Laboratory General Checklist

CAP Accreditation Program
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# Laboratory General Checklist

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ON-LINE CHECKLIST AVAILABILITY

Participants of the CAP accreditation programs may download the checklists from the CAP website (www.cap.org) by logging into e-LAB Solutions. They are available in different checklist types and formatting options, including:

- Master — contains ALL of the requirements and instructions available in PDF, Word/XML or Excel formats
- Custom — customized based on the laboratory’s activity (test) menu; available in PDF, Word/XML or Excel formats
- Changes Only — contains only those requirements with significant changes since the previous checklist edition in a track changes format to show the differences; in PDF version only. Requirements that have been moved or merged appear in a table at the end of the file.

SUMMARY OF CHECKLIST EDITION CHANGES

Laboratory General Checklist
08/17/2016 Edition

The information below includes a listing of checklist requirements with significant changes in the current edition and previous edition of this checklist. The list is separated into three categories:

1. New
2. Revised:
   - Modifications that may require a change in policy, procedure, or process for continued compliance; or
   - A change to the Phase
3. Deleted/Moved/Merged:
   - Deleted
   - Moved — Relocation of a requirement into a different checklist (requirements that have been resequenced within the same checklist are not listed)
   - Merged — The combining of similar requirements

*NOTE: The listing of requirements below is from the Master version of the checklist. The customized checklist version created for on-site inspections and self-evaluations may not list all of these requirements.*

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UNDERSTANDING THE CAP ACCREDITATION CHECKLIST COMPONENTS

All checklist requirements contain a requirement number, subject header, phase, and a declarative statement. Some requirements also contain a NOTE and/or Evidence of Compliance.

The NOTE portion of a checklist requirement provides additional detail to assist in interpreting the requirement.

Evidence of Compliance (EOC) is intended to:

- Suggest specific examples of acceptable records; some elements are required
- Assist in inspection preparation and for managing ongoing compliance
- Drive consistent understanding of requirements

If a policy or procedure is referenced within a requirement, it is only repeated in the Evidence of Compliance if such statement adds clarity. All policies or procedures covered in the CAP checklists must be a written document. A separate policy or procedure may not be needed for items in EOC if it is already addressed by an overarching policy.

The Master version of the checklist also contains references and the inspector R.O.A.D. instructions (Read, Observe, Ask, Discover), which can provide valuable insight for the basis of requirements and on how compliance will be assessed.

INTRODUCTION

The Laboratory General (GEN) Checklist applies to all sections or departments of the laboratory. It is customized based on the services reported by the laboratory to the CAP on its application.

One copy of the GEN Checklist is provided to the inspection team. One or more inspectors may be assigned to inspect with the GEN Checklist; however, all inspectors must be familiar with the GEN Checklist requirements and ensure that all areas are in compliance. For suggestions on how inspectors can assist the Laboratory General inspector, please refer to the Laboratory General (GEN) section in the Laboratory Accreditation Manual.

Note for non-US laboratories: Checklist requirements apply to all laboratories unless a specific disclaimer of exclusion is stated in the checklist.

DEFINITION OF TERMS

Addendum - Information appended to a final report without changing any of the language in the original report (original report is intact and unchanged).

Alternative assessment - A system for determining the reliability of laboratory examinations for which no commercial proficiency testing products are available, are not appropriate for the method or patient population served by the laboratory, or participation is not required by the accrediting organization.

Amended/amendment - Any change in the diagnosis, narrative text, or other content of a report that has been issued (minor or major). The change in an anatomic pathology report is usually in the diagnosis or narrative, but occasionally may involve a change in a number or some other quality.

Analytical validation - The process used to confirm with objective evidence that a laboratory-developed or modified FDA-cleared/approved test method or instrument system delivers reliable results for the intended application.
Analytical verification - The process by which a laboratory determines that an unmodified FDA-cleared/approved test performs according to the specifications set forth by the manufacturer when used as directed.

Annual - Every 12 calendar months

Biennial - Every 24 calendar months

Authority - The power to give orders or make decisions: the power or right to direct someone or control a process

Calibrator, historical - The set of archived results of a single-point calibrator that demonstrates stability of the assay over time

Check - Examination to determine the accuracy, quality or presence of any attribute of a test system

Clinical validation - The determination of the ability of a test to diagnose or predict risk of a particular health condition or predisposition, measured by sensitivity, specificity, and predictive values

Commutable - The property of a reference material that yields the same numeric result as would a patient's specimen containing the same quantity of analyte in that analytic method under discussion (i.e. matrix effects are absent).

Confirmation - Substantiation of the correctness of a value or process

Corrected/correction - Errors in test results that may include incorrect patient identification, test results, reference interval, interpretive information, or other significant information, but not minor typographical errors of no consequence.

Corrective Action - Action taken to eliminate the cause of a detected nonconformity or other undesirable situation

Correlation - Establishment of agreement between two or more measured values

Credentialing - The process of obtaining, verifying, and assessing the qualifications of a practitioner to provide care in a health care organization

Device - Any reagent, reagent product, kit, instrument, apparatus, equipment or related product, whether used alone or in combination, intended by the manufacturer to be distributed for use in vitro for the examination of human specimens

Digital image analysis - The computer-assisted detection or quantification of specific features in an image following enhancement and processing of that image, including immunohistochemistry, DNA analysis, morphometric analysis, and in situ hybridization

Equipment - Single apparatus or set of devices or apparatuses needed to perform a specific task

Examination - In the context of checklist requirements, examination refers to the process of inspection of tissues and samples prior to analysis. An examination is not an analytical test.

FDA - In the context of checklist requirements, FDA should be taken to mean the national, state, or provincial authority having jurisdiction over in vitro diagnostic test systems.

Function Check - Confirmation that an instrument or item of equipment operates according to manufacturer's specifications before routine use, at prescribed intervals, or after minor adjustment (e.g. base line calibration, balancing/zero adjustment, thermometer calibration, reagent delivery).
High complexity - Rating given by the FDA to commercially marketed in vitro diagnostic tests based on their risks to public health. Tests in this category are seen to have the highest risks to public health.

Instrument - An analytical unit that uses samples to perform chemical or physical assays (e.g. chemistry analyzer, hematology analyzer)

Instrument platform - Any of a series of similar or identical analytical methods intended by their manufacturer to give identical patient results across all models

Laboratory Director - The individual who is responsible for the overall operation and administration of the laboratory, including provision of timely, reliable and clinically relevant test results and compliance with applicable regulations and accreditation requirements. This individual is listed on the laboratory’s CAP and CLIA certificate (as applicable).

Maintenance - Those activities that prolong the life of an instrument or minimize breakdowns or mechanical malfunctions. Examples include cleaning, changing parts, fluids, tubing, lubrication, electronic checks, etc.

Moderate complexity - Rating given by the FDA to commercially marketed in vitro diagnostic tests based on their risks to public health

Modification of manufacturer’s instructions - Any change to the manufacturer’s supplied ingredients or modifications to the assay as set forth in the manufacturer’s labeling and instructions, including specimen type, instrumentation or procedure that could affect its performance specifications for sensitivity, specificity, accuracy, or precision or any change to the stated purpose of the test, its approved test population, or any claims related to interpretation of the results

Nonwaived - Tests categorized as either moderately complex (including provider-performed microscopy) or highly complex by the US Food and Drug Administration (FDA), according to a scoring system used by the FDA

Performance verification - The set of processes that demonstrate an instrument or an item of equipment operates according to expectations upon installation and after repair or reconditioning (e.g. replacement of critical components)

Policy - 1) Set of basic principles or guidelines that direct or restrict the facility's plans, actions, and decisions; 2) Statement that tells what should or should not be done

Preventive action - Action taken to eliminate the cause of a potential nonconformity or any other undesirable potential situation

Primary source verification report - A document, usually prepared by a third party agent or company that confirms that a job applicant's degree, certificate, or diploma is authentic, licenses were granted, and reported work history (company names, locations, dates and positions held) is accurate. The confirmation is obtained through direct contact with an institution, former employer, or their authorized agents.

