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Introduction

The purpose of this manual is to provide clinical laboratories with the information needed to begin using the web-based, electronic clinical laboratory information management tool, eCLEP. It includes the following major sections:

- **Getting Started: An Overview** introduces a laboratory to eCLEP.
- **Requirements for Use** provides hardware and software specifications and configuration settings required to access eCLEP.
- **Navigating in eCLEP** provides detailed directions for accessing eCLEP and entering data.
- **Reapplication Submissions** provides detailed instructions for submitting the clinical laboratory permit reapplication.
- **Open Mode Submissions** provides detailed instructions for submitting changes in facility information outside of the permit reapplication period.

Getting Started: An Overview

The New York State Department of Health (NYSDOH) has developed eCLEP to enable clinical laboratories to exchange information electronically in place of mailing paper forms. This web-based application supports the inquiry, maintenance, and reporting requirements as defined by the Wadsworth Center Clinical Laboratory Evaluation Program (CLEP) and acts as a single repository for the data. eCLEP has evolved to support the submission of permit reapplications and notification of laboratory changes, as well as provide each clinical laboratory the ability to check their laboratory licensure status 24/7.

*Note:* the eCLEP application does not service Limited Service Laboratories. Please see our website at [https://www.wadsworth.org/regulatory/clep/limited-service-lab-certs](https://www.wadsworth.org/regulatory/clep/limited-service-lab-certs) for information on Limited Service Laboratories.

eCLEP offers many advantages over existing paper-based processes, including:

- **Persistent Data** – The system displays general laboratory information as found in the Clinical Laboratory Evaluation Program’s licensure database. The most current information is displayed, eliminating redundant data entry.

- **Data Validation** – User entries are validated for incorrectly formatted and incomplete submissions at every step, eliminating submission failures and the need for follow-up communications to correct minor errors such as missing entries.

- **Delegating Submission** – The Laboratory Director may delegate the electronic submission of Laboratory information.

- **Documented Delivery** – Permit reapplications and changes to laboratory information are electronically transmitted; the time of the submission and username submitting the data is recorded.
Requirements for Use

To enter information into the eCLEP system, your laboratory must have a personal computer that is minimally configured as follows:

- Pentium processor or higher
- DSL or a broadband Internet connection (The laboratory is responsible for obtaining Internet access with an Internet Service Provider (ISP)).
- Printer (optional)

Browser Requirements and Configuration

Access to the Health Commerce System and eCLEP requires 256-bit encryption, browser setting to accept cookies and enabling of Javascript.

Supported browsers on desktop computers include: Microsoft Internet Explorer, Google Chrome and Safari (Mac OS only). Support browsers on mobile devices include: Google Chrome (iOS5.1/Android 4.0 or later) and Safari (iOS5.1 or later). The Health Commerce System supports the current and two previous versions supported browsers.

Limited support is available for the following browsers: Mozilla Firefox (desktop/mobile) and WebKit-based browsers. Microsoft Internet Explorer Mobile and Safari for Windows are not supported.
Roles and Responsibilities

This section describes the different levels of eCLEP users and their access and data submission privileges in the system. It also gives instructions on how to request access to the system.

eCLEP users at the laboratory will belong to one of two roles. Below is a description of the roles, followed by the user qualifications:

**Laboratory Director**

- View Laboratory information
- Update data
- Review the "eCLEP Summary"
- Attest to the accuracy of the entered data and submit it electronically.

**Assistant Director / Delegated Submitter**

- View Laboratory information
- Update data
- Review the "eCLEP Summary"
- Attest to the accuracy of the entered data and submit it electronically.

A Laboratory Director is an individual who is responsible for the administration of the technical and scientific operation of a clinical laboratory or blood bank, including the supervision of procedures, reporting of results and responsibilities specified in Section 19.3 of 10 NYCRR (New York Codes, Rules and Regulations) and Article 5 Title V Section 571 of the Public Health Law. Such person shall possess a Certificate of Qualification issued pursuant to Part 19 of 10 NYCRR. A Director is authorized to **view**, **enter**, **attest**, and **submit** laboratory information electronically using the eCLEP system.

An Assistant Director is a person who has been designated by the Laboratory Director to serve as an Assistant Director in one or multiple categories or subcategories of testing. Such person shall possess a Certificate of Qualification issued pursuant to Part 19 of 10 NYCRR. A responsible Assistant Director holding a Certificate of Qualification is authorized to **view**, **enter**, **attest**, and **submit** laboratory information electronically using the eCLEP system.

A Delegated Submitter is a person who has been given written authorization by the Laboratory Director to electronically submit laboratory information on behalf of the Laboratory Director. A Delegated Submitter is authorized to **view**, **enter**, **attest**, and **submit** laboratory information electronically using the eCLEP system.
HCS Access Permissions

Before logging on to eCLEP to submit data, you will need access to the New York State Health Commerce System (HCS) at https://commerce.health.state.ny.us.

The New York State Department of Health assigns a NYSDOH HCS Account ID (User ID) and password to each individual who has been granted access to the HCS.

As the HCS contains confidential information, safeguard your HCS User ID and password by not revealing them to other users. Violation of the security and use agreement (e.g. sharing your User ID and password with someone else) will result in the temporary suspension of your account privileges and repeat offenses may result in the permanent removal of the account. Also, do not leave your computer logged on to the HCS unattended. For security purposes, there are session timeouts after one hour of inactivity and system timeouts after eight hours of total connectivity.

