St. John Health System
Lab Catalog

Inheritest(R) CF/SMA Panel

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Order Name: CF/SMA WB Panel

Room Temperature

Test Number: 5194938
Revision Date: 03/21/2023

TEST NAME		METH	ODOLOGY	LOINC CODE				
Inheritest(R) CF/SMA Panel			See Test Notes					
SPECIMEN REQUIREM	MENTS							
Specimen	Specimen Volume (min)	Specimen Type	Specimen Container	Transport Environment				
Preferred	8.5 mL (3 mL)	Whole Blood	ACD Solution A or B (Yellow Top)	Room Temperature				
Alternate 1	8.5 mL (3 mL)	Whole Blood	EDTA (Lavender Top)	Room Temperature				
Alternate 2	1	Saliva	Oragene Dx saliva kit	Room Temperature				

Instructions Specimen Type: Whole blood or PurFlock buccal swab kit or Oragene Dx saliva kit

Specimen Volume: 8.5 mL whole blood or PurFlock buccal swab kit or Oragene Dx saliva kit **Mininum Volume:** 3 mL whole blood or PurFlock buccal swab kit or Oragene Dx saliva kit

Buccal swab

Collection: Standard phlebotomy. Follow PurFlock buccal swab kit or Oragene Dx 500 saliva kit collection instructions. Do not eat, drink, smoke, or

PurFlock buccal swab kit

chew gum 30 min prior to collection.

Specimen Storage: Maintain specimen at room temperature or refrigerate at 4C Do not freeze.

Special Instructions: In cases in which there is a known variant documented in the family, the physician may prefer to order Targeted Variant

Analysis, test code 482552. Test orders must include an attestation that the provider has the patient's informed consent for genetic

GENERAL INFORMATION

Alternate 3

Expected TAT 1	4 - 21	days	In some cases,	additional time m	nay be re	equired for c	onfirmatory or	reflex tests.
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Notes Clinical Questionnaire for Inheritest® Carrier Screen and GeneSeq® PLUS

Methodology

Cystic fibrosis: Next-generation sequencing to identify genetic variants, including small nucleotide variants (SNVs), insertions, deletions and copy number variants (CNVs). Spinal muscular atrophy (SMA): Copy number assessment of SMN1 exon 7 by quantitative polymerase chain reaction (qPCR). For carrier screening, when two copies of SMN1 are detected, allelic discrimination qPCR targeting c.*3+80T>G in SMN1 is performed. The presence or absence of c.*3+80T>G correlates with an increased or decreased risk, respectively, of being a silent carrier (2+0).

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