Primary specimen - The body fluid, tissue, or sample submitted for examination, study or analysis. It may be within a collection tube, cup, syringe, swab, slide, data file, or other form as received by the laboratory.

Procedure - 1) Specified way to carry out an activity of a process (also referred to by ISO as "work instructions"; 2) Set of steps performed that tells "how to do it" to achieve a specified outcome, including decisions to be made

Process - 1) Set of interrelated or interacting activities that transforms inputs into outputs; 2) Series of events, stages, or phases that takes place over time that tells "what happens" or "how it works"

Proficiency testing - Evaluation of participant (laboratory or individual) performance against pre-established criteria by means of interlaboratory comparisons. In some countries, the PT programs for clinical laboratories are called "external quality assessment" programs.
**Reagent** - Any substance in a test system other than a solvent or support material that is required for the target analyte to be detected and its value measured in a sample.

**Report errors** - A report element (see GEN.41096) that is either incorrect or incomplete

**Responsibility** - A duty or task that an individual is required or expected to do

**Secondary specimen** - Any derivative of the primary specimen used in subsequent phases of testing. It may be an aliquot, dilution tube, slide, block, culture plate, reaction unit, data extract file, image, or other form during the processing or testing of a specimen. (The aliquots or images created by automated devices and tracked by internal electronic means are not secondary specimens.)

**Section Director** - The individual who is responsible for the medical, technical and/or scientific oversight of a specialty or section of the laboratory.

**Semiannual** - Every 6 calendar months

**Subject to U.S. Regulations** - Laboratories located within the United States and laboratories located outside of the US that have obtained or applied for a CLIA certificate to perform laboratory testing on specimens collected in the US for the assessment of the health of human beings.

**Telepathology** - The practice in which the pathologist views digitized or analog video or still image(s), and renders an interpretation that is included in a formal diagnostic report or is recorded in the patient record.

**Testing personnel** - Individuals responsible for performing laboratory assays and reporting laboratory results

**Test** - A qualitative, semiquantitative, quantitative, or semiquantitative procedure for detecting the presence of, or measuring an analyte

**Test system** - The process that includes pre-analytic, analytic, and post-analytic steps used to produce a test result or set of results. A test system may be manual, automated, multi-channel or single-use and can include reagents, components, equipment or instruments required to produce results. A test system may encompass multiple identical analyzers or devices. Different test systems may be used for the same analyte.

**Waived** - A category of tests defined as "simple laboratory examinations and procedures which have an insignificant risk of an erroneous result." Laboratories performing waived tests are subject to minimal regulatory requirements.
QUALITY MANAGEMENT

The laboratory must have a written quality management program to systematically ensure the quality of laboratory services. In laboratories that are part of a larger institution (e.g. a hospital), the laboratory quality management program must be integrated with the institutional program.

Although effective organization of the laboratory and appropriate delegation of duties are part of quality management, these areas are addressed in the Team Leader Checklist. Quality management requirements pertaining to all laboratory sections are addressed in the All Common Checklist.

Inspector Instructions:

- Policy for communication of employee concerns
- Sampling of quality indicators with follow-up actions when targets are not achieved
- Annual appraisal of effectiveness of the QM Program
- Document control policy
- Record/specimen retention policy
- Error, complaint and incident logs with corrective/preventative actions
- Device-related adverse patient event procedure and records of reporting (if applicable)
- Results of the laboratory's self-evaluation and correction of deficiencies
- Sampling of records of manufacturer's recalls and records of follow-up

- CAP sign regarding the reporting of quality concerns

- How is the laboratory's QM performance communicated to other hospital departments?
- How was referring physicians'/clients'/or patients' satisfaction measured? What were the results and what actions were taken as a result of the findings?
- Is there a specific example when problems were identified that could have interfered with patient care or safety?

- If any problems are found during review of quality measurements, or when asking questions, further evaluate the laboratory's investigation and resolution, including root cause analysis and associated risk-reduction activities when applicable
- If trends in negative feedback are identified in the satisfaction survey, further evaluate investigation and corrective actions

The laboratory has a written quality management (QM) program.

NOTE: There must be a document that describes the overall QM program. The document need not be detailed, but should spell out the objectives and essential elements of the QM program. The QM plan may be based upon some reference resource such as CLSI QMS01-04; the ISO 9000 series or ISO 15189; AABB's quality program; CAP's quality management publications;
or it may be of the laboratory's own design. If the laboratory is part of a larger organization, the laboratory QM program is coordinated with the organization's QM plan.

REFERENCES

GEN.16902 QM Implementation Phase II

For laboratories that have been CAP accredited for more than 12 months, the QM plan is implemented as designed and is reviewed annually for effectiveness.

NOTE: Appraisal of program effectiveness may be evidenced by an annual written report, revisions to laboratory policies and procedures, or revisions to the QM plan, as appropriate.

Evidence of Compliance:
✓ Evidence that the plan has been implemented as designed requires all of the following:
  ● quality measurements/assessments specified in the plan are being substantially carried out;
  ● there is evidence of active review of quality measurements;
  ● if target performance levels are specified in the plan and the targets are not being met, there are records of follow-up action;
  ● any interventions/changes to operations that are specified in the plan have been carried out as scheduled, or the reason for delay recorded; AND
  ● any communication of information that is required by the plan have taken place

REFERENCES

GEN.20100 QM Extent of Coverage Phase II

The QM program covers all areas of the laboratory and all beneficiaries of service.

NOTE: The QM program must be implemented in all areas of the laboratory (e.g. chemistry, anatomic pathology, satellite, point-of-care, consultative services). The program must include all aspects of the laboratory's scope of care, such as inpatient, outpatient, and referral laboratory services.

GEN.20208 QM Patient Care Services Phase II

The QM program includes a process to identify and evaluate errors, incidents and other problems that may interfere with patient care services.

NOTE: There must be an organized process for recording of problems involving the laboratory that are identified internally, as well as those identified through outside sources such as complaints from patients, physicians or nurses. The process must be implemented in all sections of the laboratory, and on all shifts. Any problem that could potentially interfere with patient care or safety must be addressed. Clinical, rather than business/management issues, should be emphasized. The laboratory must record investigation and resolution of these problems.
Laboratories must perform root cause analysis of any unexpected event involving death or serious physical or psychological injury, or risk thereof (including “near misses” and sentinel events). Laboratories must be able to demonstrate appropriate risk-reduction activities based on such root cause analyses.

REFERENCES

**REVISED** 08/17/2016
GEN.20316 QM Indicators of Quality

The QM program includes monitoring key indicators of quality in the pre-analytic, analytic, and post-analytic phases.

NOTE: Key indicators should monitor activities critical to patient outcome or that may affect many patients. The laboratory must evaluate its indicators by comparing its performance against available benchmarks. The laboratory should also evaluate the effectiveness of each corrective action. The number of monitored indicators should be consistent with the laboratory’s scope of care. Special function laboratories may monitor fewer indicators; full-service laboratories should monitor multiple aspects of the testing process appropriate to their scope of service.

For laboratories that have implemented one or more individualized quality control plans (IQCPs), the quality management program must include a review of the ongoing monitoring of the effectiveness of each IQCP.

While there is no requirement to monitor any specific laboratory indicator, the following key quality indicators have been commonly used to measure laboratory performance in a consistent manner and are important to clinicians and patients as indices of care.