Clinical Laboratory Directors and HCS Coordinators without HCS accounts

Clinical Laboratory Directors without HCS accounts may begin the HCS account application process with the HCS Affiliation Request form available at www.wadsworth.org/regulatory/clep/clinical-labs/obtain-permit/health-commerce. After completing the form, fax it to 518-449-6901. The Laboratory Director will receive an e-mail from camu@its.ny.gov which will include a bar-coded PDF document to sign and have notarized. This form must be returned to the Commerce Account Management Unit (CAMU) to complete the affiliation process. Laboratory directors are expected to complete and submit this form promptly. An amended permit reflecting the change in directorship will not be issued until the laboratory director’s HCS account has been verified.

The HCS Affiliation Request form is also used to establish HCS Coordinators at your laboratory.

Requesting HCS Accounts for Other Individuals

The Laboratory Director or HCS Coordinator for the laboratory can electronically request an account for additional laboratory staff. The Laboratory Director or HCS Coordinator needs to log into the Health Commerce System at https://commerce.health.state.ny.us, select the Coordinator’s Account Tools (left side under My Applications), then click on the appropriate ‘Request an Account for…’ link.

Delegated Submitter

The Laboratory Director may delegate data submission privileges to a staff member who already has an HCS account by signing and completing a Delegated Submitter Request form. The form is available at www.wadsworth.org/regulatory/clep/clinical-labs/obtain-permit/health-commerce.
Accessing eCLEP and the Permit Materials Module

1. To access the eCLEP Home Page enter the following web address into an Internet browser:
   https://commerce.health.state.ny.us

2. Enter your User ID and Password into the HCS Login screen and click Sign In:

3. The HCS Homepage displays. Look for eCLEP in the left frame under My Applications:
4. Click on eCLEP in the left frame and the eCLEP Home Page will display. Click on Permit Materials, Laboratory Reapplication / Laboratory Changes area at the upper right.

5. HCS account holders affiliated with more than one laboratory will be required to enter an appropriate 4 digit numeric Permanent Facility Identifier (PFI).

(An alphanumeric PFI denotes a Limited Service Laboratory (LSL). LSLs are not serviced by the eCLEP application.)
6. Most users, however, will be brought directly to the Reapplication Mode or Open Mode page. The reapplication period occurs in April, actual dates will vary year to year. Open Mode is available the rest of the year, provided there are no laboratory information changes submissions pending. Note: Reapplication mode is denoted by the presence of green bar at the top of the screen with the dates of the reapplication period; Open mode is denoted by the presence of teal bar at the top of the screen.

7. On the Reapplication Mode or Open Mode page, you may want to review the Current Data on File (without pending changes) using the “View Summary” link and print it out to use as a worksheet.
8. Make changes to laboratory information as required using the links on the dark blue menu bar (Lab Profile, Ownership, Personnel, Permit Categories/Tests, and PSCs and Others).
Reapplication Mode Submissions

NOTE: All eCLEP submissions are reviewed by the Clinical Laboratory Evaluation Program prior to acceptance. The Program reserves the right to request additional information, request re-submission to obtain missing information, or to reject the request in total if the eCLEP submission is not acceptable. eCLEP submission does not constitute approval by the Program.

We suggest that you first review the information on file for your laboratory and make any necessary revisions prior to beginning the reapplication submission. If you have already been navigating through the sections on the blue menu bar, click on the Reapplication Center button on the green menu bar to return to the main Reapplication Mode page.

1. Alternatively, you may start the reapplication submission process before revising facility information, however; once you begin navigating through the sections indicated on the blue bar to provide required information, you must return to the main Reapplication Mode page to continue with the submission process.

Click Enter to begin the reapplication submission process.
The **Step 1: Review and Update** page displays the data on file in the Laboratory Licensure database (and any pending changes already entered via eCLEP) for your facility. Review and click **Next**. A printable version of this information is available by clicking the “Printable Summary in PDF Format” link.

<table>
<thead>
<tr>
<th>Permit Materials</th>
<th>Proficiency Testing</th>
<th>LDT Approval</th>
<th>Survey</th>
<th>Limited Labs</th>
<th>Select Facility</th>
</tr>
</thead>
</table>

**Reapplication Center**

Reapplication Period: Feb 09, 2017 through Feb 24, 2017 at 05:00 PM EST

**PFI: 0888  Name: Internal Test for EPRIS**

- **Step 1: Review and Update**

Please review the summary below for accuracy and completeness. If you need to make changes, click the appropriate link in the blue menu bar above (for example, facility address data is found in the Lab Profile area). Once in an area, you'll be able to make changes to the data.

Once you are satisfied that the information in the summary below is complete and accurate, click “Next” to continue.

<table>
<thead>
<tr>
<th>Field Name</th>
<th>Current Data</th>
<th>Changes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory PFI</td>
<td>0888</td>
<td></td>
</tr>
<tr>
<td>Contact Person (other than director):</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phone:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Email:</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Step 2: Provide Required Data** page will list sections/subsections that you will have to visit in order to complete the reapplication. Required information that must be completed before you will be able to submit include:

- laboratory contact person
- owner declaration and Disclosure of Ownership and Controlling Interest Statement upload
- facility e-mail
- test volume, if applicable
- POC testing, if applicable
- PSCs and Others tab, if applicable

You may proceed to the areas with outstanding data requirements by either method below, or a combination of these two methods.

a) Navigate to each section by clicking the links in the blue menu bar near the top, e.g. **Lab Profile, Ownership, Personnel, Permit Categories/Tests, PSCs and Others**.

b) If there are no, or few, owner/personnel/testing changes during this reapplication, you may navigate directly to the sections with outstanding data requirements by clicking the underlined links in the **How to Resolve** column.
To return to the **Step 2: Provide Required Data** page to resolve further outstanding data requirements, or to verify that all data requirements have been resolved, click the button on the green bar at the top of the screen from any page to get back to the main Reapplication Mode page.
Click Enter, and then click Next on the Step One: Review and Update page. The Step 2: Provide Required Data page will list any outstanding data entry requirements. If there are no outstanding data entry requirements, the Required Data table will read “All required data has been provided.” Only after all data requirements have been resolved will you be able to proceed to Step 3: Attest and Submit by clicking Next.

