A. Agenda

For Emergency Adoption
Addition of Section 16.70 – Body Scanners in Local Correctional Facilities

For Adoption
Addition of Section 415.32 – Nursing Home Weekly Bed Census Survey
Amendment of Part 766 – New Requirements for Annual Registration of Licensed Home Care Services Agencies
Amendment of Part 405 and Section 751.5 -- Hospital Policies for Human Trafficking Victims
Amendment of Part 19 – Clinical Laboratory Directors

Program Area
Center for Environmental Health
Division of Nursing Homes
Division of Home and Community Based Services
Office of Primary Care and Health Systems Management
Wadsworth Laboratory

Unit Representative
Alex Damiani
Sheila McGarvey
Margaret Willard
Deirdre Astin
Beverly Rauch

B. Information Announcements

1. Anyone wishing to make oral comments at this meeting should contact the Bureau of Policy and Standards Development by 11:00 A.M. on Wednesday, April 10, at (212) 417-6218 to arrange for placement on the speakers’ list. Please give your name, affiliation, if any and the agenda item(s) you wish to address. To ensure that all commenters have an opportunity to address the Committee, speakers should limit their comments to 3-4 minutes maximum.

2. All meeting attendees including Committee members are requested to sign the Attendance Sheet, which will be circulated in the meeting room.
Pursuant to the authority vested in the Public Health and Health Planning Council and the
Commissioner of Health by sections 201, 225, and 3502 of the Public Health Law, Parts 16 and
89 of Title 10 of the Official Compilation of Codes, Rules and Regulations of the State of New
York are amended, to be effective upon filing with the Secretary of State, to read as follows:

New section 16.70 is added to Part 16 to read as follows:

16.70 Use of Body Scanning.

(a) This section shall not apply in cities having a population of two million or more.

(b) Practitioners licensed under Article 35 of the Public Health Law and unlicensed personnel
employed at a local correctional facility may utilize body imaging scanning equipment that
applies ionizing radiation to humans for purposes of screening inmates committed to such
facility, solely in connection with the implementation of such facility's security program and in
accordance with the provisions of this Part.

(c) Definitions

(1) “Body imaging scanning equipment" or "equipment" means equipment that is specifically
manufactured for security screening purposes and utilizes a low dose of ionizing radiation,
with a maximum exposure per scan equal to or less than 10 µSv (1 mrem), to produce an
anatomical image capable of detecting objects placed on, attached to or secreted within a
person's body. The utilization of body imaging scanning equipment is for purposes of
screening inmates committed to such facility, in connection with the implementation of such
facility's security program.
(2) "Local correctional facility" shall mean a local correctional facility as defined in Correction Law section 2(16).

(3) “Equipment operator” or “operator” means personnel employed at the local correctional facilities that have successfully completed a training course approved by the Department.

(4) "Screening" means the sum of radiation exposures or scans necessary to image objects concealed on all sides of the body as intended by the system design under normal conditions.

(d) Equipment use and installation requirements

(1) Prior to the equipment’s first use on humans at a specific physical location or upon any major repairs that could influence image quality or exposure:

   (i) body imaging scanning equipment purchased or installed at a local correctional facility must be registered with the Department, in accordance with § 16.50 of this Part; and

   (ii) radiation protection survey, shielding evaluation and verification of image usefulness for detecting foreign objects must be completed by a licensed medical physicist.

(2) Equipment must have a clearly marked restricted area and one or more indicators when a scan is in process that is clearly visible to all security screening system operators and anyone approaching the restricted area.

(3) Equipment must be periodically inspected by the Department as described in § 16.10 of this Part.

(4) Equipment must be tested by a licensed medical physicist annually to verify the equipment is operating as designed.

(5) The facility must maintain a policy and procedure manual describing equipment operations, body scanning procedures, records and associated facility policies shall be maintained and
available upon request by the Department. The policy and procedure manual must include the following items:

(i) operating procedures appropriate for the specific equipment and intended scan types;
(ii) policy prohibiting the use of the equipment on individuals who are not inmates;
(iii) policy regarding the determination of pregnancy that has been approved by the jail physician;
(iv) emergency contact information in the event the equipment overexposes any individual or there is equipment related failure that potentially requires service prior to scanning other inmates;
(v) requirements for exposure records to be provided to an inmate upon release or transfer to another facility; and
(vi) exposure per scan for each scan protocol used.

6) Records and documentation of the program operation shall be maintained in accordance with § 16.14 of this Part and shall include, at a minimum, the following:

(i) the number of times the equipment was used on inmates upon intake, after visits, and upon the suspicion of contraband, as well as any other event that triggers the use of such equipment;
(ii) the average, median, and highest number of times the equipment was used on any inmate, with corresponding exposure levels;
(iii) the number of times the use of the equipment detected the presence of drug contraband, weapon contraband, and any other illegal or impermissible object or substance; and
(iv) the number of times an inmate has been scanned.
(e) Exposure limits and reporting requirements

(1) No person other than an inmate of a local correctional facility shall be exposed to the useful beam and then only by an individual that has met the provisions of subdivision (e) of this section.

(2) Limits on the use of equipment and exposure to inmates are:

   (i) no more than fifty percent of the annual exposure limits for non-radiation workers as specified by applicable regulations, not to exceed 0.5 mSv (50 mrem);

   (ii) inmates under the age of eighteen shall not be subject to more than five percent of such annual exposure limits, not to exceed 0.05 mSv (5 mrem); and

   (iii) pregnant women shall not be subject to scanning at any time.

(3) The following events shall be reported to the Department in writing within 30 days:

   (i) incidents or any injuries or illness resulting from the use of such equipment or reported by persons scanned by such equipment; and

   (ii) exposure that exceeds the limits set forth in this Part.

(f) Training Requirements

(1) Every equipment operator shall receive initial operator training, to be provided by the equipment manufacturer or their approved representative, or another source approved by the Department.

(2) The contents of the initial operator training must include radiation safety, equipment operations, exposure and exposure limits for occupational exposed staff and inmates; applicable regulations; and facility policies and procedures.
(3) Initial operator training must be documented and available for review by the Department upon request. Such documentation must include the names of the presenter or sources, attendees, dates and contents of the training.

(4) Every equipment operator shall receive refresher training, to be provided by the equipment manufacturer or their approved representative, or another source approved by the Department. Such training shall meet the requirements listed in paragraphs (1), (2) and (3) of this subdivision and include any changes to the policies and procedures manual or updates to the regulations.

Section 89.30 is amended by adding a new subdivision (c) to read as follows:

(c) A person employed at a local correctional facility, as defined by Correction Law section 2(16), is exempt from licensure as a radiologic technologist when operating body imaging scanning equipment that applies ionizing radiation to humans for purposes of screening inmates committed to such facility, in connection with the implementation of such facility’s security program.
REGULATORY IMPACT STATEMENT

Statutory Authority:

The Department of Health (Department) is required by Public Health Law (PHL) § 201(1)(r) to supervise and regulate the public health aspects of ionizing radiation. PHL § 225(4) authorizes the Public Health and Health Planning Council (PHHPC) to establish, amend and repeal provisions of the State Sanitary Code (SSC), subject to the approval of the Commissioner of Health. PHL §§ 225(5)(p) and (q) and 201(1)(r) authorize PHHPC to establish regulations in the SSC to protect the public from the adverse effects of ionizing radiation.

PHL § 3502 authorizes personnel employed at local correctional facilities to utilize body imaging scanning equipment that applies ionizing radiation to humans for purposes of screening inmates as part of the facilities’ screening program, provided that the use of such equipment is in accordance with regulations promulgated by the Department.

Legislative Objectives:

The legislative intent of PHL §§ 201(1)(r) and 225(5)(p) and (q) is to protect the public from the adverse effects of ionizing radiation. Establishing regulations to ensure safe and effective use of radiation producing equipment is consistent with this legislative objective.

The legislative intent of Article 35 of the PHL is to ensure that when radiation is applied to a human being it is being done appropriately and by a qualified individual. Although in general radiation should only be applied to humans for medical reasons, PHL § 3502 allows correctional facilities to utilize very low dose x-ray equipment for security screening of inmates, while protecting the health of screened inmates.
Needs and Benefits:

Effective January 30, 2019, PHL § 3502(6) permits unlicensed personnel working at local correctional facilities to utilize body imaging scanning equipment that applies ionizing radiation to humans, for purposes of screening inmates committed to such facilities, in connection with the implementation of a facility's security program. Such equipment can be an efficient method of detecting contraband, such as knives, other weapons, and illegal drugs including heroin and opioids, and will enhance the safety of both inmates and correction officers.

These regulations provide protections to the inmates and staff by establishing requirements and controls to ensure appropriate operation of the body scanning imaging equipment. These include testing of the equipment by a licensed medical physicist prior to use and annually thereafter; annual training for equipment operators to ensure proper operation and application; establishment of policies and procedures for use of the equipment; and documentation and inspection requirements to monitor and ensure that inmates are not overexposed to radiation based on the dose limits set forth in the law. The regulations will permit local correctional facilities to take advantage of the enhanced security that body imaging scanning equipment can provide, while minimizing the risk to inmates posed by exposure to ionizing radiation.

Costs:

Costs for the Implementation of, and Continuing Compliance with the Regulation to the Regulated Entity:

The regulations will impose little or no cost to regulated entities. The regulations would only apply to local correctional facilities that voluntarily choose to use body imaging scanning
equipment as part of the facility’s security program. Local correctional facilities that choose to utilize body imaging scanning equipment will be subject to equipment purchase costs; costs to hire a licensed medical physicist to test the body scanning imaging equipment annually, at a cost of approximately $500 per test; administrative costs associated with maintaining records of the use of the equipment; and annual staff training costs. County facilities must register their new x-ray equipment, but they are fee-exempt and will not be charged by the Department for registration or inspections.

**Costs to State and Local Governments:**

These regulations apply only to local correctional facilities operated by county governments that voluntarily choose to use body imaging scanning equipment as part of the facility’s security program. Such facilities will be subject to the costs described above.

**Costs to the Department of Health:**

This regulation will require an increase in inspections of no more than 60 additional facilities out of a total of approximately 11,000 currently registered facilities that are inspected by the Department’s Bureau of Environmental Radiation Protection. The Department will incur costs through preparing and disseminating guidance to the New York State Commission of Correction (NYSCOC) and the NYS Sheriffs Association as well as any local correctional facilities that wish to utilize body imaging scanning equipment. Staff time for registering, inspecting and providing guidance is expected to be handled using existing resources and staff.
Local Government Mandates:

The regulation does not impose any new programs, services, duties or responsibilities upon any county, city, town, village, school district, fire district or other special district. The regulations apply only to local correctional facilities that voluntarily choose to use body imaging scanning equipment as part of the facility’s security program. Such facilities will be subject to the costs described above.

Paperwork:

Local correctional facilities that voluntarily choose to use body imaging scanning equipment as part of the facility’s security program will be required to register the equipment and maintain records related to the policies, procedures and utilization of the equipment.

Duplication:

The regulations do not duplicate, overlap or conflict with any existing federal or state rules or regulations.

Alternatives:

There are no suitable alternatives to the regulations that would meet the requirements of PHL § 3502 while adequately protecting the health of inmates.

Federal Standards:

Not applicable. The operation of radiation producing equipment is regulated by the State only.
Compliance Schedule:

There is no compliance schedule imposed by these regulations, which shall be effective upon filing with the Secretary of State.