1. **Patient/Specimen Identification:** Percent of patient wristbands with errors, percent of ordered tests with patient identification errors, or percent of results with identification errors
2. **Test Order Accuracy:** Percent of test orders correctly entered into a laboratory computer
3. **Specimen Acceptability:** Percent of specimens accepted for testing
4. **Stat Test Turnaround Time:** Collection-to-reporting turnaround time or receipt-in-laboratory-to-reporting turnaround time of tests ordered with a “stat” priority (e.g. emergency department or intensive care unit specimens), mean or median turnaround time, or the percent of specimens with turnaround time that falls within an established limit
5. **Critical Value Reporting:** Percent of critical results with written record that results have been reported to caregivers; percent of critical results for which the primary clinician cannot be contacted in a reasonable period of time
6. **Customer Satisfaction:** Standardized satisfaction survey tool with a reference database of physician, nurse, or patient respondents
7. **Corrected Reports – General Laboratory:** Percent of reports that are corrected
8. **Corrected Reports – Anatomic Pathology:** Percent of reports that are corrected
9. **Surgical Pathology/Cytology Specimen Labeling**: Percent of requisitions or specimen containers with one or more errors of pre-defined type

10. **Blood Component Wastage**: Percentage of red blood cell units or other blood components that are not transfused to patients and not returned to the blood component supplier for credit or reissue

11. **Blood Culture Contamination**: Percent of blood cultures that grow bacteria that are highly likely to represent contaminants

Performance of indicators should be compared with benchmarks, preferably from multi-institutional studies conducted within ten years of the laboratory's use of the monitor, where such surveys are available.

Both the College of American Pathologist's Q-TRACKS Program itself and publications of Q-TRACKS studies in the Archives of Pathology provide information regarding definitions of quality indicators and demonstrate statistically valid peer-group performance standards.

For benchmark information on commonly used quality indicators, please refer to the Quality Management Quality Indicator Monitoring Guidance Document posted on the CAP Website at the following link: [http://www.cap.org/apps/docs/laboratory_accreditation/qim.pdf](http://www.cap.org/apps/docs/laboratory_accreditation/qim.pdf)

**Evidence of Compliance:**

- Listing of quality indicators that include the following:
  - indicators for pre-analytic, analytic, and post-analytic phases **AND**
  - indicators to address the scope of testing and laboratory services **AND**
  - frequency for monitoring each indicator **AND**
  - defined benchmarks for the performance of each indicator **AND**
- Quality management data and reports for quality indicator monitoring and evaluation, including, comparison against benchmark data, and corrective action when targets are not met

**REFERENCES**

1. Clinical Laboratory Improvement Amendments 42 CFR § 493.1701
Employee and Patient Quality Communication

The laboratory has a procedure for employees and patients to communicate concerns about quality and safety to management.

NOTE: The investigation and analysis of employee and patient complaints and suggestions, with corrective or preventive action as appropriate, should be a part of the laboratory quality management program and be specifically addressed in laboratory quality management records.

Evidence of Compliance:
✓ Records of employee and patient complaints (if any) with appropriate follow up

CAP Sign

The laboratory posts the official CAP sign regarding reporting of quality concerns.

NOTE: The laboratory must prominently post the official CAP sign regarding the reporting of quality concerns to CAP.

While personnel should report concerns to laboratory management, the laboratory must ensure that all personnel know that they may communicate with CAP directly if they have a concern not addressed by laboratory management, and that CAP holds such communications in strict confidence. In addition, the laboratory must have a policy prohibiting harassment or punitive action against an employee in response to a complaint or concern made to CAP or other regulatory organization regarding laboratory quality or safety.

The dedicated, confidential CAP telephone line for quality or safety concerns is 866-236-7212 (US, toll-free) and 847-832-7533 (international).

Official CAP signs may be obtained by calling 800-323-4040 option 1#.

Customer Satisfaction

The laboratory has measured the satisfaction of healthcare providers or patients with laboratory services within the past two years.

NOTE: Satisfaction metrics are important for understanding the needs of clients (physicians, patients, referring laboratories, nurses, etc.) to improve laboratory services. Experience has shown that surveys are more informative if they are conducted anonymously and allow for open ended comments. The sample size should be adequate. A numeric satisfaction scale allows for calculation of statistics.

Evidence of Compliance:
✓ Records of the design and results of satisfaction surveys

REFERENCES
GEN.20340 Notifications From Vendors

The laboratory manages notifications from vendors of defects or issues with supplies or software that may affect patient care.

NOTE: Notifications may take the form of product recalls, market withdrawals, or software patches and upgrades. The laboratory should take action on those that have the potential to affect testing results or laboratory services.

Evidence of Compliance:
✓ Records of manufacturer’s recalls received AND
✓ Records of follow-up

GEN.20351 Adverse Patient Event Reporting

The laboratory has a procedure for reporting device-related adverse patient events, as required by the FDA.

NOTE: This checklist item does NOT apply to laboratories accredited under the CAP Forensic Drug Testing program. Non-US laboratories are encouraged to comply with this checklist item, either through reporting to the FDA in the US or to their national equivalent.

When information reasonably suggests that any laboratory instrument, reagent or other device (including all instruments in the central laboratory, satellite laboratories, point-of-care testing programs, and accessory devices used for phlebotomy or specimen collection) has or may have caused or contributed to a patient death or serious patient injury, the FDA requires hospitals and outpatient diagnostic facilities, including independent laboratories, to report the event. If the event is death, the report must be made both to the FDA and the device manufacturer. If the event is serious patient injury, the report may be to the manufacturer only, unless the manufacturer is unknown, in which case the report must be submitted to the FDA. Reports must be submitted on the FDA Form 3500A (or an electronic equivalent) as soon as practical but no later than 10 days from the time medical personnel become aware of the event.

The FDA defines “serious patient injury” as one that is life threatening; or results in permanent impairment of a body function or permanent damage to a body structure; or necessitates medical or surgical intervention to preclude permanent impairment of a body function or permanent damage to a body structure. Device malfunctions or problems that are reportable may relate to any aspect of a test, including hardware, labeling*, reagents or calibration; or to user error (since the latter may be related to faulty instrument instructions or design). An adverse patient event that may have resulted from inherent limitations in an analytic system (e.g. limitations of sensitivity, specificity, accuracy, and precision) is not reportable.

The laboratory should have written procedures for 1) the identification and evaluation of adverse patient events, 2) the timely submission of MDR (medical device reporting) reports, and 3) compliance with record keeping requirements. A written record of participation in the overall institutional MDR process is required of laboratories that are part of a larger organization (e.g. hospital laboratories).

The laboratory should educate its personnel in the FDA MDR requirements.

The laboratory (or parent institution, as appropriate) must submit an annual report of device-related deaths and serious injuries to FDA, if any such event was reported during the previous year. Annual reports must be submitted on Form 3419 (for hospital-based laboratories only, or an electronic equivalent) or Form 3500 (for non-hospital-based laboratories) by January 1 of each year. The laboratory or institution must keep records of MDR reports for 2 years.

Additional information is available on the FDA website, at http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/default.htm

*In this context, “labeling” refers to all user instructions provided by the manufacturer.
Evidence of Compliance:
✓ Records of MDR reports for reportable events, if applicable

REFERENCES

**NEW** 07/28/2015
GEN.20361 CLIA Certificate Type

For laboratories subject to US regulations performing patient testing subject to CLIA, the laboratory has registered with the Centers for Medicare and Medicaid Services (CMS) and obtained a CLIA certificate that corresponds to the complexity of testing performed, as applicable.

NOTE: This requirement does not apply to laboratories that are part of the Department of Defense. Laboratories located in CLIA exempt states, such as Washington and New York, must be able to show that they have obtained a CLIA number, when appropriate.

The CLIA regulations define a laboratory as a facility that performs testing on materials derived from the human body for the purpose of providing information for the diagnosis, prevention, or treatment of any disease or impairment of, or the assessment of the health of, human beings.

