

THIS IS NOT A TEST REQUEST FORM.

Please fill out this form and submit it with the test request form or electronic packing list.

PATIENT HISTORY FOR CYSTIC FIBROSIS (CF), FRAGILE X SYNDROME (FXS), AND SPINAL MUSCULAR ATROPHY (SMA) CARRIER SCREENING

Patient Name _____ Date of Birth _____ Sex F M
 Physician _____ Physician Phone _____
 Practice Specialty _____ Physician Fax _____
 Genetic Counselor _____ Counselor Phone _____

Patient's Ethnicity (check all that apply)

- African-American Asian Hispanic Native American
 Ashkenazi Jewish Caucasian Middle Eastern Other: _____

Reason for Referral:

- Routine carrier screening; healthy patient with no family history of cystic fibrosis (CF), fragile X syndrome (FXS), or spinal muscular atrophy (SMA)
- Family history; healthy patient with a family history of CF, FXS, or SMA. Specify which condition: _____
 The relative is: healthy affected Relationship of affected relative to patient: _____
 Has DNA testing been performed for the family member(s)? No Yes Unknown
 If yes, attach a copy of the relative's DNA laboratory result. (REQUIRED for familial mutation testing)
 or describe the test results: _____
- Known carrier; patient is a known healthy carrier of CF, FXS, and/or SMA (has had previous testing):
 If yes, which condition? _____ Test result of patient: _____
- Other: (e.g., abnormal ultrasound finding). Please describe: _____

Is reproductive partner symptomatic or a known healthy carrier of CF, FXS, or SMA? Yes No NA

If yes: which condition? _____ Partner is: a healthy carrier affected/symptomatic
 Has DNA testing been performed for the partner? No Yes Unknown
 If yes, describe the test results: _____

Does reproductive partner have a family history of CF, FXS, or SMA? Yes No NA

Relationship of affected relative to partner: _____
 If yes: which condition? _____ Relative is: a healthy carrier affected/symptomatic
 Has DNA testing been performed for the partner? No Yes Unknown
 If yes, describe the test results: _____

Check the test you intend to order.

Panel that includes testing for carrier status for 3 disorders—CF, FXS, and SMA:

- 3000258 Genetic Carrier Screen, (CF, FXS, and SMA) with Reflex to Methylation:** Carrier screening for cystic fibrosis (CF), fragile X syndrome (FXS), and spinal muscular atrophy (SMA).

Panel components available as individual tests:

- 2013661 Cystic Fibrosis (CFTR) 165 Pathogenic Variants:** Carrier screening for cystic fibrosis (CF)
 2009033 Fragile X (FMR1) with Reflex to Methylation Analysis: Carrier screening for fragile X syndrome (FXS)
 2013436 Spinal Muscular Atrophy (SMA) Copy Number Analysis: Carrier screening for spinal muscular atrophy (SMA)

Targeted testing for known CF mutation (laboratory report from family member REQUIRED)

- 2001961 Familial Mutation, Targeted Sequencing.** Tests for a mutation previously identified in a family member.

Additional carrier screening options (Ashkenazi Jewish screening, hemoglobinopathy screening, or expanded carrier screening) are available. See www.aruplab.com/genetics for additional options.

Master Label

For questions, contact an ARUP genetic counselor at (800) 242-2787, ext. 2141