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Effect of Rule:

The regulation will only apply to local correctional facilities, operated by county governments, that voluntarily choose to use body imaging scanning equipment as part of the facility’s security program. This regulation will not impact local governments unless they operate such facilities. The regulation will have no impact on small businesses.

Compliance Requirements:

A local correctional facility that chooses to use body imaging scanning equipment as part of the facility’s security program will need to ensure that equipment is installed properly and is operating as designed through licensed medical physicist verification. In addition, the local correctional facility must develop and maintain policies and a procedure manual; provide all personnel who will utilize the equipment with required training; and maintain records of the utilization.

Professional Services:

A local correctional facility that chooses to use body imaging scanning equipment as part of the facility’s security program will be required to have equipment installed by qualified installers for the specific brand of body imaging scanning equipment being used. At facilities with female inmates, the jail physician will be required to develop policies regarding the determination of pregnancy and to update those policies over time as needed. Body scanning
imaging equipment will require annual testing by a licensed medical physicist with an estimated
cost of approximately $500; such testing is also required prior to use of the equipment.

**Compliance Costs:**

A local correctional facility that chooses to use body imaging scanning equipment as part
of the facility’s security program will acquire the equipment based on their own requirements.
Annual compliance costs are expected to be minimal, and will consist of the costs of refresher
training, annual testing by a licensed medical physicist, and record keeping of the inmates
scanned.

**Economic and Technology Feasibility**

This regulation is economically and technically feasible, as these regulations only impose
requirements on local correctional facilities that choose to use body imaging scanning equipment
as part of the facility’s security program. Such facilities will acquire equipment based on their
own requirements and, as described above, ongoing compliance costs are minimal.

**Minimizing Adverse Impact:**

The impact of this regulation is expected to be minimal as these regulations only impose
requirements on local correctional facility that choose to use body imaging scanning equipment
as part of the facility’s security program. To assist such facilities in minimizing any adverse
impact, the Department will provide guidance to NYSCOC and the NYS Sheriffs Association as
well as any local correctional facilities that wish to utilize body imaging scanning equipment.
Small Business and Local Government Participation:

The Department has consulted with the NYS Sheriffs' Association and the New York City Department of Health and Mental Hygiene during the development of the regulations.

Cure Period:

Chapter 524 of the Laws of 2011 requires agencies to include a “cure period” or other opportunity for ameliorative action to prevent the imposition of penalties on the party or parties subject to enforcement under the proposed regulation. This regulatory amendment governing the utilization of body imaging scanning equipment by local correctional facilities does not mandate that local correctional facilities use such equipment. Hence, no cure period is necessary.
RURAL AREA FLEXIBILITY ANALYSIS

Types and Estimated Numbers of Rural Areas:

This rule applies uniformly throughout the state, including rural areas. Rural areas are defined as counties with a population less than 200,000 and counties with a population of 200,000 or greater that have towns with population densities of 150 persons or fewer per square mile. The following 43 counties have a population of less than 200,000 based upon the United States Census estimated county populations for 2010 (http://quickfacts.census.gov).

- Allegany County
- Cattaraugus County
- Cayuga County
- Chautauqua County
- Chemung County
- Chenango County
- Clinton County
- Columbia County
- Cortland County
- Delaware County
- Essex County
- Franklin County
- Fulton County
- Genesee County
- Lewis County
- Livingston County
- Madison County
- Montgomery County
- Oneida County
- Orleans County
- Oswego County
- Otsego County
- Putnam County
- Rensselaer County
- Schoharie County
- Schuyler County
- Seneca County
- St. Lawrence County
- Steuben County
- Sullivan County
- Tioga County
- Tompkins County
- Ulster County
- Warren County
- Washington County
- Wayne County
- Wyoming County
- Yates County
- Schenectady County

The following counties have a population of 200,000 or greater and towns with population densities of 150 persons or fewer per square mile. Data is based upon the United States Census estimated county populations for 2010.

- Albany County
- Broome County
- Dutchess County
- Erie County
- Monroe County
- Niagara County
- Oneida County
- Onondaga County
- Orange County
- Saratoga County
- Suffolk County

Every county in NYS operates a local corrections facility, except Greene and Schoharie counties where the local corrections facilities are currently out of commission. They anticipate
eventually being back in operational status. Approximately 77% of local correctional facilities are in rural areas.

**Reporting, Recordkeeping and Other Compliance Requirements; and Professional Services:**

A local correctional facility that chooses to use body imaging scanning equipment as part of the facility’s security program will need to ensure that equipment is installed properly and is operating as designed through licensed medical physicist verification. In addition, the local correctional facility must develop and maintain policies and a procedure manual; provide all personnel who will utilize the equipment with required training; and maintain records of the utilization.

**Costs:**

A local correctional facility that chooses to use body imaging scanning equipment as part of the facility’s security program will acquire the equipment based on their own requirements. Annual compliance costs are expected to be minimal, and will consist of the costs of refresher training and record keeping of the inmates scanned.

**Minimizing Adverse Impact:**

The impact of this regulation is expected to be minimal as these regulations only impose requirements on local correctional facility that choose to use body imaging scanning equipment as part of the facility’s security program. To assist such facilities in minimizing any adverse
impact, the Department will provide guidance to NYSCOC and the NYS Sheriffs association as well as any local correctional facilities that wish to utilize body imaging scanning equipment.

**Rural Area Participation:**

The Department consulted with the NYS Sheriffs' Association during the development of the regulation. Sheriff’s operate all the local correctional facilities in NYS except for Westchester County and New York City. They indicated that there were no specific issues in this rule that would impact the use body scanning equipment at rural facilities.
STATEMENT IN LIEU OF
JOB IMPACT STATEMENT

A Job Impact Statement for these amendments is not being submitted because it is apparent from the nature and purposes of the amendments that they will not have a substantial adverse impact on jobs and/or employment opportunities.
EMERGENCY JUSTIFICATION

Compliance with the requirements of the State Administrative Procedure Act for filing of a regulation on a non-emergency basis including the requirement for a period of time for public comment cannot be met because to do so would be detrimental to the health and safety of inmates in local correctional facilities.

Effective January 30, 2019, Public Health Law § 3502(6) permits unlicensed personnel working at local correctional facilities to utilize body imaging scanning equipment that applies ionizing radiation to humans for purposes of screening inmates committed to such facility, in connection with the implementation of such facility's security program. Such equipment is intended to be used as an efficient method of detecting contraband, such as knives and other weapons, as well as illegal drugs including heroin and opioids, and will enhance the safety of both inmates and correction officers.

The regulations provide protections to the inmates and staff by establishing requirements and controls to ensure appropriate operation of the body scanning imaging equipment. These include testing of the equipment by a licensed medical physicist prior to use and annually thereafter; annual training for operators of the equipment to ensure proper operation and application; establishment of policies and procedures to guide use of the equipment; and documentation and inspection requirements to monitor and ensure that inmates are not overexposed to radiation based on the dose limits in the law.

Delaying these regulations would prevent local correctional facilities from enhancing security programs through the use of body imaging scanning equipment while minimizing the risks posed to inmates by exposure to ionization.
Pursuant to the authority vested in the Public Health and Health Planning Council, subject to the approval of the Commissioner of Health, by section 2803(2) of the Public Health Law, a new section 415.32 is added to Part 415 of Title 10 (Health) of the Official Compilation of Codes, Rules and Regulations of the State of New York, to be effective upon publication of a Notice of Adoption in the New York State Register:

415.32 Weekly bed census data survey.

(a) Definitions. As used in this section, the following terms shall have the following meanings:

(1) “Communications Directory” (Directory) shall mean a listing of all organizations with access to the HCS, ordered by type, and including the identity of and contact information for individuals at each organization who: (i) perform specific job functions identified by the Department; and/or (ii) have access to perform certain data exchange functions on the HCS.

(2) “HCS Coordinator” shall mean the individual designated by each organization with access to the HCS to be responsible for authorizing and managing accounts and maintaining other key information about the organization’s HCS users.

(3) “Health Commerce System” (HCS) shall mean the Department’s secure Internet portal used for communications and information exchange with organizations including nursing homes and other health care providers or any successor system used for such information exchange as required by the Department.

(4) “Health Electronic Reporting Data System” (HERDS) shall mean the data reporting application on the HCS that houses the Survey or any successor system used for such
reporting as required by the Department.

(5) “Nursing Home Data Reporter” shall mean the name of the role in the Directory that provides access to an individual designated by a nursing home to use HERDS.

(6) “Nursing Home Weekly Bed Census Survey” (Survey) shall mean an electronic survey used by each nursing home to report its bed census to the Department using HERDS.

(7) “Role” shall mean the term used to indicate in the Directory the specific job functions and HCS data exchange functions assigned to individuals by each organization.

(b) Submission of Surveys.

(1) Each nursing home shall complete the Survey on HERDS on a weekly basis by indicating, for each category of bed, the total number of certified or approved beds and the number of those beds that are available. The Survey shall be submitted on a weekly basis by individuals at the nursing home who are assigned to the Nursing Home Data Reporter role within the Directory.

(2) Nursing homes shall report bed census data reflecting the weekly census taken every Wednesday at 12:00 a.m. The nursing home’s designated Nursing Home Data Reporter shall enter and transmit the survey census data to the Department between Wednesday at 12:01 a.m. and the following Tuesday at 11:59 p.m. Instructions for the Survey will be available on the HCS.

(c) Designation of Nursing Home Data Reporters. Nursing homes shall, through their HCS Coordinators, designate a sufficient number of Nursing Home Data Reporters to ensure that the Survey is submitted to the Department in a timely manner.
REGULATORY IMPACT STATEMENT

Statutory Authority:

Public Health Law (PHL) section 2803(2)(a)(v) provides that the Public Health and Health Planning Council shall adopt rules and regulations, subject to the approval of the Commissioner of Health, governing the standards and procedures followed by nursing homes which, at a minimum, must meet federal standards.

Legislative Objectives:

The legislative objective of PHL Article 28, as set forth in PHL section 2800, includes the protection of the health of the residents of New York State through the efficient provision and proper utilization of health services of the highest quality at a reasonable cost. This proposal, which requires nursing homes to submit weekly bed census data to the Department of Health (Department) through the Department’s Health Commerce System, is consistent with that objective. Having current and accurate nursing home bed occupancy data is important in the event of natural disasters and to alert the Department to significant changes in nursing home occupancy, improving the Department’s ability to take appropriate action. While facilities have already been advised administratively that they must submit this data, including the requirement in regulation will improve compliance.
Current Requirements:

The Health Commerce System (HCS), previously known as the Health Provider Network (HPN), is a highly secure, Internet-based, electronic portal for communications and critical data sharing with organizations including nursing homes and other health care providers. Section 400.10 of Title 10 (Health) of the New York Compilation of Codes, Rules and Regulations (NYCRR) requires providers, including nursing homes, to maintain and keep updated an active HPN account.

DAL #09-02, effective April 8, 2009, was issued by the Department to require nursing homes to report weekly bed census data electronically to the Department through the HPN. The DAL provided for such data to be reported each week between Wednesday 8:00 a.m. and Friday 5:00 p.m. In 2013, via a notice sent through the HCS, the Department informed nursing homes that such data should be reported between Wednesday 12:01 a.m. and the following Tuesday at 11:59 p.m.

Needs and Benefits:

It is critical that the Department have accurate nursing home census data including occupancy and availability data by bed type. Natural events such as hurricanes and floods and other emergency events such as extended power outages could cause situations in which some nursing homes may have to transfer their residents to other facilities to ensure their safety. In those situations, the Department must be able to quickly assess the number and location of nursing home residents across the affected area, as well as the number of available beds. Furthermore, the ability to monitor a facility’s current occupancy data
improves the Department’s ability to identify a declining census and proactively take appropriate action.