Examples of laboratory activities that do not require registration with the CMS for a CLIA number include:
- Specimen collection
- Specimen preparation, including histology, tissue embedding, sectioning, and staining
- Forensic testing
- Research testing on human specimens where patient-specific results are not reported to the clinician
- Drug testing meeting SAMHSA guidelines and regulations

Laboratories must obtain the CLIA certificate type that corresponds to their highest level of complexity. The CLIA certificate types include:
- Certificate of Waiver - waived tests only*
- Certificate of Provider Performed Microscopy (PPM) Procedures - testing performed by a physician, midlevel practitioner or dentist for specific microscopy procedures (moderate complexity) during the course of a patient’s visit
- Certificate of Registration - nonwaived testing (moderate or high complexity) prior to initial laboratory inspection
- Certificate of Compliance - nonwaived testing with inspection by the State Department of Health (CLIA inspection)
- Certificate of Accreditation - nonwaived testing with inspection by a CMS-approved accrediting organization, such as the CAP’s accreditation programs.


*Any modification from the manufacturer’s instructions changes the test classification to nonwaived and requires a different type of CLIA certificate.

GEN.20374 Federal/State/Local Regulations

The laboratory has a policy for ensuring compliance with applicable federal, state and local laws and regulations.
NOTE: Applicable federal, state and local requirements may include but are not limited to the following areas: handling radioactive materials, shipping infectious or diagnostic materials, personnel qualifications, retention of specimens and records, hazardous waste disposal, fire codes, medical examiner or coroner jurisdiction, legal testing, acceptance of specimens only from authorized personnel, handling controlled substances, patient consent for testing, confidentiality of test results, and donation of blood. The checklists contain specific requirements on these areas.

The laboratory may obtain information on applicable federal, state and local laws and regulations from multiple sources, including hospital management, state medical societies and state departments of health.

REFERENCES
1) Department of Health and Human Services, Centers for Medicare & Medicaid Services. Clinical laboratory improvement amendments of 1988; final rule. Fed Register. 2004(Oct 1); [42CFR493.1101(c)]

**REVISED** 07/28/2015
GEN.20375 Document Control Phase II

The laboratory has a document control system to manage policies, procedures, and forms that are subject to CAP accreditation.

NOTE: This includes documents relating directly to laboratory testing, as well as others, such as quality management, safety, specimen collection, personnel, and laboratory information systems. The document control system must ensure that only current policies, procedures (including derivative documents such as card files and summary charts), and forms are in use and that records for approval, review, and discontinuance are available.

It is recommended that the laboratory maintain a control log listing all current policies, procedures, and forms with the locations of copies. The control log may contain other information as appropriate, such as dates when policies and procedures were placed in service, schedule of review, identity of reviewer(s), and dates when policies and procedures were discontinued and/or superseded.

Additional requirements regarding procedure manuals are found in the All Common Checklist, and in this checklist in the Collection Manual, Computer Services and Safety sections.

REFERENCES

**REVISED** 08/17/2016
GEN.20377 Record/Specimen Retention Phase II

Laboratory records and materials are retained for an appropriate time.

NOTE: Policies for retention of records and materials must comply with federal, state and local laws and regulations and with the retention periods listed below, whichever is most stringent. For testing on minors (under the age of 21), stricter state regulations may apply.

More specific requirements for certain laboratory records are found in the Anatomic Pathology, Cytopathology, Cytogenetics, Molecular Pathology, Reproductive Laboratory Medicine, and Transfusion Medicine Checklists.

<table>
<thead>
<tr>
<th>Type of Record/Material</th>
<th>Retention Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specimen requisitions (including the patient chart or medical record if used as the requisition)</td>
<td>2 years</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Accession records</td>
<td>2 years</td>
</tr>
<tr>
<td>Quality management records</td>
<td>2 years</td>
</tr>
<tr>
<td>Validation/verification of method performance specifications</td>
<td>2 years after discontinuation of the test</td>
</tr>
<tr>
<td>Proficiency testing records</td>
<td>2 years (5 years for transfusion medicine)</td>
</tr>
<tr>
<td>Policies and procedures</td>
<td>At least 2 years following discontinuance (5 years for transfusion medicine)</td>
</tr>
<tr>
<td>Quality control records</td>
<td>2 years (5 years for transfusion medicine)</td>
</tr>
<tr>
<td>Individualized Quality Control Plan (IQCP), including risk assessment and supporting data, and approval of quality control plan</td>
<td>2 years following discontinuation of the IQCP</td>
</tr>
<tr>
<td>Ongoing quality assessment data</td>
<td>2 years</td>
</tr>
<tr>
<td>Instrument maintenance* and function check records</td>
<td>2 years</td>
</tr>
<tr>
<td><strong>Personnel Records</strong></td>
<td></td>
</tr>
<tr>
<td>Competency assessment records</td>
<td>2 years (5 years for transfusion medicine)</td>
</tr>
<tr>
<td>Training records</td>
<td>2 years (5 years for transfusion medicine)</td>
</tr>
<tr>
<td><strong>Patient Specimens (stored under appropriate conditions)</strong></td>
<td></td>
</tr>
<tr>
<td>Serum, heparinized plasma, EDTA plasma, CSF, and body fluids (except urine)</td>
<td>48 hours</td>
</tr>
<tr>
<td>Whole blood specimens, including blood gas specimens</td>
<td>Not defined</td>
</tr>
<tr>
<td>Urine</td>
<td>24 hours; exceptions may be made at the discretion of the laboratory director.</td>
</tr>
<tr>
<td><strong>Clinical Pathology Slides</strong></td>
<td></td>
</tr>
<tr>
<td>Blood Films</td>
<td>7 days</td>
</tr>
<tr>
<td>Permanently stained body fluid slides</td>
<td></td>
</tr>
<tr>
<td>Permanently stained microbiology slides prepared from clinical specimens (including blood culture bottles)</td>
<td></td>
</tr>
</tbody>
</table>
### Testing Records

<table>
<thead>
<tr>
<th>Description</th>
<th>Retention Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Instrument printouts and worksheets **</td>
<td>2 years</td>
</tr>
<tr>
<td>Patient test results and reports, including original and corrected reports, and referral laboratory reports</td>
<td></td>
</tr>
<tr>
<td>Direct-to-consumer testing results, including reference ranges</td>
<td>10 years</td>
</tr>
</tbody>
</table>

**Laboratory Computer Services**

<table>
<thead>
<tr>
<th>Description</th>
<th>Retention Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Computer system validation records</td>
<td>2 years beyond the life of the system</td>
</tr>
<tr>
<td>Records of changes to software, the test library, and major functions of laboratory information systems</td>
<td></td>
</tr>
<tr>
<td>Ongoing computer system checks (e.g. calculation verification)</td>
<td>2 years</td>
</tr>
</tbody>
</table>

* Laboratories may wish to retain instrument maintenance records for longer than the two-year requirement (e.g. for the life of the instrument), to facilitate trouble-shooting.

** For data directly transmitted from instruments to the laboratory computer system via an interface (on-line system), it is not necessary to retain paper worksheets, printouts, etc., so long as the computer retains the data for at least two years. Manual computer entry of patient result data from worksheets, print-outs, etc. requires retention of all worksheets, printouts, etc. for at least two years (digitized or photographic images are acceptable). For results that are manually entered into the computer from 1) observation of an electronic display, with no paper print-out available, or 2) manually performed test methods without worksheets, the two-year retention requirement applies to the data within the computer.

### Evidence of Compliance:
- ✓ Written policy for retention of records, specimens and slides

### REFERENCES

### GEN.20425 Record Retention Phase II

The laboratory has a policy to ensure that all records, slides, blocks, and tissues are retained and available for appropriate times should the laboratory cease operation.

### REFERENCES

**NEW** 08/17/2016

### GEN.20450 Correction of Laboratory Records Phase II

The laboratory follows a written policy for the management and correction of laboratory records, including quality control data, temperature logs, and intermediate test results or worksheets.
NOTE: Laboratory records and changes to such records must be legible and indelible. Original (erroneous) entries must be visible (i.e. erasures, white and correction fluid are unacceptable) or accessible (e.g. audit trail for electronic records). Corrected data, including the identity of the person changing the record and when the record was changed, must be accessible to audit. This requirement does not apply to changes to patient reports (refer to GEN.41310).