Please read the Step 3: Attest and Submit page in its entirety and click the checkbox to signify that you have read, and agree with, the attestation; then click Next.
The Step 4: Print for Your Records page allows access to the eCLEP summary in PDF format. Click on the Submission dated [date, time] (PDF) and print or save this document for your records, if desired. Then click Finished. You will be directed to the main Read-Only mode page.
Open Mode Submissions

NOTE: All eCLEP submissions are reviewed by the Clinical Laboratory Evaluation Program prior to acceptance. The Program reserves the right to request additional information, request re-submission to obtain missing information, or to reject the request in total if the eCLEP submission is not acceptable. eCLEP submission does not constitute approval by the Program.

Enter laboratory changes as necessary by navigating the blue menu bar. Click on the Submit Changes button to begin the Open Mode submission process.

Click Enter to begin the Open Mode submission process.
The **Step 1: Review and Update** page displays the data on file in Laboratory Licensure database (and any pending changes already entered via eCLEP) for your facility. Review and click **Next**. A printable version of this information is available by clicking the “Printable Summary on PDF Format” link.
Please read the **Step 2: Attest and Submit** page in its entirety and click the checkbox to signify that you have read, and agree with, the attestation; then click **Next**.

The **Step 3: Print For Your Records** page allows access to the eCLEP summary in PDF format. **NOTE:** The eCLEP Summary is no longer required to be signed and returned to CLEP. Click on the Submission dated [date, time] (PDF) and print or save this document for your records, if desired. Then click **Finished**. You will be directed to the main Read-Only mode page.
Navigating in the Permit Materials Module

Lab Profile

General Information section

The General Information webpage allows you to make changes to the laboratory name and address, facility type and lab contact information. Note an effective date for any laboratory name and address changes is required. Enter the required information and click Save.
Regulatory Information section

The Regulatory Information webpage allows you to revise the CLIA registration and Medicaid number for the laboratory. Enter the required information and click Save.

Hours section

The Hours section allows you to change laboratory testing hours. Enter the required information and click Save. Note: The Clinical Laboratory Evaluation Program may seek clarification of information entered in the “Hours Note” field before accepting the proposed change.
Contact Person section

The **Contact Person** section allows you to change/update the contact person for the laboratory and their contact information (e-mail and phone number).

- The laboratory contact person is the individual who is designated by the laboratory director and owner(s) to communicate with the Department on matters relating to the clinical laboratory permit.
- You are required to verify/update the **Contact Person** in Reapplication mode.

**USER TIP:** More than one email address may be entered in the Contact Person Email field by separating each address with a comma.
Ownership

The Ownership section is divided into three subsections, **Owner, Declaration, and Upload**.

Labs will be required to upload a list of direct and indirect owners using the Upload Feature as part of the permit reapplication.

- **Direct ownership** means an individual or entity with an ownership interest or controlling interest in the applying facility.

- **Indirect ownership** means an individual or entity with an ownership interest, controlling interest, or corporate membership, in an entity with direct or indirect ownership in the applying clinical facility. Indirect owners who hold a ten (10) percent or greater ownership interest, controlling interest, or corporate membership, are required to be disclosed by the applying clinical facility.

**Examples of ownership structures:**

**Example 1 (Business Corporation):** ABC Lab is owned by ABC Lab, Inc. ABC Lab Inc. has two major stockholders, Mr. Smith and Mr. Hernandez. ABC Lab, Inc. is the direct owner. Mr. Smith and Mr. Hernandez are indirect owners.

**Example 2 (Business Corporation):** ABC Lab, Inc. dba ABC Lab is owned by ABC Lab, Inc. ABC Lab, Inc has two primary investors; Umbrella Corp, Inc. and Ms. Smirnov. ABC Lab, Inc., is the direct owner. Umbrella Corp, Inc. and Ms. Smirnov are indirect owners.

**Example 3 (Partnership):** Acme Lab is owned by Zhang Brothers, LLP. The partners of Zhang Brothers, LLP are Zhang Industries and Mr. Lee. Zhang Industries is owned by A. Zhang and B. Zhang. Zhang Brothers, LLP is the direct owner. Zhang Industries, Mr. Lee, A. Zhang, and B. Zhang are all indirect owners.

**Example 4 (Not-for-Profit Corporation):** Healthy Hospital Laboratory is owned by Healthy Hospital, Inc., a not-for-profit corporation. Healthy Hospital, Inc. has two corporate members, Biggie Health Systems, Inc. and Bigger Health Systems, Inc. Biggie Health Systems, Inc. and Bigger Health Systems, Inc. are considered indirect owners in Healthy Hospital Laboratory.

**Example 5: (Professional Corporation):** Neighborhood Physicians, PLLC operates a clinical laboratory. Neighborhood Physicians, PLLC is owned by Hospital Physicians, PC and Dr. Patel. Hospital Physicians, PC and Dr. Patel are indirect owners.

- **Ownership Interest** means the possession of stock, equity in the capital, or any interest in revenue of an entity.

- **Controlling interest** means the ability to direct or control the operation or management of an entity. Members on the Board of Directors or Board of Trustees for not-for-profit corporations are considered to have controlling interests. Any individual or entity with a ten (10) percent or greater controlling interest is required to be disclosed by the applying clinical facility. Licensed physicians who are included on the Board of Directors/Board of Trustees for a not-for-profit corporation are required to disclose their authority to order laboratory tests if they have greater than 10% controlling interest in the applying clinical facility.
**ECLEP Manual**

- **Corporate membership** means an individual or entity with a voting interest in a not-for-profit corporation that directly owns the applying facility. Corporate membership includes, but is not limited to, the right to vote in the election for directors of the clinical laboratory or on fundamental corporate transactions such as closing the business or amending the bylaws.

