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LIMITED SERVICE LABORATORY REGISTRATION NOTIFICATION TO ADD PERMANENT TESTING LOCATION TO A MULTI-SITE NETWORK APPLICATION INSTRUCTIONS

This application is intended for use by not-for-profit and/or government Limited Service Laboratories in order to take advantage of a multisite Limited Service Laboratory Registration option, whereby you may link multiple permanent locations performing waived and/or providerperformed microscopy procedures under a single CLIA registration number. Read and follow the instructions carefully since submission of incomplete or incorrect applications will delay processing.

A. BACKGROUND AND GENERAL INFORMATION

The New York State Department of Health's Clinical Laboratory Evaluation Program has been authorized under Section 579 of Article 5, Title V of the Public Health Law to provide oversight to non-physician office laboratories performing waived and/or provider-performed microscopy procedures in New York State. These facilities are considered Limited Service Laboratories and must register with the Department as described in this registration package in order to obtain a federal CLIA number and authorization to perform patient testing.

B. HOW TO DETERMINE IF YOUR FACILITY QUALIFIES FOR THE MULTI-SITE REGISTRATION

Not-for-profit, state or local government laboratories or programs engaged in limited public health testing not exceeding fifteen types of tests per registration may be eligible to apply for a multi-site CLIA number, sharing a common director and CLIA number. Only one application and reapplication fee would be required for all the laboratories sharing the common CLIA number. Use this form to add a secondary location to a new or existing multi-site network. All sites in the multi-site network must be operated by the same non-for-profit corporation or government entity.

Please note that the authority for the New York State Department of Health, Wadsworth Center, Clinical Laboratory Evaluation Program to request personal information from you, including identifying numbers such as federal Employer Identification Number (EIN), and the authority to maintain such information, is found in Section 5 of the New York State Tax Law. <u>Disclosure of this information by you is mandatory</u>. These numbers are routinely used only as identifiers within our Program. They may only be released for tax administration purposes and other purposes authorized by the Tax Law. The Administrator of the Clinical Laboratory Evaluation Program is responsible for maintaining the records of such information. The administrator can be reached by writing to the Clinical Laboratory Evaluation Program at the address indicated at the top of this page.

SECTION 1 - PRIMARY LABORATORY INFORMATION. Information provided is to be that of the laboratory designated as the Primary Limited Service Laboratory registrant under the CLIA & PFI Numbers referenced.

- Primary Laboratory Name: Indicate the legal name and address of the Primary Limited Service Laboratory registrant.
- Primary CLIA & PFI Numbers: If the Primary Limited Service Laboratory registrant has already been issued CLIA & PFI
 Numbers, please indicate them in the areas provided in this section. If <u>no</u> numbers have been issued previously, they will be
 assigned upon the submission of a Limited Service Laboratory Registration Application, form DOH-4081.
- Primary Laboratory Telephone & Fax Numbers, and E-mail Address: These sections are self-explanatory.

SECTION 2 – ADDITIONAL TESTING SITE INFORMATION. Information provided in this section should be that of the NEW permanent testing location to be added to the Primary Limited Service Laboratory Registration (CLIA Number) referenced in Section 1–Primary Laboratory Information.

- **Testing Site Name:** Indicate the legal name of the NEW permanent testing site to be covered under the Primary Limited Service Laboratory registration.
- County/Borough: Indicate the New York State county or borough that the NEW permanent testing site is physically located in.
- Testing Site Address: The testing site address must be the actual physical location of the NEW permanent testing site, including floor, suite and/or room, if applicable.
- Testing Site Telephone and Fax Numbers, E-mail Address: Indicate contact information for the new permanent testing site.
- Testing Site Contact Person Name, Telephone Number and E-Mail Address: Indicate contact information for the new permanent testing site.
- Testing Site Days & Hours of Testing: Indicate the days and hours when laboratory testing will be performed at the NEW
 permanent testing site.
- Laboratory Type: Select one from the list below that best describes your laboratory and enter in appropriate area on application:

