

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.

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

RASOPATHY REQUISITION FORM

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

TEST TO BE PERFORMED

Please check box(es) to order.

The panel test is performed via next-generation sequencing (NGS). CNV analysis is included when NGS data meets necessary quality standards.

Expanded RASopathy Panel (14 Genes)

Familial Variant Testing (Sequencing - first 3 variants, \$500; each additional variant, \$50.)

Familial Variant(s) OR Research Confirmation - If proband testing was performed elsewhere, please attach a copy of the original result and send positive control sample, if available.

Gene _____ Variant _____ LMM Accession #: PM- _____

Proband Name _____ Relationship to Patient _____

CLINICAL INFORMATION

Clinical status: Affected Unknown Unaffected

Purpose of study: Diagnostic Family history Prenatal Other: _____

Clinical diagnosis: Noonan LEOPARD CFC Costello NF1 Legius

(check all known/suspected clinical diagnoses)

Age at diagnosis: _____

ICD-10 Code(s): Q87.5 (Other congenital malformation syndromes with other skeletal changes)
 Q87.89 (Other specified congenital malformation syndromes, not elsewhere classified)
 Q879.8 (Other specified congenital malformation)
 Other _____

Ultrasound Finding: Cystic hygroma Increased NT - Size: _____ None
 Heart defect - Type: _____ Other _____

Congenital heart defect: Pulmonic valve stenosis Hypertrophic cardiomyopathy None
 Septal defect Other _____

Facial dysmorphism: Epicanthal folds Ptosis of the eyelids Low nasal bridge Macrocephaly
 Hypertelorism Downward eye slant Low set ears and posteriorly rotated
 Papillomas Coarseness None

Short stature: Yes - Height (%): _____ Parental Heights: _____ No

Neurodevelopment: Learning disabilities Developmental delay Intellectual Disability ADHD
 Seizures Normal

Skeletal: Pectus excavatum Pectus carinatum Scoliosis Normal

Genitourinary: Cryptorchidism (undescended testes) Normal
 Kidney malformation If yes, please describe: _____

Hair/Skin/Eye findings: Loose anagen hair Lentigines Café-au-lait spots (#): _____ Lisch nodules (#): _____
 Wide-spaced nipples Webbed neck Freckling ____ axillary ____ inguinal
 Lipomas Other _____ None

Tumor Growth: Neurofibromas (#): _____ Plexiform neurofibromas (#): _____ Optic glioma
 Osseous lesion - Type: _____ Other: _____

Malignancy: Yes No If yes, please describe: _____

Bleeding diathesis: Yes No If yes, please describe: _____

Vasculopathy: Yes No If yes, please describe: _____

Additional Features: _____

Previous Genetic Testing: Yes No Gene(s)/Tests: _____

Result (if variants detected, please elaborate): _____

PLEASE PROCEED TO PAGE 4 TO COMPLETE THE FAMILY HISTORY

RASOPATHY REQUISITION FORM

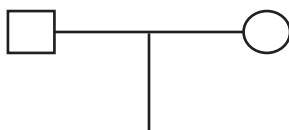
Patient Name: _____ Date of Birth: ____/____/____ (MM/DD/YYYY)

FAMILY HISTORY

Family History: Yes No (Sketch below or attach pedigree, if appropriate)

Has another family member already had genetic testing for this disease? Yes No

If yes, please describe and attach a copy of the genetic test lab report and pedigree. _____



Paternal Ancestry: _____

Maternal Ancestry: _____

Consanguinity: Yes No

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