- **Management company** means any organization that operates and manages a clinical laboratory on behalf of the owner, with the owner retaining ultimate legal responsibility for the operation of the business.

- During the **Reapplication** period, you will be required to enter any missing data and/or update information. The reapplication cannot be submitted without providing this information. You will receive error messages when you try to continue without addressing these fields. When this happens, please enter the missing data, select a dropdown option and/or click the radio button; then click **Save** again.

- During the **Open Mode**, update information as necessary to accurately reflect a laboratory change.
Owner section

This section captures information such as the owner type, Federal Employer Identification Number (EIN, aka TIN), owner name, etc. If the response to question 1 is "Yes", you will be prompted to upload a list of all laboratories in which any of the direct or indirect owners have ownership, controlling interest, or corporate membership.

PLEASE NOTE: All laboratories that share a common Federal Employer Identification Number (EIN) are considered to be owned by the same entity and disclosure of the other laboratories owned by the direct owner is required. Note that to complete this section, the applying facility should consult their administration and/or legal department. It is not necessary to include Limited Service Laboratories in this list.
During Reapplication, all laboratories are required to upload a list of direct and indirect owners of the laboratory.

The list of direct owners must include (based on ownership type):

- **Individuals**: Names, addresses, percentage of ownership, and social security numbers of individual owners.
- **Partnership**: Names, addresses, percentage of ownership, and social security numbers of all partners.
- **Government**: The governmental entity and name of the representative official (i.e., Commissioner of Health, Chancellor, etc.) who can be contacted regarding ownership issues.
- **For-Profit Corporation**: Names, addresses, percentage of ownership, and social security numbers (or EIN) for corporate officers, and/or shareholders.
- **Not-for-Profit Corporation (NFPC)**: A list of the Board of Directors/Trustees/Governors of the NFPC.
- **Other**: Names, addresses, percentage of ownership and SSN or EIN, as appropriate.

The list of indirect owners must include those individuals or entities that 1) possess ten (10) percent or more of the voting shares of an entity that directly owns/operates a clinical laboratory; 2) maintain a controlling interest of ten (10) percent or more in an entity that directly owns/operates a clinical laboratory; or 3) maintain corporate membership in a not-for-profit corporation that directly owns/operates a clinical laboratory.

The list must include (based on ownership type):

- **Individuals**: Names, addresses, percentage of ownership, and social security numbers of individual owners
- **Partnership**: Names, addresses, percentage of ownership, and social security numbers of partners
- **For-Profit Corporation**: Names, addresses, percentage of ownership, and social security numbers (or EIN) for corporate officers, and/or shareholders
- **Not-for-Profit Corporation**: A list of the Board of Directors/Trustees/Governors of the NFPC.
Declaration section

Respond to the questions presented in this section. For each “Yes” response, the laboratory will be prompted to upload supplemental documentation. These documents will be uploaded in the Upload screen described below.

1. Has the director, any assistant director(s), or those having a direct or indirect ownership or controlling interest in the applying clinical facility ever been charged with violations of local, state or federal laws, rules and regulations, including, but not limited to the Public Health Law or related statutes, concerning the provision of health care services or reimbursement for such services? To the extent that such charges are currently pending, respond “yes”.
   - Yes
   - No

   On a separate sheet, identify the individuals (directors or owners) who have sustained charges and the details of such charges. The PFI number of the laboratory must be included on this sheet. This sheet must be uploaded in the field labeled “Director/Owner Violation or Charges” on the Upload page.

2. Has the director, any assistant director(s), or those having a direct or indirect ownership or controlling interest in the applying clinical facility ever been charged with any crime, including, but not limited to any offense related to furnishing of or billing for clinical laboratory services and medical care, services or supplies, or which is considered an offense involving theft or fraud? To the extent that such charges are currently pending, respond “yes”.
   - Yes
   - No

   On a separate sheet, identify the individuals (directors or owners) who have convictions and the details of such convictions. The PFI number of the laboratory must be included on this sheet. This sheet must be uploaded in the field labeled “Director/Owner Crime Conviction” on the Upload page.

3. Are any individuals, with direct or indirect ownership or controlling interest in the laboratory or blood bank, licensed health professionals, authorized by law to order clinical laboratory tests and receive results?
   - Yes
   - No

   On a separate sheet, identify the individuals with greater than ten (10) percent controlling interest who are authorized to by law to order clinical laboratory tests. The PFI number of the laboratory must be included on this sheet. This sheet must be uploaded in the field labeled “List of Authorized Individuals” on the Upload page.
If a laboratory declares it has entered into a new management contract, a follow-up request to submit a copy of the contract to CLEP will be made, there is currently no upload feature for management contract submission.
Upload section

Depending on the laboratory’s responses to the questions on the Declaration page, users will see one or more fields requesting specific documents to be uploaded. During Reapplication, all laboratories are required to upload a list of direct and indirect owners of the laboratory. Refer to page 22 for definitions of direct and indirect owners and page 25 for specific instruction on reporting the ownership.

To upload a document, verify the document type you wish to upload matches the document type on the screen (List of Owners, List of Other Labs Owned, Director/Owner Violation or Charges, Director/Owner Crime Conviction, List of Authorized Individuals) then click Browse button to the right of the File Name space. Navigate to the electronic file on your computer, then click Open to upload.

If you accidently upload the wrong document, you may click on Browse button again and choose another document, the original uploaded document will be overwritten.