Despite the current requirement for bed census data reporting, communicated via a DAL and a subsequent HCS notice, the Department often finds itself in the position of having to call some nursing homes repeatedly to obtain this information. This proposed regulation will add a new section 415.32 to Title 10 of the NYCRR to require that nursing homes submit bed census data on a weekly basis by electronically filing the Nursing Home Weekly Bed Census Survey (Survey). This will promote compliance and ensure that the Department has access to essential, current occupancy data as necessary to protect residents.

Accordingly, the proposed regulation provides that the Survey must be submitted via the HCS Health Electronic Response Data System (HERDS) application by a facility staff person assigned a Nursing Home Data Reporter role within the HCS Communications Directory. Nursing homes shall report bed census data reflecting the weekly census taken every Wednesday at 12:00 a.m. The facility’s designated Nursing Home Data Reporter shall enter and transmit the survey census data to the Department between Wednesday at 12:01 a.m. and the following Tuesday at 11:59 p.m. Instructions for the Survey will be available on the HCS. The proposal further requires nursing homes, through their HCS Coordinators, to designate enough Nursing Home Data Reporters to ensure that the facility can submit surveys to the Department as required.
COSTS:

Costs to Private Regulated Parties:

New York State health care facilities are already required by section 400.10 of the NYCRR to have an HCS account to exchange electronic information with the Department. Moreover, nursing homes are already expected to send bed census information to the Department as communicated in the DAL. Therefore, nursing homes should not incur any additional costs related to the electronic submission of bed census information to comply with the proposed regulation.

Costs to Local Government:

This proposal will not impact local governments unless they operate a nursing home, in which case they will be impacted to the same extent as other nursing homes. As previously noted, nursing homes are not expected to incur any additional costs related to the electronic submission of bed census information.

Costs to the Department of Health:

The Department is not expected to incur any additional administrative costs as a result of the proposed regulation. The statewide HCS infrastructure and the mechanisms for nursing home bed census data collection are already in place.

Costs to Other State Agencies:

The proposed regulatory changes will not result in any additional costs to other State agencies.
Local Government Mandates:

This proposed regulation does not impose any new mandates on local governments.

Paperwork:

Nursing homes are already expected to submit bed census information via the HCS. Accordingly, the proposal should not increase paperwork.

Duplication:

This proposed regulation reiterates and strengthens the existing requirement, set forth in the DAL, that nursing homes report census data on a weekly basis to the Department. Moreover, while federal regulations require submission of bed census data to the federal Centers for Medicare and Medicaid Services (CMS) on a quarterly basis, this regulation will ensure that the Department receive this information directly and more frequently.

Alternatives:

There are no other alternatives for the Department to reliably secure current bed census data from nursing homes.

Federal Standards:

Federal regulations require nursing homes to submit quarterly census data to CMS.
**Compliance Schedule:**

These regulations will be effective upon publication of a Notice of Adoption in the New York State Register. The statewide HCS infrastructure and the mechanisms for bed census reporting for nursing homes are already in place. Consequently, regulated parties should be able to comply with the proposed regulation as of its effective date.

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STATEMENT IN LIEU OF
REGULATORY FLEXIBILITY ANALYSIS
FOR SMALL BUSINESSES AND LOCAL GOVERNMENTS

No regulatory flexibility analysis is required pursuant to section 202-(b)(3)(a) of the State Administrative Procedure Act. The proposed rule will not have a substantial adverse impact on small businesses or local governments. Nursing homes that constitute small businesses and local health departments that operate nursing homes, like all other nursing homes, are already required to have an HCS account to exchange electronic information with the Department and report bed census data.
STATEMENT IN LIEU OF
RURAL AREA FLEXIBILITY ANALYSIS

No rural area flexibility analysis is required pursuant to section 202-bb(4)(a) of the
State Administrative Procedure Act. The proposed rule will not have an impact on nursing
homes located in rural areas any differently than in any other areas. Such nursing homes
are already required to have an HCS account to exchange electronic information with the
Department and report bed census data.
STATEMENT IN LIEU OF JOB IMPACT STATEMENT

No job impact statement is required pursuant to section 201-a(2)(a) of the State Administrative Procedure Act. No adverse impact on jobs and employment opportunities is expected as a result of this proposed regulation.
Pursuant to the authority vested in the Public Health and Health Planning Council and the Commissioner of Health by section 3605(7) of the Public Health Law, sections 766.9 and 766.12 of Title 10 of the Official Compilation of Codes, Rules and Regulations of the State of New York are hereby amended to be effective upon publication of a Notice of Adoption in the New York State Register.

Subdivision (n) of § 766.9 is amended to read as follows:

(n) ensure that any franchise agreement complies with the following:

* * *

(4) An agreement which contains elements of both a franchise agreement and a management contract shall be subject to the applicable provisions of this subdivision and subdivision (m) of this section[.]; and

A new subdivision (o) is added to § 766.9 to read as follows and existing subdivision (o) re-lettered (p):

(o) ensure registration of the licensed home care services agency with the commissioner through submission of annual registration forms included in the annual statistical report;

(1) no licensed home care services agency shall be operated, provide nursing services, home health aide services, or personal care services, or receive reimbursement from any source for the provision of such services during any period of time on or after January 1, 2019, unless it has registered for the current period:
(2) a licensed home care services agency that fails to submit a complete and accurate set of all required registration materials by the annual deadline of November 16th is required to pay a fee of $500 for each month or part thereof that the licensed home care services agency is not registered;

(3) a licensed home care services agency that fails to register in the prior year by the deadline of the current year shall not be permitted to register for the upcoming registration period unless it submits any and all unpaid late fees;

(4) the department shall publish a listing of all licensed home care services agencies and their current registration status on its public website;

(5) the department shall institute proceedings to revoke the license of any licensed home care services agency that fails to register for two annual registration periods, whether or not such periods are consecutive; and

(6) the department shall pursue revocation of the license of a licensed home care services agency if it evidences a pattern of late registration over the course of multiple years without justification acceptable to the commissioner.

Subdivision (c) of § 766.12 is amended to read as follows:

(c) The home care services agency shall furnish annually to the department a copy of:

(1) statistical summaries of all health care services, including the type, frequency and reimbursement for services provided, including reimbursement from federal and state governmental agencies, on forms provided by the department;

(2) if a for-profit corporation, a list of the principal stockholders and the number and percent of the total issued and outstanding shares of the corporation held by each, duly certified by the
secretary of the corporation as to completeness and accuracy;

(3) if a not-for-profit corporation, a list of directors, officers and corporate members, if such members number 10 or fewer;

(4) the agency’s registration in a manner prescribed by the department; and

(5) other such records and reports as may be legally required by the department.

* * *
REGULATORY IMPACT STATEMENT

Statutory Authority:

This proposal will implement amendments to Public Health Law (PHL) §§ 3605-a and 3605-b requiring registration of licensed home care services agencies pursuant to Article 36.

Legislative Objective:

Public Health Law Article 36 was intended to promote the quality of home care services provided to residents of New York State and to assure adequate availability as a viable alternative to institutional care. The proposed regulation furthers this objective by developing a system for the Department of Health (Department) to identify agencies that are non-operational and aligns state regulations with the Department’s strategic plan.

Needs and Benefits:

The proposed changes to 10 NYCRR §§ 766.9 and 766.12(c)(4) implement amendments to PHL §§ 3605-a and 3605-b made by Chapter 57 of the Laws of 2018, Part B, §§ 9-c and 9-d, requiring registration of licensed home care services agencies pursuant to PHL.

Annual registration of licensed home care services agencies will allow the Department, on an annual basis, to confirm operational entities in all regions of the state. The registration will confirm the number of agencies providing services in the defined services area and the types of services provided. The information will assist the Department in identifying potential gaps in provider capacity and consumer access to services, and is important as the Department develops a need methodology for licensed home care services agencies. It will also be useful to the Department’s oversight and surveillance functions.
This will be integral in improving the overall quality of services provided to individuals who are receiving home care services.

Just as important, the information obtained from the licensed home care services agency registration will improve consumer access to information about licensed home care services agency availability. The information collected from the registration process will improve the currency and accuracy of provider-related information on the DOH public website, giving consumers meaningful information that can help them identify available options for home care services. Additionally, the public website will identify those agencies who are registered with the Department and those agencies who are not registered with the department, indicating their compliance with 10 NYCRR § 766.9.

To comply with the registration requirement, licensed home care services agencies will need to complete a section that will be added to the existing annual statistical report. These must be submitted during the annual data collection period, which commences in August of the preceding year of the registration deadline and ends by November 16th.

The proposed changes will provide a benefit to current licensed home care services agencies who complete the registration as required, as they will be listed on the public website as being currently registered and active.

Costs to Regulated Parties:

The regulated parties (providers) are not expected to incur any additional costs as a result of the proposed rule change. There are no additional costs to local governments for the implementation of and continuing compliance with this amendment. There are no additional costs to the Department of Health as a result of the proposed rule change.
Local Government Mandates:

The proposed amendment does not impose any new programs, services, duties or responsibilities upon any county, city, town, village, school district, fire district or other special district. The registration will be incorporated as part of the annual statistical reports already required to be submitted by licensed home care services agencies. Therefore, the new state regulation will require county operated agencies to complete one additional form.

Paperwork:

The registration will be incorporated as part of the annual statistical reports already required to be submitted by licensed home care services agencies. Therefore, the new state regulation will require one additional form to be completed.

Duplication:

The proposed rule is not duplicative of any known rules or regulations.

Alternatives:

There are no alternatives to this proposal, which is necessary to implement a legislative enactment requiring licensed home care services agencies to register annually with the Department.

Federal Standards:

This amendment does not exceed any minimum standards of the federal government for the same or similar subject areas.
Compliance Schedule:

There are no significant actions which are required by the affected providers to comply with the amendments, as the amendments ensure conformance with expectations that were already in effect. Those licensed home care services agencies who are operational should already be in compliance with the required annual statistical reports and should be readily able to comply. The registration will be incorporated as part of the annual statistical reports already required to be submitted by licensed home care services agencies. Therefore, the new state regulation will require one additional form to be completed. A licensed home care services agency that fails to submit a complete and accurate set of all required registration materials by the annual deadline of November 16th, established by the Commissioner of Health, is required to pay a fee of $500 for each month or part thereof that the licensed home care services agency is not registered. No licensed home care services agency shall be operated, provide nursing services, home health aide services, or personal care services, or receive reimbursement from any source for the provision of such services during any period of time on or after January 1, 2019, unless it has registered for the current period. The regulations will be effective upon publication of a Notice of Adoption in the New York State Register.

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REGULATORY FLEXIBILITY ANALYSIS FOR SMALL BUSINESSES AND LOCAL GOVERNMENTS

Effect of Rule:

Licensed home care services agencies, including those operated by county health departments, provide health services in the home pursuant to Public Health Law Article 36. There are currently 1,083 licensed operators providing home care services at 1,475 licensed sites. Local governments will not be affected by this rule except to the extent that they operate licensed home care services agencies; nor will small businesses be impacted in their routine cost of conducting business.

Compliance Requirements:

Regulated parties are expected to be in compliance beginning on and after January 1, 2019. The proposed regulations will implement the new registration requirement for licensed home care services agencies, which will be carried out through existing reporting mechanisms. The registration process is a new requirement; however, the registration process will be incorporated with existing statistical data collection requirements for licensed home care services agencies which are required annually. Therefore, compliance requirements are minimal.

