California Prenatal Screening (PNS) Program

Additional Information for Providers

About California State-Contracted cfDNA Laboratories

NATERA LABORATORY

1. **Methodology for cfDNA analysis?**
   Single-nucleotide polymorphism-based, or SNP-based, cfDNA screening reveals a baby’s risk for severe genetic disorders. The test uses a unique SNP-based technology to analyze fetal/placental DNA obtained through a blood draw from the mother. This test can differentiate between maternal and fetal DNA to assess the risk of aneuploidies. The test also screens twin pregnancies for zygosity and fetal sex of each baby.

2. **Can I have additional tests performed on the specimen I send for the Prenatal Screening Program screening through Natera?**
   a. Yes. Additional tests can be ordered using the same specimen. These are the following:
      i. Panorama CA Supplemental Panel: includes sex chromosome aneuploidies, Triploidy. Optional add-ons: 22q11.2 deletion syndrome, extended microdeletion panel
      ii. Horizon Carrier Screen:
         1. Single gene options: CF, SMA, DMD
         2. Panel options: Horizon 4, Horizon 14, Horizon 27, Horizon 106, Horizon 274

3. **How do patients consent to tests additional to the PNS Program screening panel?**
   a. The physician must sign a separate requisition form indicating that they have consented the patient for additional testing. Patient consent language is on the requisition form.
   b. Visit [www.natera.com/CAPNS](http://www.natera.com/CAPNS) to download the requisition form or call 650-446-4095.

4. **Are you CLIA-certified?** Yes - CLIA# 05D1082992
   **Are you certified by California Laboratory Field Services?** Yes - CDF# 00337104

5. **Where is the location of your lab(s) where my patients’ specimens will be processed in California?**
   The performing lab for Natera is located at:
   Natera, Inc.,
   201 Industrial Rd.
   San Carlos, CA 94070

6. **Disclaimer:**
   This test has been developed and its performance characteristics determined by the CLIA-certified laboratory performing the test. The test has not been cleared or approved by the US Food and Drug Administration (FDA). CAP accredited, ISO 13485 certified, and CLIA certified. © 2022 Natera, Inc. All Rights Reserved.