Once all documents have been uploaded, click Save.
Personnel

The Personnel section has many subsections, including Director, Assistant Director, and Responsibilities. Note that any yellow highlighted areas are required. You will need to know the Certificate of Qualification (CQ) code of any new directors or assistant directors. The CQ code (five letters followed by a number) can be found on the individual’s certificate. If you are unable to locate this document for the individual, please call (518) 485-5378 or e-mail CLEP@health.ny.gov for help in looking up CQ codes.

- During the Reapplication mode, please review each subsection for accuracy.

Director section

The Director section allows you to view and update current on-site hours for the Laboratory Director as well as appoint a new Laboratory Director. Update hours as needed and click Save. Note: The Clinical Laboratory Evaluation Program may seek clarification of the Director's work schedule before accepting the proposed change.

On-site hours for a Laboratory Director may not overlap with hours at another facility. The laboratory will be required to submit new hours to eliminate such overlaps. The other laboratory(ies) where the director is employed may also need to revise the director’s work schedule.
To appoint a new Director, enter the CQ Code of the new director and the effective dates of the change, click Next.

Note: In order to indicate a replacement for the outgoing laboratory director in eCLEP, the incoming director must hold a valid Certificate of Qualification and his/her CQ code must be entered. If the incoming director does not currently hold or has not applied for a Certificate of Qualification, please contact clepcert@health.ny.gov for alternate instructions.

- Note, when a new Laboratory Director is appointed, s/he must also complete and submit an HCS Affiliation Request form available at www.wadsworth.org/regulatory/clep/clinical-labs/obtain-permit/health-commerce.

- Note, a current Laboratory Director cannot be removed from the laboratory without identifying a replacement. If the incoming director does not currently hold a Certificate of Qualification, please contact CLEP at clepcert@health.ny.gov or (518) 485-5378 for alternate instruction.
On the following screen, enter the new director’s on-site hours, click **Next**. **Note:** The Clinical Laboratory Evaluation Program may seek clarification of the Director’s work schedule before accepting the proposed change.

On-site hours for a Laboratory Director may not overlap with hours at another facility. The laboratory will be required to submit new hours to eliminate such overlaps. The other laboratory(ies) where the director is employed may also need to revise the director’s work schedule.

Review the additional places of employment for the new director; add additional facilities as needed; click **Next**.

Indicate the permit categories in which the new director will have responsibilities, click **Finish**. The check box list includes all categories either held or in applied status for the laboratory.

**Note:** The new director must hold a Certificate of Qualification in the corresponding category to allow assignment of responsibility for a permit category. If the laboratory director does not hold the appropriate corresponding category on his/her Certification of Qualification, the request for assignment of responsibility for the permit category will be rejected. An individual may not serve as laboratory director unless s/he is assigned responsibility for at least one permit category.

This page also allows the laboratory to request one additional category by choosing a category from the New Category dropdown below the check box list, then click **Finish**.
The next page will display the new director change. Review the information for accuracy and click Save.

**Assistant Director section**

The **Assistant Director** section allows you to view the current assistant directors, update assistant director on-site hours, add a new assistant director(s), and remove an assistant director(s).
To add an Assistant Director, please follow the steps as presented above for appointing a new Laboratory Director.

- To update the on-site hours for an Assistant Director, either click on the individual’s name in the View page (see above) or choose the individual from the drop down list presented on the Update Hours page, click Next.

- On the next screen, update the hours as needed, click Save. Note: The Clinical Laboratory Evaluation Program may seek clarification of the Assistant Director’s work schedule before accepting the proposed change. On-site hours for an Assistant Director may not overlap with hours at another facility. The laboratory will be required to submit new hours to eliminate such overlaps.
To remove an Assistant Director, either click the remove link next to the individual's name on the View page; or choose the individual from the dropdown list presented on the Remove page, click Next.

On the following page, enter the effective date of the Assistant Director's departure, click Remove.
Note: If the departing assistant director is the sole individual responsible for a permit category(ies), the Clinical Laboratory Evaluation Program will notify the director that the laboratory is in jeopardy of losing an approved (or pending) permit category unless a timely arrangement is made for assigning a qualified person (current or new) to be responsible for the permit category.
Responsibilities Section

This section allows the laboratory to view all the permit categories and the corresponding CQ holders with responsibility. On the “View” screen, clicking on a Director’s name will allow you to edit the responsibilities for that individual.

Form the “Update” screen, choose a Director from the dropdown to make edits to responsibilities.
Existing permit category responsibilities are indicated by a check mark. Additional permit categories can be requested by adding a check mark next to the desired category and clicking 'Save'.
Permit Categories/Tests
The Permit Categories/Tests sections allows you to:
- add permit categories to the laboratory permit;
- change permit category responsibilities for the laboratory director and/or assistant director(s);
- remove permit categories from the laboratory permit;
- enter test volumes (required for laboratories located in NYS during permit reapplication).

Responsibilities section
Under the Responsibilities section, you may view the laboratory’s current permit categories, the status of each category, and the laboratory director (DI) /assistant director (AD) responsible for each permit category.
- Click on the permit category name to view the current DI/AD responsible for the category and to add or remove individuals as responsible.
Alternatively, choose the category to update from the Responsibilities Update page, click Next. This dropdown menu will include all categories that the laboratory has applied for (pending) and those already held (approved). This will take you to the same page as above.

**Note:** Personnel changes still pending review by the Department will not appear as available for responsibility assignment (e.g., changes entered but not yet submitted in eCLEP). Only Certificate of Qualification holders already associated with the laboratory will be listed. A new Assistant Director must be added through the Personnel section.

On the following page, indicate the effective date of the individual’s new responsibility, click Add.
Category Upload – Cytopathology Proficiency Testing Enrollment

During permit reapplication, laboratories holding the category of **Cytopathology – Gynecological Testing** are required to upload proof of enrollment in a CMS-approved proficiency testing (PT) program. **Acceptable documentation is an enrollment confirmation from the PT program.** Purchase orders and order forms are not acceptable.