01-24 Ambulance	14- <i>01</i> Hospital
02-3B Ambulatory Surgery Center	15-11 Independent
03-02 Ancillary Testing Site in Health Care Facility/Hospital	16-12 Industrial* (Include Bureau Lic. Number with application)
Extension Clinic	17-13 Insurance
04-25 Assisted Living Facility	18-14 Intermediate Care Facility for the Mentally Retarded
05- 26 Blood Bank	19- <i>15</i> Mobile Laboratory
06-3A Community Clinic	20-16 Pharmacy
07-04 Comprehensive Outpatient Rehabilitation Facility	21-19 Physician Office
23-06 Correctional Facility	22-20 Practitioner Other
08-3C End Stage Renal Disease Dialysis Facility	24-27 Public Health Laboratory
09-3D Federally Qualified Health Center	25-3D Rural Health Clinic
10-08 Health Fair	26-17 School/Student Health Service
11-07 Health Maintenance Organization	27-18 Skilled Nursing Facility or Nursing Facility
12-08 Home Health Agency	28-28 Tissue Bank/Repositories
13-09 Hospice	29-99 Other* (Specify Laboratory Type)

SECTION 2 – ADDITIONAL TESTING SITE INFORMATION (continued). Information provided in this section should be that of the NEW permanent testing location to be added to the Primary Limited Service Laboratory Registration (CLIA Number) referenced in Section 1–Primary Laboratory Information.

Community Screening: Indicate whether your laboratory or laboratory network will perform community screening events.
 Laboratories seeking approval to operate community screening events must maintain a protocol describing in detail how laboratory testing will be performed.

SECTION 3A – WAIVED TEST PROCEDURES REQUESTED. For each *Waived* test that you wish to perform at the NEW permanent testing site, you must provide the following information:

Indicate the *Waived* test procedure (i.e. blood glucose, dipstick urinalysis, fecal occult blood, etc.) that you wish to perform and provide the combined estimated annual test volume for <u>all</u> *Waived* test procedures indicated. **Waived* testing includes tests performed using a kit, device or procedure, which has been designated by the Food and Drug Administration (FDA) as *Waived* for the purposes of CLIA '88. Non-DOT breath alcohol testing must be performed using an FDA approved IVD Over-The-Counter device. Sites performing these tests shall maintain documentation that the tests in use have been so designated. Listings of waived tests are available at the following websites:

To Search By Test System: www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfCLIA/testswaived.cfm

To Search By Analyte: www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfCLIA/analyteswaived.cfm

To Search a Particular Kit/Mfr.: www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfCLIA/search.cfm

To Search FDA's IVD Over-The-Counter Lab Test Database: www.accessdata.fda.gov/scripts/cdrh/cfdocs/cflVD/Search.cfm

**SPECIAL NOTE (Regarding COVID-19 Testing): You must specify the category that you are requesting: COVID-19 Antigen, COVID-19 Molecular, and/or COVID-19 Antibody. Understand that COVID-19 testing may only be performed using a device approved for use in Limited Service Laboratories. The current list of approved devices is posted on our website under the tab entitled "COVID-19 Response for Limited Service Laboratory Registration Requests and Additions" at: https://www.wadsworth.org/regulatory/clep/limited-service-lab-certs. This list will be revised as new tests are approved.

Additional guidance with protocol development for lead, and/or rapid HIV testing is available at the following websites:

For Lead Testing: www.wadsworth.org/regulatory/clep/limited-service-lab-certs For HIV Testing: www.health.state.ny.us/diseases/aids/testing/rapid/index.htm

SECTION 3B – PROVIDER-PERFORMED MICROSCOPY (PPM) PROCEDURES REQUESTED. For each Provider-performed Microscopy (PPM) Procedure that you wish to perform at the NEW permanent testing site, you must provide the following information: Indicate the Provider-performed Microscopy (PPM) Procedures (i.e. Wet Mounts, KOH Preps, etc.) that you wish to perform and provide the

Indicate the Provider-performed Microscopy (PPM) Procedures (i.e. Wet Mounts, KOH Preps, etc.) that you wish to perform and provide the combined estimated annual test volume for all PPM procedures indicated. *Provider-performed Microscopy (PPM) Procedures includes tests personally performed as part of physical examinations by health care providers, licensed and currently registered in New York State, including physicians, dentists, podiatrists, physician assistants, nurse practitioners and certified midwives operating within the scope of practice for their profession and which have been designated as PPM Procedures by the Centers for Disease Control. Sites performing these tests shall maintain documentation that the tests in use have been so designated.