The Department does not intend to publish a small business regulation guide in connection with this regulation. Although a number of licensed home care services agencies are small businesses, the impact is expected to be minimal. Additional guidance will be posted on the web as needed after the regulation is promulgated.
**Professional Services:**

No additional professional staff are expected to be needed as a result of the regulations. Record keeping and compliance requirements could be handled by existing staff, as it is the expectation that the administrator complete the registration.

**Compliance Costs:**

There are no capital costs associated with these proposed rules. Any costs are already incurred by agencies under the existing regulations.

**Economic and Technological Feasibility:**

The Department has considered feasibility and believes there will be minimal, if any, economic and technological impact. The registration will be incorporated as part of the annual statistical reports already required to be submitted by licensed home care services agencies. Therefore, the new state regulation should not affect the routine cost of doing business, unless agencies have been non-compliant with existing requirements.

**Minimizing Adverse Impact:**

While the Department has considered the options of State Administrative Procedure Act (SAPA) § 202-b(1) in developing this rule, flexibility does not exist for any particular entity since the new requirements are consistent with requirements that are already in effect.

**Small Business and Local Government Participations:**

The Department will meet the requirements of SAPA § 202-b(6) in part by publishing a
notice of proposed rulemaking in the State register with a comment period. The Department has
not solicited input prior to publication as the proposed amendments are required by statute, do
not change existing procedures in any substantive manner and will, therefore, have no
deleterious effect on small businesses and local governments.

**Rules that Either Establish or Modify a Violation or Penalties Associated with a Violation:**

A licensed home care services agency which fails to submit a complete and accurate set
of all required registration materials by the deadline established by the Commissioner shall be
required to pay a fee of $500 for each month or part thereof that the licensed home care services
agency is in default. The statute allows for the LHCSA to register at any time, however, the fines
will continue to be incurred.

A licensed home care services agency that failed to register in the prior year by the
deadline of the current year shall not be permitted to register for the upcoming registration period
unless it submits any unpaid late fees.

A licensed home care services agency is prohibited from providing nursing services,
home health aide services, or personal care services, or receive reimbursement from any source
for the provision of such services during any period of time on or after January 1, 2019, unless it
has registered with the Department.

The Department shall institute proceedings to revoke the license of any licensed home
care services agency that fails to register for two annual registration periods, whether or not such
periods are consecutive. The Department shall have the discretion to pursue revocation of the
license of a licensed home care services agency on grounds that it evidences a pattern of late
registration over the course of multiple years.
The registration will be incorporated as part of the annual statistical reports already required to be submitted by licensed home care services agencies. Therefore, the new state regulation will require one additional form to be completed. A licensed home care services agency that fails to submit a complete and accurate set of all required registration materials by the annual deadline of November 16th is required to pay a fee of $500 for each month or part thereof that the licensed home care services agency is not registered. No licensed home care services agency shall be operated, provide nursing services, home health aide services, or personal care services, or receive reimbursement from any source for the provision of such services during any period of time on or after January 1, 2019, unless it has registered for the current period.
STATEMENT IN LIEU OF RURAL AREA FLEXIBILITY ANALYSIS

All counties in New York State (NYS) have rural areas with the exception of seven (7) downstate counties. Approximately 80% of licensed home care services agencies are licensed to serve counties with rural areas. No rural area flexibility analysis is required pursuant to § 202-bb(4)(a) of SAPA. The proposed amendment does not impose an adverse impact on facilities in rural areas and it does not impose additional reporting, record keeping or other compliance requirements on facilities in rural areas. The proposed amendment to require licensed home care agencies to complete registration seeks information regarding operational agencies and to assure home care availability in rural areas as an alternative to institutional care.
STATEMENT IN LIEU OF

JOB IMPACT STATEMENT

A Job Impact Statement for these amendments is not being submitted because it is apparent from the nature and purposes of the amendments that they will not have a substantial adverse impact on jobs and/or employment opportunities.
Pursuant to the authority vested in the Public Health and Health Planning Council and Commissioner of Health by sections 2803(2)(a) and 2805-y(4) of the Public Health Law, sections 405.9, 405.18, 405.19, 405.20, 407.5, and 751.5 of Title 10 of the Official Compilation of Codes, Rules and Regulations of the State of New York (NYCRR) are hereby amended, to be effective upon filing of a Notice of Adoption in the New York State Register:

Subparagraph (ii) of paragraph (11) of subdivision (b) of section 405.9 of Title 10 is amended to read as follows:

(ii) If a patient eligible for transfer to a hospital operated by the Veteran's Administration requests such transfer, hospital staff shall make such arrangements. Transfer shall be effected in accordance with paragraph [(g)(7) (h)(7)] of this section.

Subdivision (g) is relettered as (h) and a new subdivision (g) is added to section 405.9 of Title 10 to read as follows:

(g) Human Trafficking. The hospital shall provide for the identification, assessment, and appropriate treatment or referral of individuals who are suspected to be human trafficking victims, as that term is defined in section 483-aa of the Social Services Law and used in Article 10-D of the Social Services Law. The hospital shall establish and implement written policies and procedures, which shall apply to all service units of the hospital and, at a minimum, shall meet the following requirements:
(1) Policies and procedures shall provide for the identification, assessment, and appropriate treatment or referral of individuals who are suspected to be human trafficking victims;

(2) In the case of individuals who are suspected to be human trafficking victims and are under eighteen years old, policies and procedures shall provide for the reporting of such persons as an abused or maltreated child if required under Title 6 of Article 6 of the Social Services Law;

(3) The hospital shall inform individuals who are suspected to be human trafficking victims of services that may be available, including those referenced in Article 10-D of the Social Services Law. Referrals also may be made to other health care providers, appropriate state agencies, and/or other providers of services as appropriate. Such information may be provided verbally and/or in writing as appropriate;

(4) The hospital shall post the human trafficking hotline poster issued by the National Human Trafficking Resources Center, or a variation of such poster created by the Office of Temporary and Disability Assistance (OTDA) consistent with section 483-ff of the Social Services Law, whichever OTDA makes available on its website. Posters shall be placed in conspicuous locations near primary public entrances and where other posters and notices are posted; and

(5) The hospital shall establish and implement training, which may be incorporated into current training programs, for all individuals licensed or certified pursuant to Title 8 of the Education Law who provide direct patient care, and for all security personnel, regarding the policies and procedures established pursuant to this subdivision. Such training shall include training in the
recognition of indicators of a human trafficking victim and the responsibilities of such personnel in dealing with persons suspected as human trafficking victims.

Subdivision (h) of section 405.9 of Title 10 is relettered as (i) and subparagraph (ii) of paragraph (7) of the former subdivision (g), now relettered as subdivision (h), of section 405.9 of Title 10 is amended to read as follows:

(ii) Patients discharged from the hospital by their attending practitioner shall not be permitted to remain in the hospital without the consent of the chief executive officer of the hospital except in accordance with provisions of subdivision [(h)] (i) of this section.

Subparagraph (vi) of paragraph (2) of subdivision (b) of section 405.18 of Title 10 is amended to read as follows:

(vi) In accordance with the provisions of section [405.9(g)] 405.9(h) of this Part, rehabilitation therapy staff shall work with the attending practitioner, the nursing staff, other health care providers and agencies as well as the patient and the family, to the extent possible, to assure that all appropriate discharge planning arrangements have been made prior to discharge to meet the patient's identified needs.

New paragraph (6) is added to subdivision (c) of section 405.19 of Title 10 to read as follows, and existing paragraphs (6) through (10) are renumbered (7) through (11):
(6) The emergency service shall provide for the identification, assessment, and appropriate
treatment or referral of individuals who are suspected to be human trafficking victims, as
described in subdivision (g) of section 405.9 of this Part.

Paragraph (5) of subdivision (c) of section 405.20 of Title 10 is amended, paragraph (6) is
renumbered (7) and a new paragraph (6) is added to read as follows:

(5) identification, assessment, and referral of individuals with documented substance use
disorders or who appear to have or be at risk for substance use disorders, as that term is defined
in section 1.03 of the Mental Hygiene Law, as described in subdivision (f) of section 405.9 of
this Part; [and]

(6) compliance with the human trafficking provisions pertaining to the identification, assessment,
and appropriate treatment or referral of individuals who are suspected to be human trafficking
victims, as described in subdivision (g) of section 405.9 of this Part; and

Paragraph (6) of subdivision (b) of section 407.5 of Title 10 is amended to read as follows:

(6) Discharge/transfer. Hospitals shall comply with the provisions of paragraph (1) of
subdivision [(h)][(i) of section 405.9 of this Title concerning discharge/transfer. In addition,
PCHs and CAHs shall comply with the following:

*  *  *
A new paragraph (8) is added to subdivision (a) of section 751.5 of Title 10, and paragraphs (8) through (16) are renumbered (9) through (17), to read as follows:

(8) the identification, assessment, and appropriate treatment or referral of individuals who are suspected to be human trafficking victims, as that term is defined in section 483-aa of the Social Services Law and used in Article 10-D of the Social Services Law; training in the recognition of indicators of a human trafficking victim and the responsibilities of such personnel in dealing with persons suspected as human trafficking victims, the reporting of individuals who are suspected to be human trafficking victims and are under eighteen years old as abused or maltreated children if required under Title 6 of Article 6 of the Social Services Law; and the posting of the human trafficking hotline poster issued by the National Human Trafficking Resources Center, or a variation of such poster created by the Office of Temporary and Disability Assistance (OTDA) consistent with section 483-ff of the Social Services Law, whichever OTDA makes available on its website, in conspicuous locations near primary public entrances and where other posters and notices are posted;
REGULATORY IMPACT STATEMENT

Statutory Authority:

Public Health Law (PHL) § 2803(2)(a) authorizes the Public Health and Health Planning Council (PHHPC) to adopt and amend rules and regulations, subject to the approval of the Commissioner of Health (Commissioner), to implement PHL Article 28 and establish minimum standards for health care facilities.

PHL § 2805-y(4) authorizes the Commissioner to issue regulations, in consultation with the Office of Temporary and Disability Assistance (OTDA) and the Office of Children and Family Services (OCFS), to implement the section, which requires “subject facilities” (general hospitals, public health centers, diagnostic centers, treatment centers, or outpatient departments) to develop, maintain, and train staff in policies and procedures for the identification, assessment, treatment, and referral of human trafficking victims.

Legislative Objectives:

This proposal will implement PHL § 2805-y, added by Chapter 408 of the Laws of 2016, to require general hospitals and diagnostic and treatment centers (D&TCs), which encompass the entities referenced as “subject facilities” in the statute, to establish policies and procedures for the identification, assessment, treatment, and referral of human trafficking victims and to train staff in such policies and procedures. The policies and procedures must include the posting of a human trafficking hotline poster consistent with the objectives of Social Services Law (SSL) § 483-ff, added by Chapter 311 of the Laws of 2016.

As explained below, a 2007 law established new crimes related to human trafficking and made various health and social services available to victims. More recent enactments reflect a
legislative desire to combat this growing issue by requiring that general hospitals and D&TCs adopt procedures to identify victims, treat and/or refer them for other services as appropriate, and post a hotline number in public areas where victims may be present.