Evidence of Compliance:
✓ Records of corrections to laboratory records following the policy

GEN.23584 Interim Self-Inspection Phase II

The laboratory conducts an interim self-inspection and records efforts to correct deficiencies identified during that process.

NOTE: The interim self-inspection is an important aspect of continuing education and laboratory improvement. The use of a variety of mechanisms for self-inspection (residents, technologists or others trained to perform inspections) is strongly endorsed. Self inspection by personnel familiar with, but not directly involved in, the routine operation of the laboratory section to be inspected is a best practice. Records of performance of the interim self-inspection with correction of deficiencies is a requirement for maintaining accreditation. The laboratory must have a record to demonstrate that personnel responsible for each laboratory section have reviewed the findings of the interim self-inspection.

Evidence of Compliance:
✓ Written evidence of self-inspection findings with records of corrective action

REFERENCES

**REVISED** 07/28/2015

GEN.26791 Terms of Accreditation Phase II

The laboratory has a policy that addresses compliance with the CAP terms of accreditation.

NOTE: The CAP terms of accreditation are listed in the laboratory’s official notification of accreditation. The policy must include notification of CAP regarding the following:

1. Investigation of the laboratory by a government entity or other oversight agency, or adverse media attention related to laboratory performance; notification must occur no later than two working days after the laboratory learns of an investigation or adverse media attention. For laboratories subject to US regulations, this notification must include any complaint investigations conducted or warning letters issued by any oversight agency (e.g. CMS, State Department of Health, The Joint Commission, FDA, OSHA).
2. A facility must notify the CAP as soon as it finds itself to be the subject of a validation inspection
3. Discovery of actions by laboratory personnel that violate national, state or local regulations
4. Change in laboratory test menu prior to beginning that testing or the laboratory permanently or temporarily discontinues some or all testing
5. Change in laboratory directorship, location, ownership, name, insolvency, or bankruptcy; notification must occur no later than 30 days prior to the change(s); or, in the case of unexpected changes, no later than two working days afterwards. Laboratories subject to US regulations must also notify the US Department of Health and Human Services.
In addition, the policy must address:

6. Provision of a trained inspection team comparable in size and scope if requested by CAP at least once every two-year accreditation period
7. Cooperation with CAP and HHS when the laboratory is subject to a CAP or HHS complaint investigation or validation inspection
8. Adherence to the Terms of Use for the CAP Certification Mark of accreditation
9. For laboratories subject to US regulations, availability, on a reasonable basis of the laboratory’s annual proficiency testing results upon request of any person

Evidence of Compliance:
✓ Records of notification, if applicable

GEN.30000 Monitoring Analytic Performance

There is a written quality control program that clearly defines policies and procedures for monitoring analytic performance.

NOTE: There must be a written overall quality control program for the entire laboratory. It must include general policies and assignment of responsibilities. There must be clearly defined, written procedures for ongoing monitoring of analytic performance, including (1) number and frequency of controls; (2) establishment of tolerance limits for control testing; and (3) corrective actions based on quality control data. Quality control records should be well-organized with a system to permit regular review by appropriate supervisory personnel (laboratory director, supervisor or laboratory quality control coordinator).

SPECIMEN COLLECTION, HANDLING, AND REPORTING

Specimen collection, handling, and results reporting are critical. Specific instructions for the proper collection and handling of specimens must be made available to laboratory personnel and to anyone collecting patient test materials that are sent to the laboratory.

Inspector Instructions:

- Follow a patient specimen beginning with test ordering through patient identification, phlebotomy/collection, labeling, transport, receipt and processing, delivery to test area, analysis, result review, and reporting. Determine if practice matches related policies and procedures.
SPECIMEN COLLECTION INSTRUCTIONS

Inspector Instructions:

- Sampling of specimen collection policies and procedures
- Specimen handling policies and procedures for referral of testing
- Specimen collection manuals (available)

**REVISED** 07/28/2015
GEN.40016 Specimen Collection Procedure Review Phase II
There are records of review of the specimen collection/handling procedures by the current laboratory director or designee at least every two years.

**REVISED** 07/28/2015
GEN.40032 New Specimen Collection Procedure Review Phase II
The laboratory director reviews and approves all new specimen collection and handling procedures, as well as substantial changes to existing procedures before implementation.

NOTE: Current practice must match written procedures.

GEN.40050 Distribution of Manuals Phase I
The specimen collection manual is distributed to all specimen-collecting areas within the hospital (nursing stations, operating room, emergency room, out-patient areas) AND to areas outside the main laboratory (such as physicians’ offices or other laboratories).

NOTE: It is acceptable for this information to be electronically available to users rather than in book format; there is no requirement for a paper-based specimen collection manual. Indeed, electronic manuals have the advantage of more accurately reflecting current requirements.

REFERENCES

GEN.40100 Specimen Collection Manual Elements Phase II
The specimen collection manual includes instructions for all of the following elements, as applicable.

1. Preparation of the patient
2. Type of collection container and amount of specimen to be collected
3. Need for special timing for collection (e.g. creatinine clearance)
4. Types and amounts of preservatives or anticoagulants
5. Need for special handling between time of collection and time received by the laboratory (e.g. refrigeration, immediate delivery)
6. Proper specimen labeling
7. Need for appropriate clinical data, when indicated

NOTE: Because of the importance of clinical information in the practice of surgical pathology and cytopathology, requisitions for such specimens should include pertinent clinical data, as well as pre-operative and/or post-operative diagnosis. Written instructions should be available for all applicable tissue and cytologic specimens, including biopsies, resections, PAP tests, sputum washings, brushings, body fluids, fine needle aspirations, etc. Instructions must include proper fixation of slides and tissue specimens. A variety of tests in clinical pathology also require specific clinical information (e.g. maternal AFP screening, TDM peak and trough measurements, and antibiotic therapy) or special instructions for collection, preservation, and storage (e.g. timed or 24-hour urine specimens).

REFERENCES
2) Burton JL, Stephenson TJ. Are clinicians failing to supply adequate information when requesting a histopathological investigation? J Clin Pathol. 2001;54:806-808

**REVISED** 07/28/2015

GEN.40125 Handling of Referred Specimens Phase II

For specimens sent to referral laboratories, the referring laboratory properly follows all requisition, collection and handling specifications of the referral laboratory.

NOTE: Pre-analytic variables must be closely controlled to maintain specimen integrity. These include specimen temperature, transport time, and the interval before separation of blood cells from serum/plasma. For coagulation tests, important considerations include proper filling of the collection tube, the use of waste tubes, and, if blood must be drawn through an indwelling line, flushing of the line. For surgical pathology and cytopathology, specimens must be preserved by proper fixation or refrigeration. Twenty-four-hour urine specimens may require special preservatives for specific tests. Also, it may be necessary to collect specific patient information required by the testing laboratory (e.g. menstrual history for cytopathology, gestational age for prenatal neural tube defect screening, preoperative diagnosis for surgical pathology, and bleeding history for specialized coagulation assays).

For microbiology specimens, guidelines for the timing of specimen collection, collection techniques, and selection of appropriate collection devices and transport media must be followed as stipulated by the referral laboratory.

For newborn screening specimens, the specimen collection, application and drying of blood spots, and submission of specimens to the referral laboratory must follow the designated newborn screening laboratory’s instructions and be in compliance with the most recent edition of the CLSI Document NBS01 and state or local regulations. Specimens should be transported after they are dry and no later than 24 hours after collection or following the instructions provided by the designated newborn screening laboratory. Delays in specimen transportation from the collection facility to the testing laboratory may compromise the integrity of the specimen and results and could critically impact the newborn.