SECTION 4 - CERTIFICATION

This section must be completed & signed by the Laboratory Director responsible for the technical and clinical direction of laboratory testing at the Primary Limited Service Laboratory under the CLIA & PFI numbers indicated in Section 1–Primary Laboratory Information and the individual completing the application (if different). This individual assumes responsibility as the laboratory director for testing performed at all sites within the network. Please Note: All signatures must be original. SIGNATURE STAMPS WILL NOT BE ACCEPTED.

OUR MAILING ADDRESS

Application documents must be returned to our office at the address below:

Regular Mail

Clinical Laboratory Evaluation Program Wadsworth Center New York State Department of Health Empire State Plaza Albany, NY 12237

Express Mail

Clinical Laboratory Evaluation Program Wadsworth Center New York State Department of Health Empire State Plaza P1 South - Loading Dock J Albany, NY 12237

LIMITED SERVICE LABORATORY MULTI-SITE NETWORK REGISTRATION

Once your application is approved, the Primary Site will be sent registration documents, which will serve to verify your enrollment with this program and will also provide documentation of your CLIA registration number. Registrations will be valid for two years from the date issued. Approximately three months before it expires, the Primary Site will receive an application to renew the registration for the entire network. Multi-site network registrants may only perform the tests listed on the registration certificate issued to the Primary Site.

CHANGES IN STATUS

Once approved, the Primary Site must keep our Program informed of any changes which may affect your registration status (i.e. laboratory name, address, director, test menu, owner, additional testing sites, etc.). Please be advised that Limited Service Laboratory registrations are void upon change in the laboratory location or the owner, and you must inform our Program of any change in location or laboratory director within 30 days of the change. The Limited Service Laboratory Change forms may be downloaded from our website at:

www.wadsworth.org/regulatory/clep/limited-service-lab-certs

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LIMITED SERVICE LABORATORY REGISTRATION NOTIFICATION TO ADD PERMANENT TESTING LOCATION TO A MULTI-SITE NETWORK APPLICATION

FOR OFFICE USE ONLY: I____ PFI:_____SITE:____Gaz Code: ____

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same non-for-profit corporation	or government entity.				,	
Are you adding an additional sit	e to an existing CLIA registrat	ion nur	mber?			
\square Yes (Complete this docum	ent in its entirety)					
☐ No (Complete a <i>Limited Se</i> If this is a new facility, indicate the	ervice Laboratory Registration, for projected opening date:				rvice Laboratory Registration)	
1. PRIMARY LABORATORY INF Limited Service Laboratory regi	ORMATION: This section is to be strant under the CLIA & PFI Number			ety by the laborato	ry designated as the Primary	
Laboratory Name (Limited to 70 Characters):				CLIA Number:		
				PFI Number:		
Laboratory Address (Physical Locatio	n of Laboratory):					
City				State	ZIP Code	
Telephone Number:	FAX Number:		Laboratory E-Ma	il Address:	1	
2. ADDITIONAL TESTING SITE be added to the Multi-Site Limite	INFORMATION: Complete this s d Service Laboratory Registration					
Site Name (Limited to 70 Characters)					County/Borough:	
Site Address (Physical Location of Sit	e):				1	
City:				State:	ZIP Code:	
Telephone Number:	Fax Number:		Site Contact Per	son Name:	1	
Site E-Mail Address:			Telephone Number:			
			E-Mail Address:			
Indicate the Days & Hours when testing	ng will be performed (Please clarify h	hours as	AM and/or PM):			
MOto TU_	to W	/E	to	TH	to	
FRto SA_	to S	SU	to	-		
Indicate Laboratory Type Code From List Located in Instructions			ate whether your la	ether your laboratory or laboratory network will perform community		
(form DOH-4081MSi):		scree	ening events:	☐ Yes	□ No	