**Needs and Benefits:**

The scale of the human trafficking problem constitutes a public health crisis impacting people and their families throughout New York. Legislation enacted in 2007 greatly expanded the tools available to address the issue, but human trafficking nevertheless remains prevalent. A recent study found that 69 percent of survivors surveyed indicated they had accessed health care services at some point during their trafficking. Chapter 408 of the Laws of 2016 recognized this additional opportunity to support human trafficking victims by requiring general hospitals and D&TCs to establish and implement policies to identify, assess, and treat or refer individuals suspected of being victims. Similarly, Chapter 311 of the Laws of 2016 sought to publicize information about resources for human trafficking victims in public areas where victims are likely to be present, including hospitals and clinics.

The New York State Anti-Trafficking Statute, Chapter 74 of the Laws of 2007, was enacted in light of the growing problem of human trafficking for “forced labor, involuntary domestic servitude, or sexual exploitation.” The sponsor’s memorandum noted that victims – frequently children – may be trafficked within or into the United States and New York often serves as a hub of such activity. Among other things, the law added Penal Law §§ 135.35 and 230.34 to establish the crimes of labor trafficking and sex trafficking, respectively.

The 2007 enactment, as amended in 2015, also added SSL Article 10-D providing for services to human trafficking victims. SSL § 483-aa(a) defines a “human trafficking victim” as a
victim of sex trafficking or labor trafficking under the above-referenced Penal Law sections. SSL § 483-bb provides that OTDA may contract with non-governmental entities to make available services, including case management, emergency temporary housing, health care, mental health counseling, and drug addiction screening and treatment, to “pre-certified” human trafficking victims. SSL § 483-aa(b) defines “pre-certified victim of human trafficking” as a person with a pending application for federal certification as a victim of a severe form of trafficking in persons as defined in section 7105 of title 22 of the United States Code (Trafficking Victims Protection) but has not yet obtained such certification, or a person who has reported a crime to law enforcement and it reasonably appears to law enforcement that the person is such a victim.

SSL § 483-cc sets forth procedures for confirming an individual’s status as a human trafficking victim. Under that section, a law enforcement agency or district attorney’s office that encounters a person who reasonably appears to be a human trafficking victim must notify OTDA and the Division of Criminal Justice Services (DCJS) that the individual may be eligible for services under SSL Article 10-D. To activate this process, a law enforcement agency or district attorney’s office must use the New York State Referral of Human Trafficking Victim Form available on the OTDA website at http://otda.ny.gov/programs/bria/trafficking.asp. Providers of social or legal services designated by an applicable state agency (OTDA, the Office for the Prevention of Domestic Violence, or the Office of Victim Services) that encounter a person who reasonably appears to be a human trafficking victim may submit the form if the individual consents.

Upon receipt of the form, DCJS, in consultation with OTDA and the referring agency or office, assesses whether the person meets the criteria for certification as a victim of a severe form
of trafficking in persons as defined in 22 U.S.C. § 7105 or appears to be otherwise eligible for any federal, state or local benefits and services. If so, OTDA reports such finding to the victim and the referring entity and may assist the victim in receiving services. This finding is referred to as “confirmation” as a victim of human trafficking.

Chapter 311 of the Laws of 2016 added a new SSL § 483-ff requiring OTDA to make available on its website the hotline poster issued by the National Human Trafficking Resources Center (NHTRC) or a version created by OTDA. The section provides for OTDA to consult with other state agencies to encourage that the posters be placed where human trafficking victims may be present, including hospitals and urgent care centers, in conspicuous places near primary public entrances or where posters and notices are customarily placed.

Chapter 408 of the Laws of 2016 added new PHL § 2805-y to require “subject facilities” to establish and implement policies and procedures pertaining to victims of human trafficking. New PHL § 2805-y(1) defines key terms such as “subject facilities,” defined to mean general hospitals, public health centers, diagnostic centers, treatment centers or outpatient departments, and provides that the requirements of PHL § 2805-y applies to all service units that include emergency services, pediatrics, obstetrics and gynecology, orthopedics, internal medicine, family medicine, radiology, surgery, psychiatry and dental services to the extent the facility maintains a dental clinic, center, or department on site of the facility.

New PHL § 2805-y(2) requires subject facilities to establish and implement written policies and procedures for the identification, assessment, and appropriate treatment or referral of persons suspected of being human trafficking victims, as that term is defined by SSL § 483-aa. Further, policies and procedures must provide for referral of human trafficking victims under the
age of 18 to the Statewide Central Register of Child Abuse and Maltreatment (SCR) established pursuant to SSL Title 6, Article 6 if required by that law.

New PHL § 2805-y(3) also requires subject facilities to require all “subject facility personnel” – defined as nursing, medical, social work and other clinical care personnel as well as security personnel – to complete training regarding such policies and procedures. This must include training in the recognition of indicators of a human trafficking victim and the responsibilities of such personnel in dealing with persons suspected of being victims.

Finally, new PHL § 2805-y(4) authorizes the Commissioner to identify organizations or providers that could provide training for general hospitals consistent with the new provisions. The subdivision also authorizes the issuance of regulations, in consultation with OTDA and OCFS, as necessary to carry out the new section.

Consistent with these requirements, this proposal will amend 10 NYCRR §§ 405.9, 405.19, 405.20, and 751.5 to require general hospitals and D&TCs to establish written policies and procedures for the identification, assessment, and appropriate treatment or referral of individuals who are or appear to be a human trafficking victim and train staff in such policies and procedures. Referrals may be provided verbally and/or in writing as appropriate. Policies, procedures and training must include information about the referral process overseen by OTDA and DCJS. While the proposed regulations do not mandate that hospitals and D&TCs use the New York State Referral of Human Trafficking Victim Form, they are strongly encouraged to do so when they can secure the victim’s consent.

In addition, there are other sources of assistance that the victim can be referred to, such as the NHTRC hotline, that provide confidential assistance to those victims who do not feel comfortable being referred to OTDA and DCJS. Further, the proposed regulation requires
posting of the NHTRC hotline poster or other variation developed by OTDA in conspicuous locations, which is consistent with the objectives of SSL § 483-ff. The poster designated for such purpose by OTDA is available at http://otda.ny.gov/programs/bria/trafficking.asp.

Under the law, policies and procedures and training must also include the reporting of human trafficking victims under 18 years of age to the SCR if required under SSL Title 6, Article 6. Medical and hospital personnel already serve as mandated reporters who are required to make reports to the SCR if they suspect child abuse or maltreatment. As reiterated by Chapter 408, if an individual appears to be a human trafficking victim under the age of 18, mandated reporters in hospitals and D&TCs must make a report if required under SSL Title 6, Article 6.

COSTS:

Costs to Private Regulated Parties:

While current regulations do not specifically refer to individuals who are human trafficking victims, general hospitals and D&TCs are already required to have written policies and procedures for various operational requirements, train staff in such policies and procedures, and refer patients to appropriate follow-up care. The proposed regulations do require additional effort to ensure that the policies and training include the identification, assessment, and appropriate treatment or referral of individuals who are suspected victims of human trafficking, consistent with PHL § 2805-y. However, the additional costs are expected to be minimal given the existing training infrastructure in general hospitals and D&TC’s. In addition, these efforts are expected to assist individuals in obtaining treatment critical for their overall health and well-being and could help such individuals avoid future emergency room visits and hospital admissions. Therefore, the cost of implementing the proposed regulations is likely to be offset
by a reduction in care provided at no, or low, cost to victims of human trafficking.

**Costs to Local Government:**

This proposal will not impact local governments unless they operate a general hospital or a D&TC, in which case the impact would be the same as outlined above for private parties.

**Costs to the Department of Health:**

The proposed regulatory changes will not result in any additional costs to the Department.

**Costs to Other State Agencies:**

The proposed regulatory changes may result in additional costs to other state agencies if referrals increase and more victims access available services, but this would be consistent with the objectives of the statute. OTDA, OCFS, and DCJS have existing materials related to human trafficking available on their websites.

**Local Government Mandate:**

The proposed regulations do not impose any new programs, services, duties or responsibilities upon any county, city, town, village, school district, fire district or other special district, unless such local government operates a hospital or D&TC.
**Paperwork:**

General hospitals and D&TCs are already required to establish written policies and procedures related to various operational requirements, train staff, and refer patients. Therefore, the proposed regulations should not significantly increase their paperwork.

**Duplication:**

Existing regulations require hospitals to make appropriate referrals for patients to a variety of services, but do not specifically reference human trafficking victims. There otherwise are no relevant State or federal regulations which duplicate, overlap or conflict with the proposed regulations.

**Alternatives:**

There are no alternatives to the proposed regulations related to hospital policies and procedures, which are necessary to implement the provisions of PHL § 2805-y, added by Chapter 408 of the Laws of 2016, and SSL § 483-ff, added by Chapter 311 of the Laws of 2016.

**Federal Standards:**

There are currently no federal requirements for hospitals to adopt policies and procedures for the identification, assessment, treatment, and referral of human trafficking victims.

**Compliance Schedule:**

The regulations will be effective upon publication of a Notice of Adoption in the New York State Register.
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REGULATORY FLEXIBILITY ANALYSIS
FOR SMALL BUSINESSES AND LOCAL GOVERNMENTS

Effect of Rule:

The proposed regulatory provisions related to human trafficking will apply to all general hospitals and diagnostic and treatment centers (D&TCs) in New York State. This proposal will not impact local governments or small business unless they operate a general hospital or D&TC, in which case the requirements will be the same as for those entities.

Compliance Requirements:

These regulations will require general hospitals and D&TCs to develop, maintain and disseminate written policies and procedures for the identification, assessment, and appropriate treatment or referral of victims of human trafficking. These facilities will be required to train their licensed and certified clinical staff members as well as security staff members in such policies and procedures. In addition, the policies must incorporate the posting of a poster with human trafficking hotline information, available on the Office of Temporary and Disability Assistance website, in conspicuous places.

Professional Services:

While the current regulations do not specifically refer to individuals who are human trafficking victims, general hospitals and D&TCs are already required to establish written policies and procedures related to various operational requirements, train staff in such policies and procedures and refer patients to appropriate follow-up care. As such, the Department
anticipates that no additional professional services will be required for general hospitals and D&TCs to comply with this proposed regulation.

**Compliance Costs:**

While the current regulations do not specifically refer to individuals who are or may be victims of human trafficking, general hospitals and D&TCs are already required to have written policies and procedures related to various operational requirements, train staff in such policies and procedures and refer patients to appropriate follow-up care. The proposed regulations do require additional effort to ensure that the policies and training include the identification, assessment and referral of individuals who are suspected victims of human trafficking, consistent with the requirements of PHL § 2805-y. However, the additional costs are expected to be minimal given the existing training infrastructure in general hospitals and D&TC’s. In addition, these efforts are expected to assist individuals in obtaining treatment critical for their overall health and well-being and could help such individuals avoid future emergency room visits and hospital admissions. Therefore, the cost of implementing the proposed regulations is likely to be offset by a reduction in care provided at no, or low, cost to victims of human trafficking.

**Economic and Technological Feasibility:**

This proposal is economically and technically feasible. Although existing regulations do not specifically refer to human trafficking victims, general hospitals and diagnostic and treatment centers are already required to establish written policies and procedures related to various operational requirements, train staff in such policies and procedures and refer patients to appropriate follow-up care.
Minimizing Adverse Impact:

The impact of this proposal is expected to be minimal as general hospitals and D&TCs are already required to have written policies and procedures related to various operational requirements, train staff in such policies and procedures and refer patients to appropriate follow-up care.

