



**GENETICS REQUISITION FORM (CYP2C19, CF, MCC, SMA)**

PATIENT INFORMATION (Please Print):				REFERRING PHYSICIAN INFORMATION (Please Print):			
First Name:		MI:	Last Name		Physician:		
Date of birth: (MM/DD/YYYY)		Sex: <input type="checkbox"/> M <input type="checkbox"/> F		MRN #:		Address:	
Address			City:		City:		State:
State:		Zip Code:		Phone:		Phone:	
						Fax:	
Ancestry (check all that apply): <input type="checkbox"/> White/Caucasian <input type="checkbox"/> Black/African American <input type="checkbox"/> Hispanic <input type="checkbox"/> Asian <input type="checkbox"/> Ashkenazi Jewish <input type="checkbox"/> Eastern/Central Europe <input type="checkbox"/> Western/Northern Europe <input type="checkbox"/> Central/South American <input type="checkbox"/> Middle Eastern <input type="checkbox"/> Native American <input type="checkbox"/> Other: (specify) _____				NPI#:			
				Additional Report To:		Fax:	
				Other Medical Professional:		Fax:	
SPECIMEN INFORMATION				INFORMED CONSENT & STATEMENT OF MEDICAL NECESSITY			
A requisition form <b>MUST</b> accompany each specimen.  Date/Time of sample obtained Date: _____ Time: _____:_____  <input type="checkbox"/> Peripheral Blood in EDTA (5-6 ml in lavender top tube) <input type="checkbox"/> Extracted DNA Source of Extracted DNA <input type="checkbox"/> CVS <input type="checkbox"/> Amniotic Fluid <input type="checkbox"/> Peripheral Blood				I affirm each of the following: I have provided genetic information to the patient and the patient has consented to genetic testing. This test is medically necessary for the diagnosis of a disease or syndrome. The result will be used in the patient's medical management and treatment decisions. The person listed as the Ordering Physician is authorized by law to order the test(s) requested herein.  <b>REQUIRED</b> Signature of Requesting Physician _____			
INDICATION FOR STUDY:							
Reason for Referral :					ICD-10 Code(s):		
<input type="checkbox"/> <b>Diagnosis/Suspected diagnosis</b> List clinical findings: _____ _____ <input type="checkbox"/> <b>Carrier screening</b> <input type="checkbox"/> <b>Positive family history</b> Relationship to patient: _____ Gene variant(s) known in family: _____ <input type="checkbox"/> <b>Abnormal fetal ultrasound</b> List findings: _____ _____ <input type="checkbox"/> <b>Other (list):</b> _____ Comments/ Special Instructions _____							
TEST(S) REQUESTED:							
<input type="checkbox"/> CYP2C19 Genotyping <input type="checkbox"/> Cystic fibrosis (CFTR) 144 variant panel <input type="checkbox"/> Cystic fibrosis (CFTR) sequencing <input type="checkbox"/> Cystic fibrosis (CFTR) targeted variant analysis Variant(s) to be analyzed: _____ <input type="checkbox"/> Maternal cell contamination (MCC) <input type="checkbox"/> Spinal Muscular Atrophy (SMA) (Carrier screen - SMN1 dosage) <input type="checkbox"/> Spinal Muscular Atrophy (SMA) (Diagnostic test - SMN1 dosage & SMN2 dosage)							