WAIVED TEST PROCEDURES REQUEST annual test volume.	ED: Check off all Waived tests that you intend	d to perform and indicate the comb	ined estimated				
Adenovirus	☐ Erythrocyte Sedimentation Rate (ESR)	☐ Nicotine					
☐ Aerobic/Anaerobic Organisms-Vaginal	□ Ethanol	☐ Occult Blood					
☐ Alanine Aminotransferase (ALT)	☐ Glucose	☐ Ovulation Tests					
☐ Aspartate Aminotransferase (AST)	☐ Glycosylated Hemoglobin	□ pH					
☐ B-Type Natriuretic Peptide (BNP)	☐ HDL Cholesterol	□ Platelet Aggregation					
☐ Bacterial Vaginosis, Rapid	☐ Helicobacter Pylori	☐ Potassium					
☐ Blood Urea Nitrogen (BUN)	☐ Hematocrit	☐ Pregnancy Test (Urine)					
☐ Breath Alcohol (FDA OTC Devices Only)	☐ Hemoglobin	☐ Protime					
☐ Calcium	☐ HCV, Rapid	☐ RSV (Respiratory Syncytia	l Virus)				
☐ Carbon Dioxide	☐ HIV, Rapid <i>(*Submit Protocol w/App.)</i>	☐ Saliva Alcohol	,				
☐ Chloride	□ Influenza	☐ Sodium					
☐ Cholesterol	☐ Ketones	☐ Strep A Test (<i>Rapid</i>)					
☐ COVID-19 Antigen	☐ Lactic Acid (Lactate)	☐ Thyroid Stimulating Hormo	ne <i>(TSH</i>)				
☐ COVID-19 Molecular	☐ LDL Cholesterol	☐ Trichomonas, Rapid	, ,				
☐ COVID-19 Antibody	☐ Lead <i>(*Submit Protocol w/App.)</i>	☐ Triglycerides					
☐ Creatinine	☐ Microalbumin	☐ Urinalysis					
☐ Drugs of Abuse	☐ Mononucleosis	☐ Other:					
Indicate the combined estimated appual test w	huma for all Waived Test Dreadures indicated ab	•					
4. PROVIDER-PERFORMED MICROSCOPY	lume for all Waived Test Procedures indicated at						
	ctitioners, nurse midwives and physician ass		na to perform.				
 □ Direct wet mount preparations for the pres of bacteria, fungi, parasites, and human of present the present parasites in the present parasites. □ Fecal Leukocyte examinations □ Fern tests □ Nasal smears for granulocytes 	ellular elements mucous □ Potassium hydro □ Qualitative semen	qualitative examinations of vagina kide (KOH) preparations in analysis (limited to the presence action of motility)					
☐ Pinworm examinations	☐ Urine sediment e	xaminations					
Indicate the combined estimated annual test vo	lume for <u>all PPM Procedures indicated above:</u>						
5. CERTIFICATION. I understand that by signing this application form, I agree to any investigation made by the Department of Health to verify or confirm the information provided herein or adjunctive to this application, and any investigation in connection with my laboratory registration, a complaint or incident report made known to the Department. Registration under this subdivision may be denied, limited, suspended, revoked or annulled by the Department upon a determination that a laboratory services registrant: (i) failed to comply with the requirements of this subdivision; (ii) provided services that constitute an unwarranted risk to human health; (iii) intentionally provided any false or misleading information to the Department relating to registration or performing laboratory services; or (iv) has demonstrated incompetence or shown consistent errors in the performance of examinations or procedures. If additional information is requested, I will provide it. Further, I understand that, should this application or my status be investigated at any time, I agree to cooperate in such an investigation. Laboratory test registrants shall: (i) provide only the tests and services listed on the registration issued by the Department hereunder; (ii) advise the Department of any change in the registrant's name, ownership, location or qualified health care professional or laboratory director designated to supervise testing within thirty days of such change; (iii) provide the Department with immediate access to all facilities, equipment, records, and personnel as required by the department to determine compliance with this subdivision; (iv) comply with all public health law and federal requirements for reporting reportable diseases and conditions to the same extent and in the same manner as a clinical laboratory; (v) perform one or more tests as required by the Department to determine the proficiency of the persons performing such tests; and (vi) designate a qualified health care professional or qualified individual holdin							
application.							
Print Name of Multi-Site Network Laboratory Directo	Signature of Multi-Site Network Laboratory	Director	Date				
Print Name of Person Completing this Form	Signature of Person Completing this Form		Date				