To assist hospitals and D&TCs with the development of their policies, procedures and training materials, several state agencies have provided resources that are free of charge to the public. For example:

- A course entitled "NYSDOH Human Trafficking Awareness Training," available on the Department's NYLearnsPH.com Learning Management System at [https://www.nylearnsph.com/public](https://www.nylearnsph.com/public);

In addition, these efforts are expected to assist individuals in obtaining treatment critical for their overall health and well-being and could help such individuals avoid future emergency room visits and hospital admissions. Therefore, the cost of implementing the proposed regulations is likely to be offset by a reduction in care provided at no, or low, cost to victims of human trafficking.

Small Business and Local Government Participation:

Organizations representing health care providers and other stakeholders, including organizations whose members include general hospitals or diagnostic and treatment centers that
are operated by local governments or that constitute small businesses, were consulted on the proposed regulations.

**Cure Period:**

Chapter 524 of the Laws of 2011 requires agencies to include a “cure period” or other opportunity for ameliorative action to prevent the imposition of penalties on a party subject to enforcement when developing a regulation or explain in the Regulatory Flexibility Analysis why one is not included. As this proposed regulation does not create a new penalty or sanction, no cure period is necessary.
RURAL AREA FLEXIBILITY ANALYSIS

Types and Estimated Numbers of Rural Areas:

This rule applies uniformly throughout the state, including rural areas. Rural areas are defined as counties with a population less than 200,000 and counties with a population of 200,000 or greater that have towns with population densities of 150 persons or fewer per square mile. The following 43 counties have a population of less than 200,000 based upon the United States Census estimated county populations for 2010 (http://quickfacts.census.gov).

Approximately 17% of small health care facilities are located in rural areas.

Allegany County  Greene County  Schoharie County
Cattaraugus County  Hamilton County  Schuyler County
Cayuga County  Herkimer County  Seneca County
Chautauqua County  Jefferson County  St. Lawrence County
Chemung County  Lewis County  Steuben County
Chenango County  Livingston County  Sullivan County
Clinton County  Madison County  Tioga County
Columbia County  Montgomery County  Tompkins County
Cortland County  Ontario County  Ulster County
Delaware County  Orleans County  Warren County
Essex County  Oswego County  Washington County
Franklin County  Otsego County  Wayne County
Fulton County  Putnam County  Wyoming County
Genesee County  Rensselaer County  Yates County
Schenectady County

The following counties have a population of 200,000 or greater and towns with population densities of 150 persons or fewer per square mile. Data is based upon the United States Census estimated county populations for 2010.

Albany County  Monroe County  Orange County
Broome County  Niagara County  Saratoga County
Dutchess County  Oneida County  Suffolk County
Erie County  Onondaga County
There are 47 general hospitals, approximately 90 diagnostic and treatment centers, 159 nursing homes, and 92 certified home health agencies in rural areas.

**Reporting, Recordkeeping, Other Compliance Requirements and Professional Services:**

The proposed regulation is applicable to those general hospitals and diagnostic and treatment centers located in rural areas and is expected to impose only minimal costs upon hospitals, which are already required to establish written policies and procedures related to various operational requirements, train staff in such policies and procedures and refer patients to appropriate follow-up care. Because the proposed regulatory requirements can be incorporated into existing processes, they are not expected to substantially increase the administrative burden on these entities.

**Costs:**

While the current regulations do not specifically refer to individuals who may be victims of human trafficking, general hospitals and diagnostic and treatment centers (D&TCs) are already required to have written policies and procedures related to various operational requirements, train staff in such policies and procedures and refer patients to appropriate follow-up care. The proposed regulations do require additional effort to ensure that the policies and training include the identification, assessment and referral of individuals who are suspected victims of human trafficking, as well as the provision of information related to appropriate services, consistent with the requirements of the statute. However, the additional costs are expected to be minimal given the existing training infrastructure in general hospitals and D&TC’s. In addition, these efforts are expected to assist individuals in obtaining treatment
critical for their overall health and well-being and could help such individuals avoid future emergency room visits and hospital admissions. Therefore, the cost of implementing the proposed regulations is likely to be offset by a reduction in care provided at no, or low, cost to victims of human trafficking.

Minimizing Adverse Impact:

The impact of this proposal is expected to be minimal as general hospitals and D&TCs are already required to have written policies and procedures related to various operational requirements, train staff in such policies and procedures and refer patients to appropriate follow-up care.

To assist hospitals and D&TCs with the development of their policies, procedures and training materials, several state agencies have provided resources that are free of charge to the public. For example:


In addition, these efforts are expected to assist individuals in obtaining treatment critical for their overall health and well-being and could help such individuals avoid future emergency room visits and hospital admissions. Therefore, the cost of implementing the proposed regulations is likely to be offset by a reduction in care provided at no, or low, cost to victims of
human trafficking.

**Rural Area Participation:**

Organizations that include as members general hospitals and D&TCs located in rural areas were consulted on the proposed regulations.
STATEMENT IN LIEU OF JOB IMPACT STATEMENT

No job impact statement is required pursuant to section 201-a(2)(a) of the State Administrative Procedure Act. No adverse impact on jobs and employment opportunities is expected as a result of these proposed regulations.
SUMMARY OF EXPRESS TERMS

Pursuant to the authority vested in the Public Health and Health Planning Council and the Commissioner of Health by section 573 of the Public Health Law, Part 19 of Title 10 (Health) of the Official Compilation of Codes, Rules and Regulations of the State of New York is amended, to be effective upon publication of a Notice of Adoption in the New York State Register, as follows:

Section 19.1 is amended to include definitions for “assistant director,” “board certified,” “earned doctoral degree,” “training,” and “experience.” The definitions of “acceptable laboratory” and “category” are also revised and clarified. Section 19.1 is further revised to expressly recognize physicians and dentists who are licensed in the countries in which they practice as being able to qualify as directors or assistant directors of clinical laboratories or blood banks.

Section 19.2 is amended to recognize additional accrediting boards for purposes of certifying that applicants meet the educational and training requirements needed to be a director or assistant director of a clinical laboratory or blood bank.

Section 19.3 is amended to provide the Department more flexibility in updating the certificate of qualification categories. Amendments to this section will also allow the Department to issue certificates of qualification with limitations based on an applicant’s specific experience. In addition, this section is amended to include additional director responsibilities, such as ensuring staff competency, specifying in writing the responsibilities and duties of all laboratory personnel, having standard operating procedure manuals, and participating in acceptable proficiency testing.
Section 19.4 is amended for clarity and to remove references to New York City laboratory permits, which are obsolete.
Pursuant to the authority vested in the Public Health and Health Planning Council and the Commissioner of Health by section 573 of the Public Health Law, Part 19 of Title 10 (Health) of the Official Compilation of Codes, Rules and Regulations of the State of New York is amended, to be effective upon publication of a Notice of Adoption in the New York State Register, as follows:

19.1 Definitions.

(a) [Clinical laboratory director] Director means the individual responsible for administration of the technical and scientific operation of a clinical laboratory or blood bank, including supervision of test procedures, the reporting of results, and the duties and responsibilities specified in section 19.3 of this Part. If a clinical laboratory or blood bank employs more than one director, the laboratory owner(s) shall designate in writing one such individual as the director of record for the laboratory.

(b) Assistant director means a director who has been designated by the owner(s) of the laboratory as having shared responsibility with a director for the technical and scientific operation of the clinical laboratory or blood bank in one or more categories and/or subcategories.

[(b)] (c) Acceptable laboratory means [a clinical laboratory or blood bank of a hospital, health department, university, medical research institution, independent clinical laboratory or blood bank, or other facility providing equivalent training and/or experience in patient specimen testing, which has a director who meets or would meet the requirements of this Part and which meets or would meet the commissioner’s standard as outlined in Part 58 of this Title.] a facility, operating lawfully, that meets the definition of a clinical laboratory or blood bank as defined in Section 571 of the Public Health Law and which has a director who meets or would meet the
requirements of this Part, including the anatomic and clinical pathology facilities of a hospital or health department, a clinical testing unit of a university or medical research institution, an independent clinical laboratory or blood bank, a privately operated forensic testing laboratory, or a facility providing training and/or experience in the testing of human specimens.

[(c)] (d) Accredited means having the approval (accreditation) conferred on schools, institutions or programs by an accrediting agency or association recognized by the United States Secretary of Education and verified as such by the [commissioner] department.

[(d)] (e) Physician means a physician who is licensed and currently registered to practice medicine in New York State or in the state or the country in which he or she practices and is not subject to any disciplinary or non-disciplinary order by the applicable state or country except as otherwise allowed by the department.

[(e)] (f) Dentist means a dentist who is licensed and currently registered to practice dentistry in New York State or in the state or the country in which he or she practices and is not subject to any disciplinary or non-disciplinary order by the applicable state or country except as otherwise allowed by the department.

[(f)] (g) Certificate of qualification means a credential issued by the department to applicants [meeting] determined by the department to meet the requirements set forth in this Part.

[(g)] (h) Grandfathered laboratory director means a laboratory director who qualified for and received a certificate of qualification in one or more categories of testing prior to the amendment of this regulation which became effective January 25, 1988.

[(h)] (i) Category means an area, [procedure, or specialty of laboratory medicine specified in section 19.3(d) of this Part.] field, or discipline of laboratory medicine or laboratory science in
which a certificate of qualification is issued. The department may issue certificates of qualification in a specified subpart of a category, including, but not limited to, a subcategory, technology, method, or specific procedure, based on the applicant’s education, training, and experience and the applicant’s ability to demonstrate that tests performed under their direction generate reliable results. The department shall make available a list of: categories and subcategories in which certificates of qualification are issued; minimum qualifications for each category; and the corresponding categories of testing authorized by a laboratory permit.

[(i) Blood banking-collection means collection of blood or blood components, or processing of blood or blood products.

(j) Referring physician means a physician or other person authorized by law to order laboratory tests and receive reports, as specified in Subpart 58-1 of this Title.

(k) Virology means isolation and other characterization of virus.

(l) Diagnostic immunology means application of immunologic techniques to detect the presence of antigens in biologic fluids and determine host-antibody responses.

(m) Transfusion service means a service which issues blood or blood components for administration into a person, but does not include a limited transfusion service, as defined in section 58-2.1(k) of this Title.

(n) Genetic testing means enzyme, substrate, and DNA-based analyses, or qualitative and/or quantitative measurement of other body analytes, undertaken to determine the genetic status (carrier or disease) of a person.]

(j) Department means the New York State Department of Health.

(k) Board certified means having completed all requirements set forth by an accrediting board
acceptable to the department, including a passing score on any qualifying examination and completion of all the requirements for recertification whenever the certifying board mandates recertification, provided such requirements are determined by the department to provide the applicant with the ability to effectively discharge the responsibilities described in Parts 10 and 58 of this title.

(l) Earned doctoral degree means a doctor of philosophy, doctor of science, or equivalent degree as determined by the department.

(m) Training includes participation in a residency, fellowship, or post-doctoral position, or participation in a training course approved by a board acceptable to the department.

(n) Experience includes post-doctoral employment or voluntary participation in an acceptable laboratory where the applicant performed, supervised or directed testing of human clinical specimens. Teaching experience directly related to a medical technology program, clinical laboratory sciences program, or a clinical laboratory section of a residency program is also considered acceptable experience.