- The enrollment confirmation must reference the laboratory name and address.
- The PFI number of the laboratory must be handwritten on the paper if the CLIA number is not already included.
- If the laboratory personnel participate in PT at another site, the order confirmation for “paper enrollment” must be provided.

---

![Image of category upload process](image_url)
Add a Category

To request to add a permit category, click on the **Add New** hyperlink from the left panel under Category.

Choose the desired permit category from the dropdown menu and choose a responsible director or assistant director for the new category using the individual’s CQ code. If you do not see the individual’s CQ code in the list, you must add the individual under the Personnel tab before proceeding with the Add Category request. Please note that both Permit Category and CQ Code fields are mandatory.

Once the Permit Category and CQ code have been chosen, click the Next button. When requesting to add a category that includes analytes/test that are described in CLIA Subpart I (42 CFR 493 Subpart I), you will be required to indicate the CMS-approved proficiency test provider and product that will be used to satisfy proficiency testing requirements.
If you are unsure of what category is required for the testing that will be offered by the laboratory, you can use the search engine to search for category by test name. Please make sure the browser you are using is not blocking pop-ups, otherwise your search result will not be displayed.

---

**ECLEP Manual**

If your laboratory is preparing to offer laboratory-developed tests (LDT), in the new permit category, you must submit the materials specified in the Test Approval section of the Clinical Laboratory Evaluation Program's public website, Test Approval, for each LDT and receive explicit approval prior to initiating patient testing.

**Add New Category**

Please select the category you wish to add to the clinical laboratory permit. If you are unsure of the permit category for the test you wish to offer, please search for the category by entering the test name in the Search field. You may also review the Program Guide for permit category descriptions at our website at www.wadsworth.org/regulatory/c挑剔.

Once a permit category is chosen, the Certificate of Qualification code (CQ Code) of the responsible Director or Assistant Director must be entered. Please note the Director and/or Assistant Director assigned to this new category must hold the relevant corresponding category on his/her Certificate of Qualification or be in the process of adding the category to the CQ.

Note: The Clinical Laboratory Evaluation Program assumes the laboratory is prepared to meet applicable requirements for permit approval on the date the new permit category request is submitted. These requirements may include successful participation in an on-site survey, enrollment, and successful participation in proficiency testing, and review and approval of validation materials for laboratory-developed tests.
Indicate Tests Offered on NYS Specimens

This page provides a list of tests that are described in CLIA Subpart I that are included under the new category. Please indicate whether you offer these tests or not by selecting an option from the drop-down menu.

The hyperlink Category Specific Help provides additional Proficiency Testing guidance by category.

All fields in this page are mandatory. Click Next button to proceed to the next page.
Designate PT Provider and Product

This page displays the tests that have been marked as “Test Offered” offered on the previous page.

Please provide the **PT Provider** and **Product** for each test and then click **Save** to proceed.

All fields in this form are mandatory.
**View Designation** page is a summary of the Proficiency Testing information that had been entered. Review and click on Next button to complete the process.

<table>
<thead>
<tr>
<th>View Designation</th>
<th>Page is a summary of the Proficiency Testing information that had been entered. Review and click on Next button to complete the process.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Permit Materials</td>
<td>Proiciency Testing</td>
</tr>
</tbody>
</table>

**Tests Offered**

<table>
<thead>
<tr>
<th>Category</th>
<th>Test</th>
<th>Provider</th>
<th>Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacteriology</td>
<td>Gram stains</td>
<td>Medical Laboratory Evaluation</td>
<td>Bacteriology 2 - 640</td>
</tr>
<tr>
<td></td>
<td>Group A Streptococcus direct detection</td>
<td>American Academy of Family Physicians</td>
<td>Group A Strep - 783</td>
</tr>
<tr>
<td></td>
<td>Identification of bacteria by culture</td>
<td>AABB Proficiency Testing Service</td>
<td>Gonadal Culture - 2009523</td>
</tr>
<tr>
<td></td>
<td>Identification of bacterial meningitis pathogens by molecular methods</td>
<td>American Proficiency Institute</td>
<td>Meningitis Panel - 371</td>
</tr>
<tr>
<td></td>
<td>Identification of genital pathogens (bacterial) by molecular methods</td>
<td>College of American Pathologists</td>
<td>Vaginitis Screen - VS</td>
</tr>
<tr>
<td></td>
<td>Identification of respiratory bacterial pathogens by molecular methods</td>
<td>College of American Pathologists</td>
<td>Infectious Disease Respiratory Panel - IDR</td>
</tr>
<tr>
<td></td>
<td>Susceptibility (bacterial) testing (AST)</td>
<td>Acucare Inc</td>
<td>Bacterial Identification - BACT435</td>
</tr>
</tbody>
</table>

**Tests Not Offered**

<table>
<thead>
<tr>
<th>Category</th>
<th>Test</th>
<th>Provider</th>
<th>Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacteriology</td>
<td>Chlamydia/Nelseria gonorrhoea by direct detection</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Clostridium difficile direct detection</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bacteriology</td>
<td>Identification of blood pathogens (bacterial) by molecular methods</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Identification of gastrointestinal bacterial pathogens by molecular methods</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| Next |
Adding More Than One CQ Holder to a New Category

To add multiple CQ holders to a new category, first add the new Category, then go to View under Responsibilities and select the newly added Category.