Evidence of Compliance:
✓ Written procedure for submission of specimens to referral laboratories, consistent with the referral laboratory collection and handling requirements

REFERENCES
SPECIMEN COLLECTION AND LABELING

Accurate and precise laboratory data are dependent on properly collected clinical specimens.

Inspector Instructions:

- Specimen collection (patient identification, specimen labeling, correction of labeling, and adverse event) policies and procedures
- Sampling of phlebotomy/clinical specimen collection training records
- Paternity/forensic collection policies and procedures

- Sampling of phlebotomy supplies, collection devices, transport media (expiration date, storage)
- Specimen collection at one or more sites within the institution.

- How is feedback related to specimen quality provided to the individuals collecting patient specimens, including non-laboratory staff, as applicable?

- If specimen collection errors are a recurring problem, further evaluate the laboratory’s investigation of how the errors occurred and the corrective actions that were implemented

GEN.40460 Specimen Collection Supplies Phase II

Specimen collection supplies such as blood collection tubes and collection devices (e.g. heel lancets, culture swabs, and transport media) are used within their expiration date and stored per manufacturer's instructions.

NOTE: For newborn screening collection cards, if the expiration date is not printed on the individual cards, another mechanism, such as serial number, may be used for tracking.

REFERENCES
1) Clinical and Laboratory Standards Institute (CLSI). Blood Collection on Filter Paper for Newborn Screening Programs. CLSI Standard NBS01-A6. Clinical and Laboratory Standards Institute, 940 West Valley Road, Suite 2500, Wayne, Pennsylvania 19087-1898 USA

**REVISED** 07/28/2015

GEN.40470 Specimen Collection Training Phase II

There are records that all personnel collecting patient specimens have been trained in collection techniques and in the proper selection and use of equipment/supplies and are knowledgeable about the contents of the specimen collection procedures.
NOTE: This applies to laboratory employees, including those at remote sites that are owned and operated by the laboratory.

It applies to all personnel who collect and test samples under the laboratory’s CAP number, such as for point-of-care testing and for blood gas analysis. It does not apply to the collection of specimens sent to the laboratory by hospital personnel or from outside sources. All types of specimen collection techniques (e.g. phlebotomy, capillary, arterial, in-dwelling line, phlebotomy during intravenous infusion), as well as non-blood specimens, must be included in the training in accord with the individuals’ duties. If the laboratory uses prepackaged kits for specimen collection, any special instructions that accompany the kit must be part of the training.

REFERENCES

1) Galen HJ. Complications occurring from diagnostic venipuncture. J Fam Pract. 1992;34:582-584
6) Strasinger SK, di Lorenzo MA. Phlebotomy handbook, 2nd ed. Norwalk, CT: Appleton & Lange, 1996
21) Burns ER, Yoshikawa N. Hemolysis in serum samples drawn by emergency department personnel versus laboratory phlebotomists. Lab Med. 2002;33:378-380

**REVISED** 08/17/2016

Patient Identification

The individual collecting the specimen positively identifies the patient before collecting a specimen and labels the specimen in the presence of the patient.

NOTE: Personnel must confirm the patient’s identity by checking at least two identifiers before collecting a specimen. For example, an inpatient’s wristband may be checked for name and unique hospital number; an outpatient’s name and birth date may be used. The patient’s room number may not be used as an identifier. The patient’s identity should be verified by asking the patient to identify him- or herself, when it is practical to do so*. The intent of this requirement is to ensure a written, consistently followed system for correct patient and specimen identification at the point of collection.

*For example, verbal verification is not necessary if obtaining the services of a translator would delay specimen collection.

Evidence of Compliance:

✓ Written collection procedure, including criteria for patient identification

REFERENCES

All primary specimen containers are labeled with at least two patient-specific identifiers.

NOTE: A primary specimen container is the innermost container that holds the original specimen prior to processing and testing. This may be in the form of a specimen collection tube, cup, syringe, swab, slide or other form of specimen storage. Data files received from other laboratories for analysis are considered a specimen and must contain acceptable patient identifiers. Criteria for acceptable specimen labeling and the handling of sub-optimal specimens must be defined in laboratory policy.

Examples of acceptable identifiers include but are not limited to: patient name, date of birth, hospital number, social security number, requisition number, accession number, unique random number. A location (e.g. hospital room number) is not an acceptable identifier. Identifiers may be in a machine readable format, such as a barcode.

In limited situations, a single identifier may be used if it can uniquely identify the specimen. For example, in a trauma situation where a patient's identification is not known, a specimen may be submitted for testing labeled with a unique code that is traceable to the trauma patient. Other examples may include forensic specimens, coded or de-identified research specimens, or donor specimens labeled with a unique code decryptable only by the submitting location.

Obtaining uniform compliance with this requirement may be difficult when specimens are collected by non-laboratory personnel. The laboratory should 1) Provide a list of acceptable identifiers to all specimen collectors; 2) Communicate with specimen collectors regarding the importance of this requirement; and 3) Have a procedure for following up with specimen collectors when inadequately labeled specimens are received. Communication and follow-up may be through QM reports, written memoranda, phone calls, visits by client service personnel, or other means of disclosure.

Evidence of Compliance:
✓ Written policy with criteria for acceptable labeling of primary specimen containers AND
✓ Specimen collection procedures with defined labeling specifications AND
✓ Records of audits for compliance with specimen labeling policies and procedures

REFERENCES
The laboratory has a written policy regarding correction of information on specimen labels.

NOTE: If laboratory personnel become aware of a potential error in patient identification or other information (e.g. initials of individual collecting the specimen, date/time of collection) on a specimen label, best practice is to recollect the specimen. However, there may be circumstances when recollection is not possible or practical (e.g. for specimens that are impossible or difficult to recollect, such as cerebrospinal fluid). The laboratory should define the circumstances under which correction of the information on specimen labels is permitted. A record of all such corrections should be maintained. The laboratory should investigate errors in specimen labeling, and develop corrective action as appropriate, including education of personnel who collect specimens.

Evidence of Compliance:
✓ Records of corrections to specimen labels and corrective action

REFERENCES

GEN.40493 Specimen Labeling for Pretransfusion Testing Phase II

All blood specimens collected for pretransfusion testing are labeled at the time of specimen collection in the presence of the patient with:

1. Patient's first and last name
2. Unique identification number
3. Date of collection
4. A method to identify the individual collecting the specimen

NOTE: Blood specimens collected for pretransfusion testing must be positively and completely identified and labeled before leaving the patient. Acceptable practices for positive identification of patient and blood specimen labels must be defined in the procedure manual and may include visual inspection and/or an electronic system to read the identifying information contained in barcodes or radio-frequency identification (RFID) microchips or the patient's wristband. Acceptable practices for generating specimen labels must be defined in the procedure manual and may include electronic devices utilizing information encoded in barcodes or RFID microchips. There must be a dependable method to identify the individual who collected the blood specimen, such as initials or another identifier on the tube, or an electronic record.

Evidence of Compliance:
✓ Written procedure defining labeling requirements of specimens for pretransfusion testing
AND
✓ Written procedure defining system identifying the individual collecting pretransfusion testing specimens

REFERENCES
3) Sandler SG, Langeberg A, Carty K, Dohnalek LJ. Bar codes and radio-frequency technologies can increase safety and efficiency of blood transfusions. LabMedicine 2006;37:436-439

GEN.40497 Paternity/Forensic Data Phase II

If the laboratory collects specimens for paternity/forensic identity testing, the following data are obtained:

1. Place and date of specimen collection
2. Identity of person collecting the specimen
3. Photograph, or photocopy of a picture identification card for each individual tested
4. Signed record of information (including name, race, relationship) for each individual tested
5. Date of birth of child
6. Synopsis of case history/investigation, sample source
7. Record of informed consent

NOTE: If the laboratory uses prepackaged kits for specimen collection, any additional instructions that accompany the kit must be followed.

