complete and submit the New Institution Add Form*</small> Bill to Name/Department: _____ Address: _____ _____ City: _____ State: _____ Zip Code: _____ Country: _____ Phone: _____ Contact Person: _____
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CONGENITAL HEART DISEASE (CHD) REQUISITION FORM

Patient Name: _____ Date of Birth: ____/____/____ (MM/DD/YYYY)

SEQUENCING PANEL AND SINGLE GENE TESTS

Please check box(es) to order. If ordering reflex testing, please indicated order (i.e. 1, 2, 3). Testing can be conducted concurrently if desired.

Isolated Nonsyndromic Congenital Heart Disease (Septal and Cyanotic Defects with or without Conduction System Disease)

- Congenital Heart Disease Panel A (*GATA4, NKX2-5 and JAG1*) \$1,300
- GATA4* Gene Sequencing \$750
- NKX2-5* Gene Sequencing \$600
- JAG1* Gene Sequencing \$1,100

Isolated Supravalvular Aortic Stenosis

- ELN* Gene Sequencing (*Not to be ordered in place of 7q11 deletion testing for Williams syndrome*) \$1,300

Familial Variant Testing

- Familial Variant(s)** OR Research Confirmation** \$500

If proband testing was performed elsewhere, please attach a copy of the original result and send positive control sample, if available

Gene _____ Variant _____
 Proband Name _____
 Relationship to Patient _____ LMM Accession #: PM- _____

CLINICAL INFORMATION

Clinical Status: Affected Unknown (no cardiac imaging/screening) Unaffected (all cardiac imaging/screening normal)

ICD-10 Code(s): _____

Septal Defects:

- ASD (Atrial Septal Defect)
- VSD (Ventricular Septal Defect)
- AVSD (Atrioventricular Septal Defect)
- Other _____

Cyanotic Defects:

- TOF (Tetralogy of Fallot)
- TOF/PA (TOF w/ Pulmonary Atresia)
- PDA (Patent Ductus Arteriosus)
- TGAV (Transp. of Great Arteries/Vessels)
- Other _____

Conduction Defects:

- Arrhythmia
- AV Block (Atrioventricular Block)
- Other _____

Obstructive Defects:

- SVAS (Supravalvular Aortic Stenosis)
- Other _____

Other Features:

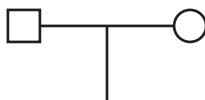
- Cutis Laxa
- _____

Previous testing for genetic/syndromic causes: Yes No

- | | | | |
|---------------------------|---|-------------------|---|
| Chromosome Abnormality | <input type="checkbox"/> nl <input type="checkbox"/> abnl <input type="checkbox"/> not done | Williams Syndrome | <input type="checkbox"/> nl <input type="checkbox"/> abnl <input type="checkbox"/> not done |
| 22q11/10p13 Deletion | <input type="checkbox"/> nl <input type="checkbox"/> abnl <input type="checkbox"/> not done | Alagille Syndrome | <input type="checkbox"/> nl <input type="checkbox"/> abnl <input type="checkbox"/> not done |
| Noonan Spectrum Disorders | <input type="checkbox"/> nl <input type="checkbox"/> abnl <input type="checkbox"/> not done | Other _____ | <input type="checkbox"/> nl <input type="checkbox"/> abnl |

FAMILY HISTORY

Family History: Yes No *If yes, please list affected family members and features or sketch below* _____



Paternal Ancestry: _____
 Maternal Ancestry: _____
 Consanguinity: Yes No

○ = Female □ = Male ◇ = Gender Unspecified
 ● ■ ◆ = Affected Individual ⊙ = Carrier