<table>
<thead>
<tr>
<th>Permit Categories</th>
<th>Responsibilities</th>
<th>Responsible Directors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Category status</td>
<td>Pending Add</td>
<td>Approved</td>
</tr>
<tr>
<td>Radiology</td>
<td>Pending</td>
<td>Todd Lee</td>
</tr>
<tr>
<td>Blood Gas and Co2</td>
<td>Pending</td>
<td>Todd Lee</td>
</tr>
<tr>
<td>Clinical Chemistry</td>
<td>Pending</td>
<td>Todd Lee</td>
</tr>
<tr>
<td>Virology</td>
<td>Pending</td>
<td>Todd Lee</td>
</tr>
</tbody>
</table>

Then proceed to add additional CQ holders to the new Category:
Pending Changes page displays the list of all unsubmitted requests.

To cancel an Add Category request: select a change request by clicking the box to the left of the category name and press the Cancel Selected Changes button.

To modify the Add Category request: Click on the hyperlink PT Changes. This will allow user to modify ONLY the Proficiency Testing information entered.
Remove a Category

Under the **Category** subsection, you may remove a permit category from the laboratory's permit. Select the category to remove from the dropdown list, click **Delete**.

![Image of the Delete Permit Category screen](image)

**Note:** When a permit category is removed, the director's and/or assistant director(s) assigned responsibility for that permit category will also be removed.

On the following page, indicate the effective date of the permit category deletion, and then click **Delete**.

![Image of the Delete Permit Category screen with effective date](image)
Test Volume

Note: this section is only visible to laboratories located in New York.

Laboratories located in New York are required to report Test Volume for each category of testing. The Test Volume section allows you to view the volumes of testing entered during the previous reapplication period and, in the Reapplication mode, enter the previous year’s testing volumes for each permit category of testing. A Guidelines for Reporting Test Volume is available in the Tools Section of eCLEP. Please contact CLEP at CLEP@health.ny.gov or call (518) 485-5378 for questions on reporting test volumes.

- In the Open mode, you can view the current information in the database.

- In Reapplication mode, you can view the information currently in the database as well as enter the previous year’s testing volumes. Enter volumes for each permit category held on the laboratory permit. Use the scroll bar to view all categories.
  - If you indicate “No tests performed this year”, you must provide a reason.
To obtain a pdf version of the previous year's test volume, access the previous year's Reapplication Submission from the Reapplication Center page and print or save as needed.

Reapplication Mode

It's time to reapply for your facility's permit. Click the 'Enter' button below to complete the reapplication process and ensure you receive your new permit by July 1.

Enter

If you would like to view the data currently on file for your facility, or view submissions your facility has made, use the links below.

Current Data on File (without pending changes)

View Summary

Electronic Submissions

- Submission dated Aug 30, 2013 12:44:49 PM EDT (PDF)
- Submission dated Apr 23, 2013 1:50:40 PM EDT (PDF)
- Reapplication Submission dated Mar 29, 2013 2:37:07 PM EDT (PDF)
- Submission dated Mar 1, 2013 2:31:20 PM EST (PDF)
- Reapplication Submission dated Jul 31, 2012 2:43:05 PM EDT (PDF)
POC Testing

This section is visible only to laboratories at hospitals, Article 28 facilities, correctional facilities, etc., located in New York.

The Point-of-Care (POC) Testing section allows you to manage locations and testing performed at the point of care, rather than the laboratory proper, at the facility.

- Under Manage Locations, you may add or delete Point-of-Care Testing (POCT) locations.

- The Add page allows you to add a test to a POCT location. Choose a POCT location from the dropdown list, choose the test being performed from the dropdown list, enter the instrument used and finally choose the staff performing the testing by selecting the check box next to the appropriate staff description. Click Save.
The Delete/Update page allows you to update or delete a test from a POCT location.

- Choose a test by clicking the appropriate radio button, then click Update or Delete, as appropriate.
- Clicking Delete will automatically remove the test from the list.
- Clicking Update will bring you back to the Add page, where you can revise the appropriate information, click Save.
The **Point-Of-Care Contact Person** page allows you to indicate a POCT Coordinator for the laboratory. Enter the appropriate contact information and click **Save**.
PSCs and Others

The PSCs and Others tab allows the laboratory to request approval to operate a patient service center (PSC) and/or health fair (HF). This area also allows you to update the PSC and HF information (location, phone number, etc.) and complete the annual reapplication process for both.

Note: This feature was introduced for the 2016 reapplication.

PSC Reapplication

During the reapplication period each Spring, laboratories currently operating an approved patient service center (PSC) should review the current data on file with the Department and update such information as appropriate. To review, click on the PSC link on the left of the screen.
On the next screen, choose ‘Stations on File’ from the menu on the left to view all stations associated with the laboratory.

A list of all patient service centers is viewable and printable from a new screen under the PSC section. Click on the Print PSC Listing link to print or save the list.

To make updates to an existing PSC, click on the ‘New/Select’ link from the PSC menu on the left. Then choose the desired PSC from the dropdown box and click ‘Next’.
If an existing PSC location has been selected, the menu of links on the left of the screen will now look different and the PSC address screen will be shown. Users can update the Address, Contact and Hours screens. Click **Save** after making changes. Address changes require an effective date.

Also during the reapplication period each Spring, the laboratory will be requested to attest that the relevant NYS regulations and standards for the operation of a PSC and/or HF have been reviewed to ensure compliance by the laboratory. Click the check box next to the highlighted text to indicate this, then click **Save**.
Request a New PSC

To request approval to operate a patient service center (PSC), click on the PSC link on the left of the screen. On the next screen, click on the New radio button, then click Next.

On the next screen, fill in the requested information and click Save. Please allow at least two weeks for processing; enter the expected opening date accordingly. Please be reminded that the PSC cannot operate without explicit approval from the Department.
Once you click Save, the links on the left will change.

To complete the application process, a self assessment must be completed and requested documents (i.e., floor plan and lease) must be uploaded. Click on **Self Assessment**. Answer the questions provided.