REFERENCES
1) Standards for Parentage testing laboratories. American Association of Blood Banks. Standards for parentage testing laboratories. Bethesda, MD: 2003:5.2.4

GEN.40498 Specimen Labeling - Paternity/Forensic ID

For paternity/forensic identity testing, the information about each individual and the accuracy of the specimen label is verified by that individual or the legal guardian.

Evidence of Compliance:
✓ Records of information and label verification by patient or legal guardian

GEN.40505 Specimen Collection Feedback

There is a mechanism to provide feedback to the collectors of specimens on issues relating to specimen quality and labeling.

NOTE: The accuracy of an analytic result depends upon the initial quality of the specimen. Proper collection techniques are essential.

Evidence of Compliance:
✓ Written procedure defining methods for providing feedback to specimen collectors AND
✓ Records of communication of specimen collection issues, such as QM reports, staff meeting minutes OR records of employee counseling

GEN.40508 Phlebotomy Adverse Reaction

The laboratory has procedures to care for patients who experience adverse reactions from phlebotomy.

NOTE: Minor adverse reactions include hematomas, abrasions, nausea, and fainting. Serious injuries include vomiting, nerve damage, seizures and injuries. Training of phlebotomists should emphasize injury prevention. Serious reactions must be recorded in an incident log.

Evidence of Compliance:
✓ Written instructions to phlebotomists AND
✓ Training records

SPECIMEN TRANSPORT AND TRACKING

This section applies to laboratories that send specimens to referral or other laboratories for testing, whether or not the specimen collection is performed by the laboratory staff. It also applies to referral laboratories that receive specimens from other laboratories or remote locations outside of the facility for testing.
While transportation of clinical specimens may not be the responsibility of personnel under the control of the laboratory director, issues of tracking and specimen quality must be addressed to ensure quality laboratory results.

Inspector Instructions:

- Sampling of specimen packing and shipping policies and procedures
- Sampling of packaging and shipping of infectious materials training records

- How do you know specimens sent from remote sites are actually received?
- What is your course of action when specimens received from remote sites are unacceptable?

**REVISED** 07/28/2015

**GEN.40511** Specimen Tracking/Labeling

All specimens are properly packaged and labeled to indicate the general nature of the materials transported.

Evidence of Compliance:

✓ Written procedure defining criteria for packaging and labeling

REFERENCES


4) Tarapchak P. In 'shipping' shape. Advance/Lab. 2000;9(7):48-59

**GEN.40512** Infectious Material Packing/Shipping

The laboratory packages and ships infectious material in accordance with applicable federal, state and local regulations.

Evidence of Compliance:

✓ Written procedures for packaging and shipping that comply with regulations

REFERENCES


**GEN.40515** Transport Personnel Training

Transport personnel are trained in appropriate safety and packaging procedures suitable to specimen type and distances transported, including training for personnel involved in packaging and shipping infectious substances.

NOTE: Training should include issues such as adherence to regulations for transport of biohazards, use of rigid containers where appropriate, temperature control, notification procedures in case of accident or spills, etc.
All personnel who package infectious specimens for shipment must satisfactorily complete training in these requirements. Federal and international regulations mandate the proper packaging and transportation of infectious substances, also termed "etiologic agents." It is the laboratory's responsibility to determine whether specimens that are to be shipped are subject to the regulations, or are exempt. For US laboratories, specific requirements are set forth by the US Public Health Service, the US Department of Transportation and the US Postal Service. These apply to domestic transportation by land, air or sea, and to international air transportation. Recurrent training is required every 3 years. The laboratory should check with its local department of transportation or state health department for any recent revisions to these requirements.

Laboratories outside of the US must comply with their national regulations.

These requirements for packaging and shipping of infectious substances do not apply to private couriers.

The laboratory may send personnel to courses for training, or may obtain materials to train its personnel in-house. Resources for training are available from many sources, including state health departments, vendors of shipping materials, and the CDC National Laboratory Training Network (NLTN).

Evidence of Compliance:
✓ Records of training for all personnel involved in transport of specimens

REFERENCES
1) Title 49, Code of Federal Regulations, Part 172.704 Training Requirements

**REVISED** 07/28/2015

GEN.40530 Specimen Tracking Phase II

For specimens submitted to the laboratory from remote sites, there is a tracking system and record to ensure that all specimens are actually received.

NOTE: Records should include time of dispatch and receipt, as well as condition of specimens upon receipt. An example of an acceptable tracking system is submission of a packing list (prepared by the client or courier) with each batch of client specimens, which may be checked against the specimens received by the laboratory. Some laboratory tests (e.g. coagulation assays) have limitations on time and temperature conditions between collection and analysis. This requirement applies to couriers/transportation systems that are within the laboratory organization or are contracted by it. It does not apply to couriers unrelated to the laboratory.

Evidence of Compliance:
✓ Specimen shipping/transport logs AND
✓ Records of follow up for specimens not received

GEN.40535 Specimen Transport QM Phase I

There is a process for monitoring the quality of submitted specimens, correcting problems identified in specimen transportation, and improving performance of clients or sites that frequently submit specimens improperly.

Evidence of Compliance:
✓ Records of corrective action OR communications with clients that frequently submit specimens incorrectly

GEN.40545 Newborn Screening Specimen Tracking Phase I
For specimens being submitted to a remote testing laboratory for newborn screening for congenital disorders, there is a tracking system and records to ensure that all specimens are submitted in compliance with timing requirements and that a result or other appropriate notification is received indicating that the specimens were actually received.

**NOTE:** Tracking records should include time of dispatch and condition of specimens upon submission. An example of an acceptable tracking system is the use of a packing list (prepared by the submitting site or courier) with each batch of specimens that is checked against the specimens received by the remote testing laboratory. Newborn screening laboratory specimens have limitations with time and humidity conditions between collection and analysis. This requirement applies to couriers/transportation systems that are part of the laboratory organization and to outside courier systems.

**Evidence of Compliance:**
- Records showing results/notifications received on all specimens **AND**
- Records of follow up for specimens not received at the remote laboratory

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**REQUISITIONS AND SPECIMEN RECEIPT/HANDLING/PROCESSING**

**Inspector Instructions:**

- Sampling of specimen receipt and handling policies and procedures
- Sampling of specimen requisitions
- Sampling of temperature logs (refrigerator, freezer)

- How do you know what date/time a specimen is received in your laboratory? How are specimens accessioned once received by the laboratory?
- What is your course of action regarding verbal orders?
- How do you know your specimen containers do not contribute to analytic interference?

- If lost specimens are a recurring problem, further evaluate the laboratory’s investigation of where/how in the process the specimen was lost and the corrective actions that were implemented

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**GEN.40700  Requisitions  Phase II**

**All specimens are accompanied by an adequate requisition.**

**NOTE:** In computerized settings, there may not be a paper requisition that is physically attached to the specimen container.

**REFERENCES**

Test requisition data elements are entered accurately into the laboratory information or record system.

NOTE: Data elements include patient demographic data; the name and location of the individual or entity ordering the test, as well as other elements needed for the final report (see GEN.41096). The laboratory must have an ongoing mechanism to ensure the accuracy of manual entries. For test orders crossing an interface to the LIS, requirements for interface integrity apply.

REFERENCES

**REVISED** 07/28/2015

GEN.40750 Requisition Elements

The paper or electronic requisition includes all of the following elements, as applicable.

1. Adequate patient identification information (e.g., name, registration number and location, or a unique confidential specimen code if an alternative audit trail exists)
2. Patient sex
3. Patient date of birth or age
4. Name and address (if different than the receiving laboratory) of the physician, legally authorized person ordering the test, or name and address of the laboratory referring the specimen
5. Tests requested
6. Last menstrual period (for gynecologic specimens)
7. Date of specimen collection, and if appropriate, time of collection
8. Source of specimen, when appropriate
9. Clinical information, when appropriate

NOTE: Specimen source may be particularly important for microbiology, surgical pathology, and cytopathology specimens. Surgical pathology specimens must be labeled and requisitions prepared in the room where the surgical procedure is performed. The patient's chart or medical record may be used as the test requisition or authorization.