19.2 Clinical laboratory or blood bank; qualifications of laboratory director.

[The] A director and any assistant director of a clinical laboratory or blood bank [must] shall possess training and/or experience acceptable to the department, obtained within the previous six years, [in generally accepted and currently used methods and techniques] in one or more categories, [listed in section 19.3(d) of this Part, and] Additionally, the applicant must meet one of the following requirements:

(a) be a physician who is currently certified by [the American Board of Pathology in]:

(1) the American Board of Pathology in:
[(1)(i)] clinical pathology; or
[(2)(ii)] anatomic pathology; or
[(3) An area of special competence relevant to the certificate of qualification sought; or] (iii) dermatopathology; or

(2) the American Osteopathic Board of Pathology in:

(i) laboratory medicine; or

(ii) anatomic pathology; or

(iii) dermatopathology; or

(b) be a physician in the State of New York who:

(1) is currently certified by the American Board of Pathology in Blood Banking and Transfusion Medicine; or

(2) is currently certified by the American Board of Pathology in Clinical Pathology or the American Board of Internal Medicine in Hematology, and possesses six months of training and/or experience in transfusion services; or

(3) possesses four years of training and/or experience in an acceptable laboratory including two or more years of training and/or experience in transfusion services and in general laboratory management.

[(b)(c)] be a dentist who is currently certified by the American Board of Oral and Maxillofacial Pathology; or

[(c)(d)] be a physician, or hold an earned doctoral degree from an accredited institution with a relevant chemical, physical or biological science major, and:

(1) is currently certified by one of the following boards and meets any supplemental requirements for experience as specified by the department:
(i) [the American Board of Medical Microbiology] the American Board of Bioanalysis as a High Complexity Laboratory Director, provided the applicant has obtained a minimum of four years of post-doctoral experience equivalent to paragraph (2) of this subdivision; or

(ii) the American Board of Clinical Chemistry in clinical chemistry; or

(iii) the American Board of Clinical Chemistry in toxicological chemistry; or

(iv) the American Board of Dermatology; or

[(iv)](v) the American Board of Forensic Toxicology, provided the applicant has an earned doctoral degree; or

[(v) the American Board of Medical Laboratory Immunology; or]

(vi) the American Board of Internal Medicine in hematology or hematology and medical oncology; or

(vii) the American Board of Medical Laboratory Immunology; or

(viii) the American Board of Medical Microbiology; or

(ix) dual certification by the American Board of Pathology in either Anatomic Pathology or Clinical Pathology, and Molecular Genetic Pathology; or

(x) the American Board of Pathology in Medical Microbiology; or

(xi) the National Registry for Certified Chemists, provided the applicant has obtained a minimum of four years of post-doctoral experience equivalent to paragraph (2) of this subdivision; or

(2) subsequent to receiving a doctor of medicine, doctor of [osteopathy] osteopathic medicine or earned doctoral degree has had, and has documented to the department, four years of training and/or experience in an acceptable laboratory, including two or more years of training and/or experience in methods and techniques currently in use in the certificate category or categories
sought and in general laboratory management, or an equivalent combination of training and/or experience as verified by the commissioner department.

[(d) A transfusion facility director shall be a physician licensed to practice medicine in the State of New York.]

19.3 Director of a clinical laboratory or blood bank; certificate of qualification issuance, duties and responsibilities.

(a) Certificate required. [A]An individual serving as a director or assistant director of a clinical laboratory or blood bank must hold a certificate of qualification issued after the commissioner department has determined that the applicant meets the requirements specified in sections 19.2 and 19.3[(e)] of this Part, and has demonstrated, in accordance with subdivision (c) of this section and section 19.4(a) of this Part, that he or she possesses the character, competence, training, and ability to direct the technical and scientific operation of a clinical laboratory or blood bank, and ensure the proper supervision or performance of test procedures, adherence to the department's quality control standards, and accurate reporting of findings of tests.

(b) An applicant for a certificate of qualification must submit a complete, original, signed, and sworn application in such form and manner as may be required by the department, and must supply such additional information as may be required by the department. An individual seeking renewal of a certificate of qualification must submit an application no later than 90 days prior to expiration of the current certificate.

(c) [To function effectively in fulfilling his or her duties and responsibilities,] To qualify for, and maintain, a certificate of qualification, a laboratory director and any assistant director [should
possess a] **shall demonstrate that he or she possesses** knowledge of basic clinical laboratory sciences and operations, and [should] **shall** have the training and/or experience and physical capability to discharge the following responsibilities:

(1) provide advice to referring [physicians] **health care providers** regarding the significance of laboratory findings and ensure that reports of test results include pertinent information required for the interpretation of laboratory data;

* * *

(3) define, implement, and monitor standards of performance [in quality control and quality assurance] for the laboratory and for other ancillary laboratory testing programs in conformance with the department’s clinical laboratory standards of practice;

* * *

(5) assure that the laboratory participates in monitoring and evaluating the quality and appropriateness of services rendered, within the context of [the quality assurance program] a quality management system, regardless of where the testing is performed;

* * *

(7) [set goals and develop and allocate resources within the laboratory] ensure that policies and procedures are established for monitoring staff to assess competency and, whenever necessary, to provide remedial training to improve skills;

(8) [provide effective and efficient administrative direction of the laboratory, including budget planning and controls in conjunction with the individual(s) responsible for financial management of the laboratory] specify in writing the responsibilities and duties of all laboratory personnel;

(9) provide [educational direction] **continuing education** to laboratory staff;

(10) [select all reference laboratories; and] **ensure that a current and complete procedure manual**
is available to all personnel;

(11) [promote a safe laboratory environment for personnel and the public.] set goals, develop and allocate resources within the laboratory;

(12) provide effective administrative direction of the laboratory, in conjunction with the individual(s) responsible for financial management of the laboratory, to ensure adequate resources are available to operate the laboratory in a manner consistent with all state and federal requirements;

(13) select all reference laboratories for services not offered by the laboratory;

(14) promote a safe laboratory environment for personnel and the public; and

(15) ensure that the laboratory, when applicable, is enrolled in a proficiency testing program acceptable to the department for the testing performed and that the laboratory adheres to the proficiency testing program’s administrative and technical requirements.

[(d) Certification. Certificates of qualification are issued in one or more of the following categories, procedures or specialties:

(1) one or more of the subspecialties of microbiology: bacteriology, virology, mycology, mycobacteriology, diagnostic immunology, and parasitology;

(2) hematology;

(3) immunohematology, excluding testing performed solely for transfusion purposes;

(4) one or more of the subspecialties of clinical biochemistry: clinical chemistry, blood pH and gases, endocrinology, and therapeutic substance monitoring/quantitative toxicology;

(5) histopathology, and/or the subspecialties: oral pathology and dermatopathology;

(6) cytopathology;]
(7) cytogenetics;
(8) histocompatibility;
(9) cellular immunology;
(10) oncofetal antigens, and/or the subspecialties: tumor markers, maternal serum, and amniotic fluid;
(11) genetic testing;
(12) transfusion services, including all pre-transfusion testing;
(13) blood banking collection-comprehensive, including all tests required in Subpart 58-2 of this Title;
(14) blood banking collection-limited, including collection of autologous blood for transfusion and excluding testing for transmissible disease markers;
(15) one or more of the subspecialties of clinical toxicology: drug analysis, blood lead, erythrocyte protoporphyrin, and chlorinated hydrocarbons;
(16) forensic toxicology; or
(17) other specific categories, procedures, or specialties designated by the department.]

[(e)(d) Required qualifications.

(1) Applicants for a certificate of qualification in bacteriology, mycobacteriology, mycology, and/or parasitology must qualify under section 19.2[(a)(1), (c)(1)(i), or (c)(2)](a)(1)(i), (a)(2)(i), (d)(1)(i), (d)(1)(viii), (d)(1)(x) or (d)(2) of this Part.

(2) Applicants for a certificate of qualification in virology must qualify under section 19.2[(c)(1)(i) or (c)(2)](d)(1)(viii), (d)(1)(x) or (d)(2) of this Part. Applicants for a certificate of qualification in virology limited to antigen detection and molecular methods must qualify under
(3) Applicants for a certificate of qualification in diagnostic immunology must qualify under section 19.2[(a)(1), (c)(1)(i), (c)(2), or (c)(1)(v)](a)(1)(i), (a)(2)(i), (d)(1)(i), (d)(1)(vii), (d)(1)(viii), (d)(1)(x) or (d)(2) of this Part.

(4) Applicants for certificate of qualification in hematology must qualify under section 19.2[(a)(1), (c)(1)(vi), or (c)(2)](a)(1)(i), (a)(2)(i), (d)(1)(vi) or (d)(2) of this Part. Applicants qualifying under section 19.2[(c)(1)(vi)](d)(1)(vi) of this Part must document that the required training and/or experience includes or is supplemented by six months' training and/or experience in an acceptable laboratory.

(5) Applicants for a certificate of qualification in immunohematology must qualify under section 19.2[(a)(1) or (c)(2)](a)(1)(i), (a)(2)(i), or (d)(2) of this Part.

(6) Applicants for a certificate of qualification in [one or more of the subspecialties of clinical biochemistry] clinical chemistry, blood pH and gases, endocrinology, or therapeutic substance monitoring - quantitative toxicology must qualify under section 19.2[(a)(1), (c)(1)(ii), or (c)(2)](a)(1)(i), (a)(2)(i), (d)(1)(i), (d)(1)(ii), (d)(1)(xi) or (d)(2) of this Part.

(7) Applicants for a certificate of qualification in histopathology and/or cytopathology must qualify under section 19.2[(a)(2)](a)(1)(ii) or (a)(2)(ii) of this Part.

(8) Applicants for a certificate of qualification in oral pathology must qualify under section 19.2[(a)(2) or (b)](a)(1)(ii), (a)(2)(ii), or (c) of this Part.

(9) Applicants for a certificate of qualification in dermatopathology must qualify under section 19.2[(a)(2) or (a)(3)](a)(1)(ii), (a)(1)(iii), (a)(2)(ii), (a)(2)(iii) or (d)(1)(iv) of this Part.

(10) Applicants for a certificate of qualification in cytogenetics, histocompatibility, cellular immunology, [oncofetal antigens, and/or] genetic testing, fetal defect markers, forensic identity,
oncology, parentage/identity testing, trace elements, and/or transplant monitoring must qualify under section 19.2[(c)(2)](d)(2) of this Part.

(11) Applicants for a certificate of qualification in transfusion services must be physicians and must qualify under section 19.2[(a)(3) or (c)(2)](b)(1), (b)(2) or (b)(3) of this Part[; or under section 19.2(a)(1) or (c)(1)(vi) of this Part including or supplemented by at least six months' training and/or experience in transfusion services].

(12) Applicants for a certificate of qualification in blood banking collection-comprehensive must qualify under section 19.2[(c)(2)](d)(2) of this Part. Required experience in blood services must include at least one year's training and/or experience in collection and testing of blood for [homologous] allogeneic transfusion.

(13) Applicants for a certificate of qualification in blood banking collection-limited must qualify under section 19.2[(a)(1), (c)(1)(vi) or (c)(2)](a)(1)(i), (b)(1)(i), or (d)(1)(vi) of this Part.

(14) Applicants for a certificate of qualification in [one or more of the subspecialties of] clinical toxicology must qualify under section 19.2[(a)(1), (c)(1)(iii), (c)(1)(iv), or (c)(2)](a)(1)(i), (b)(1)(i), (d)(1)(ii), (d)(1)(iii), (d)(1)(xi), or (d)(2) of this Part.

(15) Applicants for a certificate of qualification in forensic toxicology must qualify under section 19.2[(c)(1)(iii), (c)(1)(iv), or (c)(2)](d)(1)(iii), (d)(1)(v), or (d)(2) of this Part.

(16) Applicants for a certificate of qualification in andrology must qualify under section 19.2(d)(1)(i) or (d)(2) of this Part; or under section 19.2(a)(1)(i) or (b)(1)(i) of this Part including or supplemented by at least six months' training and/or experience in andrology.