Once all questions have been answered, click **Save**.
After all the questions have been answered and the responses have been saved, click **Upload** on the left of the screen to upload a copy of the PSC floor plan and lease. Click on **Browse** to navigate to the electronic file on your computer, then click **Open** to upload. Once both documents have been uploaded, click **Save**.
Health Fair Reapplication

During the reapplication period each Spring, laboratories currently holding a health fair permit should review the current data on file with the Department and update such information as appropriate. To review, click on the Health Fair link on the left of the screen.

The Health Fair screen will appear. Review and update information as required. If changes are made, click Save.
Using the links on the left of the screen, review the tests associated with the health fair. Click on a test name.

Review and update the test information as needed. If changes are made, click \textbf{Save}.
Request a Health Fair Permit

To request approval to operate health fairs, click on the **Health Fair** link on the left of the screen.

Click on **Add New** on the left of the screen.

Enter the requested information, click **Next**.
Enter the requested information about the tests to be associated with the health fair, click Save.

Add additional health fair tests by using the Add New link under Health Fair Tests on the left of the screen.
Remove a Health Fair Permit

To remove approval to operate health fairs, click on the Health Fair link on the left of the screen.

Click on Remove on the left side of the screen.
Remove a Health Fair Test

To remove a test from an approved Health Fair permit, click Health Fair of Health Fair Test from the left side of the screen.

Click Remove under Health Fair Tests on the left side of the screen.

Choose the test to remove from the dropdown menu and enter effective date of removal. Click Next.
Miscellaneous

Error Messages

2. Error messages are bordered in red and will appear at the top of the screen after you click Save or Next or Finish, as appropriate. Most text fields without pre-populated information will require a response in order for the page to be saved. Error messages will also prompt you to provide information in the appropriate format, e.g. telephone numbers need to be entered in this format: 123-456-7890.
Pending Changes

3. Saved changes are displayed in the beige/mustard area at the top. It is possible to cancel previously entered changes by selecting one or more of them (click in white box next to name of change) and clicking **Cancel Selected Changes**.

Note: **Pending Changes** are saved so that the reapplication may be continued at a later date/time. To continue a reapplication at a later date/time repeat steps in Steps 1-6 to in **Accessing eCLEP and the Permit Materials Module** of this manual.

If changes are entered but not submitted within one week, the laboratory will begin receiving reminder emails every Monday until the change is either cancelled or submitted.
**ECLEP Manual**

**Request to Re-Open eCLEP**
To re-open the eCLEP system from Read-Only mode to either the Reapplication mode or Open mode, please contact CLEP and please have the four digit PFI number available.

**Telephone:**

518-485-5378

or

**E-mail:**

clep@health.ny.gov

Be sure to have your PFI number when calling or emailing! For emails, please indicate “Re-Open eCLEP Permit Materials” in the subject line.

**HCS Timeout**
For security reasons, there are session timeouts after one hour of inactivity and HCS timeouts after eight hours of total connectivity. These timeouts occur without warning. Timeouts take you back to the login page and force you to re-enter your User ID and Password. If a timeout occurs before you hit Save on the data entry page, you will lose all your data entry. It is recommended to hit Save often while working on long data entry forms.

**Exiting eCLEP**
There are two ways to exit eCLEP:

1. Close your browser by selecting **File** and **Close** from the browser's menu.
2. Click **Logout** at the top right.
   a. The **You are now logged off** message page displays.

**Technical Support**
Technical Support is available for eCLEP and for the NYSDOH Health Commerce System (HCS) in the following areas:

**Help with HCS Enrollment**

For additional assistance contact the Commerce Account Management Unit (CAMU) Help Desk:

(866) 529-1890 (Mon-Fri 8am – 4:45pm)

camu@its.ny.gov

**Help with eCLEP**

For additional assistance contact the Clinical Laboratory Evaluation Program:

- Telephone support at (518) 485-5378
- E-mail support at CLEP@health.ny.gov.
Glossary

Certificate of Qualification (CQ) – a certificate issued by NYSDOH to an individual after the applicant has documented that s/he meets the minimum qualifications as a Laboratory Director set forth in Part 19 of 10NYCRR.

CLEP – Clinical Laboratory Evaluation Program

Delegated Submitter – a person who has been given written authorization by the Laboratory Director to electronically submit facility information on behalf of the Director. A Delegated Submitter will be authorized to enter and submit data electronically using the eCLEP system.

DOH – Department of Health

eCLEP – Electronic Clinical Laboratory Evaluation Program application located on the HCS

Enter Data – Filling out the forms for eCLEP

HCS – Health Commerce System – the Department of Health’s secure Internet network that provides data interchange between health care providers and the NYSDOH.

HCS Coordinator – An individual at the laboratory, designated by the laboratory director, who has the responsibility of requesting additional HCS accounts for data entry individuals. The HCS Coordinator also affiliates HCS User IDs with the laboratory for new users and removes the affiliations for users who have left the laboratory.

Laboratory Director – an individual who is responsible for the administration of the technical and scientific operation of a clinical laboratory or blood bank, including the supervision of procedures, reporting of results and responsibilities specified in Subpart 19.3 of 10 NYCRR and Article 5 Title V Section 571 of the Public Health Law. Such person shall possess a Certificate of Qualification issued pursuant to Part 19 of 10 NYCRR. A Director will be authorized to enter, submit and attest to information entered using the eCLEP system.

NYCRR – New York Codes, Rules and Regulations

NYSDOH – New York State Department of Health


Persistent Data – Data which is saved in the database and displayed in eCLEP, such as

PFI – Permanent Facility Identifier that identifies a laboratory

Submit Data – Confirming that the data entered is accurate and submitted.

User ID – An identification for logging on to the HCS