

Napa-Solano-Yolo-Marin-Mendocino County Public Health Laboratory

2201 Courage Drive, MS 9-200

Fairfield, California 94533

(707) 784-4410 FAX (707) 423-1979, Email: ngha.permits@solanocounty.com

Beatrix Kapusinszky, PhD, PHLD (ABB), Laboratory Director

**NONDIAGNOSTIC GENERAL HEALTH ASSESSMENT (NGHA)
ANNUAL REGISTRATION APPLICATION**

This registration form must be completed and received by the Napa-Solano-Yolo-Marin-Mendocino County Public Health Laboratory *at least 30 days* prior to operation of a program of nondiagnostic general health assessment (NGHA).

PART 1: ADMINISTRATION

A. Name of Organization or Operator: _____

Permanent Address: _____

_____ City _____ Zip Code

Business Phone: () _____ Fax: () _____

CLIA #: _____ Exp.: _____

B. Name of Owner: _____

Address (if different than above): _____

_____ City _____ Zip Code

Business Phone: () _____ Fax: () _____

C. Supervisory Committee Members:

Name of Physician: _____

Address: _____

_____ City _____ Zip Code

Business Phone: () _____ Fax: () _____

CA Medical License #: _____ Exp.: _____

Name of Clinical Laboratory Scientist: _____

Address: _____

_____ City _____ Zip Code

Business Phone: () _____ Fax: () _____

CA Clinical Laboratory Scientist License #: _____ Exp.: _____

D. Record Storage:

All operators must have a permanent address where records of testing and protocols shall be stored and NSYMM County Public Health Laboratory must be notified in writing within 30 days of any change in record storage location.

Record Storage Address: _____

_____ City _____ Zip Code

Business Phone: () _____ Fax: () _____

PART 2: COMPLIANCE

A. This assessment program must be operated per Section 1244 of the California Business and Professions Code. Please answer each of the following questions. To comply with current California law, you must be able to answer yes to all questions **and supporting documentation must be submitted with this application.**

YES NO

- This program will be a nondiagnostic health assessment program (NGHA), whose purpose will be to refer individuals to licensed sources of care as indicated.
- This program will utilize only those devices, which comply with all of the following:
 - A. Meet applicable state and federal performance standards pursuant to Section 26605 of the Health and Safety Code.
 - B. Are not adulterated as specified in Article 2 (commencing with Section 26610) of Chapter 6 of Division 21 of the Health and Safety Code.
 - C. Are not misbranded as specified in Article 3 (commencing with Section 26630) of Chapter 6 of Division 21 of the Health and Safety Code.
 - D. Are not new devices unless they meet the requirements of Section 26670 of the Health and Safety Code.
- This program maintains a supervisory committee consisting of at a minimum, a California licensed physician and surgeon and a Laboratory Clinical Scientist licensed pursuant to the California Business and Professions Code.
- The supervisory committee for the program has adopted written protocols, which shall be followed in the program. (Include a copy of your written protocols with this application.)
- The protocols contain provisions of written information to individuals to be assessed. (Include a copy of all written information that will be provided to individuals as part of this program.)
- Written information to individuals includes the potential risks and benefits of assessment procedures to be performed in the program.
- Written information includes the limitations, including the nondiagnostic nature, of assessment examinations of biological specimens performed in the program.
- Written information includes information regarding the risk factors or markers targeted by the program.
- Written information includes the need for follow up with licensed sources of care for confirmation, diagnosis, and treatment as appropriate.
- Written protocols contain the proper use of each devices utilized in the program. Protocols must include the operation of analyzers, maintenance of equipment and supplies, and performance of quality control procedures including the determination of both accuracy and reproducibility of measurements in accordance with instructions provided by the manufacturer of the assessment device used.
- Written protocols contain the proper procedures to be employed when drawing blood, if blood specimens are to be obtained.
- Written protocols contain procedures to be employed in handling and disposing of all biological specimens to be obtained and material contaminated by biological specimens.
- Written protocols contain proper procedures to be employed in response to fainting, excessive bleeding, or other medical emergencies.
- Written protocols contain procedures for reporting of assessment results to the individual being assessed (please attach a copy of your report form).
- Written protocols contain procedures for referral and follow up to licensed sources of care as indicated.
- The written protocols adopted by the supervisory committee shall be maintained for at least one year following completion of the assessment program during which period, they shall be subject to review by the county health officer or designee.

B. If a skin puncture to obtain a blood specimen is to be performed:

YES NO

[] [] The individual(s) performing skin punctures shall be authorized to do via (a) their professional scope of practice or (b) meet California phlebotomy regulations as identified in the California Business and Professions Code, Sections 1242.5, 1246, and 1282.2; California Code of Regulations, Title 17, Sections 1029.31–1029.35, 1031.4, 1031.5, and 1034; and Health and Safety Code, Section 120580 and possess a current phlebotomy license issued by the CA Dept. of Public Health, Laboratory Field Services Program. (Documentation of professional license(s) must be submitted with each event application.)

[] [] It is understood that “skin puncture” as related to this program means the collection of a blood specimen by the finger stick method only and does not include venipuncture, arterial puncture, or any other procedure for obtaining a blood specimens.

PART 3: FEES

- Annual registration fee: \$100
- Event permit, per site: \$50
- Consultation (per hour): \$150

Make Checks Payable To: NSYMM Public Health Laboratory
Return Application To: Napa-Solano-Yolo-Marin-Mendocino County Public Health Laboratory
 NGHA Program
 2201 Courage Drive, MS 9-200
 Fairfield, CA 94533

PART 4: PERMIT APPLICATION AND POSTING

An event permit application must be submitted for each event and each location. The event permit is valid for a 7-day timeframe, starting from the first day the event will be held. The permit for the specific event location address must be posted during operation of a nondiagnostic general health assessment program.

PART 5: CONTACT INFORMATION AND SIGNATURE

Name of Person Requesting Registration: _____

Address: _____

_____ City _____ Zip Code _____
 Business Phone: () _____ Fax: () _____

I certify that the above information is accurate and complete and that I am aware of the laws and regulations that apply to nondiagnostic testing in the State of California and in the County of Solano, Napa, Yolo, Marin, or Mendocino in which testing is to be performed.

 Applicant’s Signature Date of Application

FOR OFFICIAL USE ONLY

Reviewed by: _____ Date: _____

Date Issued: _____ Expiration Date: _____

Fees Received / Credit applied*: _____ Date Received: _____

*Credits may be used for up to 12 months after issue