(17) Applicants for a certificate of qualification in blood lead must qualify under section 19.2(a)(1)(i), (b)(1)(i), (d)(1)(i), (d)(1)(iii), (d)(1)(v), (d)(1)(xi), or (d)(2) of this Part.

[(f)] (e) Scope and limitations.
(1) The requirements for qualification set forth in section 19.2 of this Part shall apply to all laboratory directors, regardless of prior grandfathered status, upon expiration of current certificates of qualification[,] if the laboratory director is no longer employed in a laboratory or in the field of laboratory medicine.

(2) Additional categories of testing may not be added to a certificate of qualification issued on a grandfathered basis. Such a certificate [may] will not be renewed if allowed to lapse[, unless extenuating circumstances prevent timely reapplication and specific departmental approval is obtained].

19.4 Denial of an application for a certificate of qualification.

(a) In determining whether to deny an application for a certificate of qualification in whole or in part, the department shall consider: the applicant's education, experience, and licensure as required in sections 19.2 and 19.3 of this Part; the applicant's demonstrated ability to discharge the responsibilities set forth in section 19.3(c) of this Part; the character and competence of the applicant and the laboratory or laboratories directed; and any other factors the department considers relevant, including, but not limited to:

* * *

(3) false representation or omission of any material fact in making an application in any state or city of the United States for any license, permit, certificate, or registration related to a profession or business, or in making an application for a certificate of qualification or laboratory permit to New York State [or New York City];

* * *

(6) on the part of any laboratory, category, or subcategory directed by the applicant, a pattern of
repetitive failures of required proficiency testing performance in one or more proficiency testing categories, excluding failure for administrative reasons such as late result submission;

(7) on the part of any laboratory, category, or subcategory directed by the applicant, a pattern of deficiencies on onsite inspection, especially in areas of quality control, quality assurance, laboratory management, and handling of regulated medical waste and radioactive materials, including refusal or inability to produce records as requested by department employees, which deficiencies are not corrected from inspection to inspection or which recur at each [annual] inspection despite written notice of violations by a state or Federal licensing or auditing agency and which jeopardize the quality of test results and resulting patient care, even if interim corrections have occurred;

(8) on the part of any laboratory, category, or subcategory directed by the applicant, performance of any laboratory procedures not authorized by the laboratory permit issued pursuant to article 5, Title V of the Public Health Law; or operation or direction of a laboratory without a permit; or continuing operation or failure to notify the department after a change in director, ownership, or location has voided the permit;

[(9) unless the laboratory is owned and operated by the State of New York, performance of tests on specimens collected in New York City while the laboratory directed by the applicant lacks a New York City permit to perform such tests;]

[(10)](9) on the part of any laboratory, category, or subcategory directed by the applicant, referral of specimens collected in New York State [outside of New York City] to laboratories which do not possess a New York State permit;

[(11)](10) on the part of any laboratory, category, or subcategory directed by the applicant, knowing acceptance of specimens or requisitions for laboratory examination from, or issuance of
reports to, a person or persons not authorized by law to submit such specimens or requisitions, or receive such reports;

[(12)](11) on the part of any laboratory, category, or subcategory directed by the applicant, issuance of reports on laboratory work, including both patient samples and proficiency testing, actually performed in another laboratory, without designating the fact that the examinations or procedures were performed in another laboratory; and/or testing and reporting results on unsatisfactory specimens as defined by the department, including unlabeled specimens or specimens of insufficient quantity to conduct the analyses requested;

[(13)](12) on the part of any laboratory, category, or subcategory directed by the applicant, failure to establish and ensure that employees follow procedures for disposal or handling of specimens or infectious or radioactive medical waste, in violation of applicable state and Federal laws, rules and regulations, or in a manner which endangers the public, the laboratory's employees, or the environment;

[(14)](13) employment of unqualified or unlicensed technical personnel or an insufficient number of such personnel;

[(15)](14) failure of the [laboratory director] applicant to be responsible for adequately supervising laboratory personnel to ensure the proper performance of all tests conducted in the laboratory; and

[(16)](15) any other factor having a direct bearing on the applicant's ability to provide or supervise the provision of high quality laboratory services, or to ensure compliance with statutory and regulatory requirements.

* * * *
REGULATORY IMPACT STATEMENT

Statutory Authority:
Public Health Law (PHL) section 573 establishes the authority of the Department to promulgate criteria for the issuance of a certificates of qualification. PHL section 573(2) specifically states that the Department shall issue a certificate of qualification to any person who meets such minimum qualifications and who otherwise demonstrates to the Department that he or she possesses the character, competence, training and ability to administer properly the technical and scientific operation of a clinical laboratory or blood bank, including supervision of procedures and reporting of findings of tests.

Legislative Objectives:
The legislature enacted PHL section 573 to protect the health and safety of the public by requiring that only properly educated and experienced individuals be issued certificates of qualification and subsequently assigned responsibility as clinical laboratory directors. Such directors are responsible for the proper operation of clinical laboratories to ensure accurate and reliable results for clinical testing. Part 19 of Title 10 of the Official Compilation of Codes, Rules and Regulations of the State of New York (NYCRR), in its original adoption and all subsequent revisions, has been crafted to ensure that applicants have the necessary education, training and experience to effectively direct a laboratory. The proposed amendment is consistent with this legislative objective as it will include the recognition of additional accrediting boards that have been developed since the last regulatory amendment, in response to changes and advances in clinical laboratory testing.
**Needs and Benefits:**

Part 19 regulates the issuance of certificates of qualification. An individual must hold such certificate to be a clinical laboratory director or assistant director at a clinical laboratory or blood bank permitted by New York under the authority of PHL section 572. The intent of these regulations is to ensure that individuals who are granted certificates of qualification have the necessary education, experience, and training to effectively operate a clinical laboratory. Successful applicants for a certificate of qualification must demonstrate both experience in laboratory management, such as management of resources (e.g. budget allocation, staffing), implementation of a quality management system, development of standard operating procedures; and experience specific to a category of testing defined in Part 19.

Several revisions to the regulatory definitions are proposed. Most notable are the inclusion of a definition for assistant director and revision to the definition of category. Assistant directors are jointly accountable with the laboratory director for the categories of testing on the laboratory permit. In many instances, however, the assistant director may be the only individual qualified to supervise testing in a specific category on the laboratory permit.

Language is proposed in sections 19.2 and 19.3 that clarifies the role and responsibilities of assistant directors of clinical laboratories. With these revisions, assistant directors will be held to the same standards as laboratory directors.

The definition of “category” was revised to strengthen the Department’s authority to limit the approval of a certificate of qualification to a subcategory, technology, method or specific
procedure based on the applicant’s documented experience. Extensive experience in a single method of testing does not necessarily translate to breadth of knowledge across an entire category of testing. Indeed, as innovations in laboratory medicine continue, an individual’s experience in a proven technology may quickly become obsolete without continued education and training. The proposed revisions to the definition of category allow the Department to ascertain an individual’s specific breadth of experience upon each application and re-application for a certificate of qualification.

The definitions of the following terms are being proposed for the first time; board certified, earned doctoral degree, training, and experience.

A review of the accrediting boards currently recognized in Part 19 and those included in the proposed revisions was performed to ensure that the requirements for each board were consistent with the rules set forth in federal regulation. This included a review of both the educational and training requirements for the accrediting board. As noted in the proposed revisions, certain boards mandate the appropriate educational requirement of a doctoral degree, but do not specify that the candidate for the board demonstrate the required four years of post-doctoral experience. Therefore, language clarifying the post-doctoral degree experience required by the Department has been proposed for these boards (American Board of Bioanalysts High Complexity Laboratory Director and the National Registry of Clinical Chemists) to ensure that the requirements for all applicants are consistent.
The duties and responsibilities of laboratory directors and assistant directors set forth in subdivision 19.3(c) were revised to provide clarity and introduce new responsibilities. Of note are the added responsibilities of ensuring the availability of procedures for monitoring staff competency and improvement of skills. These new responsibilities are currently included in the New York State Clinical Laboratory Standards of Practice; however, formal codification in regulation is desired.

Finally, subdivision 19.3(d) has been removed since the certificate of qualification categories are repeated in the current subdivision 19.3(e), and therefore 19.3(d) was considered redundant. The Department currently maintains a list of certificate of qualification categories on its publicly accessible website, and revisions were made in proposed subdivision 19.1(i) to outline the necessary contents of this list.

Costs:

Costs to Regulated Parties:

The proposed amendment will not impose costs on regulated parties. The current regulation already requires clinical laboratories and blood banks to have directors who hold certificates of qualification.

Costs to the Agency, State and Local Governments:

The proposed amendment will not impose additional costs to the New York State Department of Health, the program responsible for oversight of clinical laboratories, or to local governments. The program responsible for the oversight of clinical laboratories is a well-established program
operated at the State level and the new language does not impact the costs of the oversight program.

**Local Government Mandates:**
The proposed regulations impose no new mandates on any county, city, town or village government; or school, fire or other special district.

**Paperwork:**
The proposed revisions to Part 19 do not require any additional forms or paperwork from applicants. All candidates are required under the current rule to provide a complete application, a curriculum vitae, and proof of licensure for physicians or granting of an earned doctoral degree. Additionally, candidates must submit proof of any accreditation by a recognized board and/or letters from third parties attesting to the candidate’s training and experience. The proposed revisions expand the list of recognized accrediting boards, which may in fact reduce the paperwork needed for candidates holding those accreditations.

**Duplication:**
The federal government also recognizes clinical laboratory directors. The Department has applied and been approved for an exemption from the federal government continuously since 1995 that grants the Department the authority to act as the primary accrediting body for clinical laboratories and clinical laboratory directors operating in New York.
 Alternatives:

The alternative to this proposal would be to maintain the existing regulatory requirements. However, the proposed amendments are necessary to update the regulations to include new definitions, update the list of acceptable accrediting boards, and clarify and expand the responsibilities of laboratory directors and assistant directors.

 Federal Standards:

The Federal Code of Regulations (CFR) sets forth rules for the education and experience of clinical laboratory directors (CFR 493.1443). The proposed revisions to Part 19 will incorporate several of the accrediting boards that are already recognized under the federal rule.

 Compliance Schedule:

Regulated parties are expected to comply with the proposed regulation by its effective date.

 Contact Person:

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STATEMENT IN LIEU OF REGULATORY FLEXIBILITY ANALYSIS
FOR SMALL BUSINESS AND LOCAL GOVERNMENTS

No regulatory flexibility analysis is required. The proposed amendment does not impose an adverse economic impact on small businesses or local governments, and it does not impose reporting, record keeping or other compliance requirements on small businesses or local governments. At present the regulations require clinical laboratories and blood banks to be directed by individuals who hold a certificate of qualification. This proposed amendment would update and expand the list of acceptable accrediting boards for obtaining a certificate of qualification and is therefore anticipated to have a positive impact by increasing the number of individuals who may qualify for a certificate of qualification.
STATEMENT IN LIEU OF RURAL AREA FLEXIBILITY ANALYSIS

No rural area flexibility analysis is required pursuant to § 202-bb(4)(a) of the State Administrative Procedure Act. The proposed amendments will not impose an adverse impact on facilities in rural areas, and will not impose any significant new reporting, record keeping or other compliance requirements on facilities in rural areas.
STATEMENT IN LIEU OF JOB IMPACT STATEMENT

No job impact statement is required pursuant to § 201-a(2)(a) of the State Administrative Procedure Act. No adverse impact on jobs and employment opportunities is expected as a result of these proposed regulations.