REFERENCES

GEN.40825 Specimen ID

There is a system to positively identify all patient specimens, specimen types, and aliquots at all times.

NOTE: Each specimen container must identify the patient uniquely. This may be text-based, numeric, bar-coded, etc. The form of this system is entirely at the discretion of each laboratory, so long as all primary collection containers and their aliquots have a unique label which one can audit back to full particulars of patient identification, collection date, specimen type, etc. Practical considerations of container size may limit the extent of such details. There must be an appropriate, consistently applied accessioning system.

REFERENCES

GEN.40900 Specimen Date Received

Phase II
The date (and time, if appropriate) that the specimen was received by the laboratory is recorded.

REFERENCES
1) Department of Health and Human Services, Centers for Medicare and Medicaid Services. Clinical laboratory improvement amendments of 1988; final rule. Fed Register. 2003(Oct 1);1034 [42CFR493.1242(b)]

GEN.40930 Authorized Requestor

The laboratory has a mechanism to ensure that specimens are analyzed only at the request of an authorized person.

NOTE: The laboratory must perform tests only at the written or electronic request of an authorized person. In some US states and other countries, individuals may order some laboratory tests without a physician's referral (direct-to-consumer testing).

Evidence of Compliance:
✓ Written policy requiring test orders by authorized persons, if applicable in the jurisdiction in which the laboratory is located

REFERENCES

GEN.40932 Verbal Test Authorization

For laboratories subject to US regulations, the laboratory solicits written or electronic authorization for verbal orders within 30 days.

NOTE: The laboratory must retain the written authorization or record of efforts made to obtain a written authorization. In a managed office where the staff assistants are not employees of the physician/clinician, the staff should not sign a test requisition for the physician without some type of provider services agreement. This agreement must specify how the clinician has accepted responsibility for the tests ordered from the off-site laboratory. (This situation is different from the hospital environment, where the physician has personally signed the order sheet.)

Evidence of Compliance:
✓ Records of follow-up to obtain written order

REFERENCES
1) Department of Health and Human Services, Centers for Medicare and Medicaid Services. Clinical laboratory improvement amendments of 1988; final rule. Fed Register. 2003(Jan 24);7162 [42CFR493.1241(a),1241(b)]

GEN.40935 Test Order Read Back

The laboratory has a policy that personnel receiving verbal or phone orders read back the entire order to verify accuracy of transcription.

GEN.40938 Unclear Test Order

The laboratory has a policy on confirmation of test orders that may be unclear (e.g. orders using non-standard or non-specific terms).

**REVISED** 07/28/2015
GEN.40942 Specimen Container Analytic Interference

Phase I

Phase II
The laboratory director or designee evaluates significant changes to specimen containers to ensure that they do not contribute to analytic interference in the assays to be performed and approves them for use.

NOTE: Significant changes include new container types, a different container type (e.g. a plain container to one with an additive), and when changing to a different vendor. To ensure that the specimen containers do not contribute to analytic interference, the laboratory director or designee must review clinical literature, as available, and evaluate information from specimen collection container and instrument/method manufacturers. Based on the information reviewed and the test systems that will be impacted, the laboratory director or designee determines whether verification by the laboratory is indicated.

Manufacturers of collection containers must perform studies to demonstrate safety and efficacy of the product prior to marketing their products. However, it is not feasible for manufacturers to evaluate all assays on all instrument and methods. The CLSI Guideline GP34-A, Validation and Verification of Tubes for Venous and Capillary Blood Specimen Collection, recommends performing a comparative tube evaluation when changing to a different type of tube (e.g. gel, additive, different vendor). A sample protocol for end user evaluation is provided in the CLSI guideline.

Evidence of Compliance:

✓ Records of specimen container evaluation for analytic interference with approval for use

REFERENCES

15) Gaillard C, Strauss F. Eliminating DNA loss and denaturation during storage in plastic microtubes. Am Clin Lab. 2001;20(2);52-54
18) Hashim IA. Blood samples collected in serum-separator tubes give higher free triiodothyronine levels using the Immulite assay. Clin Chem. 2001;47(suppl);A14
Evidence of Compliance:
✓ Records of verification of operating speeds at least annually

REFERENCES

**REVISED** 08/17/2016
GEN.41042 Refrigerator/Freezer Temperatures Phase II

Refrigerator/freezer temperatures are checked and recorded daily using a calibrated thermometer.

NOTE: This checklist requirement applies to refrigerators/freezers containing reagents or patient/client specimens. “Daily” means every day (7 days per week, 52 weeks per year). The laboratory must define the acceptable temperature ranges for these units. If temperature(s) are found to be outside of the acceptable range, the laboratory must record appropriate corrective action, which may include evaluation of contents for adverse effects.

Temperatures may be recorded either manually, or using a recording device or system by: 1) recording the numerical temperature, or 2) placing a mark on a graph that corresponds to a numerical temperature. If temperatures are recorded manually, the identity of the individual recording the temperature(s) must be recorded (the initials of the individual are adequate).

If an automated (including remote) temperature monitoring system is used instead of manual temperature monitoring, laboratory personnel must have ongoing immediate access to the temperature data, so that appropriate corrective action can be taken if a temperature is outside of the acceptable range. System records must demonstrate the daily functionality of the system.

If a minimum/maximum thermometer is used to perform continuous monitoring of temperatures between daily temperature readings or following a laboratory downtime (e.g. laboratory closure for weekend or holiday), both the low and high temperatures must be recorded. To ensure correct temperature readings, the minimum/maximum thermometer device must be reset prior to the monitoring period.

A frost-free freezer may be used to store reagents and controls provided that the function of these materials is not compromised. Storage conditions must remain within the specifications of the manufacturer of the reagent or control. Temperatures may be recorded using a continuous monitoring system or a maximum/minimum thermometer. Thermal containers within the freezer may be used.

Patient samples may be stored in frost free freezers only if protected from thawing. The laboratory must maintain records showing that the temperatures stay within the defined range.

REFERENCES

RESULTS REPORTING AND REFERRAL OF TESTING

The laboratory must provide useful clinical data. Data must be legible, accurate, reported in clearly designated units of measurement, and promptly reported to persons authorized by law to receive and use medical information.

A referral laboratory is any outside location to which the referring laboratory submits specimens or material for testing [CLSI guideline QMS05-A2]. In the requirements that follow, the phrase “referral laboratory” means an independent, external enterprise. The phrase “off-site location” is used when part of the testing essential for a final result is performed at a closely affiliated or satellite laboratory. Off-site locations include offices where images or data files are reviewed and interpreted with frequency (i.e. recurrent or on a regular basis). The
addition of an electronic signature to a final report is not off-site laboratory testing, nor is the rendering of a consultative opinion that does not involve a specimen submitted for testing.

Inspector Instructions:

- Sampling of reporting policies and procedures
- Sampling of paper or electronic laboratory reports
- Sampling of referral laboratory's patient reports
- Patient confidentiality policies and procedures

- How does the laboratory director ensure that the content of laboratory reports effectively communicates patient test results?
- How does the laboratory protect patient information?
- What is your course of action if laboratory testing is delayed? How frequently does this occur?
- What is your process for selecting referral laboratories?
- How does your laboratory determine who is authorized to receive results?
- How does your laboratory archive test results for comparison with later results?

- If instances of delayed test reporting are frequent, further evaluate laboratory director leadership's investigation, corrective actions, and resolution

GEN.41067  Content/Format Report Review  Phase I

An individual meeting CAP laboratory director qualifications reviews and approves the content and format of paper and electronic patient reports at least every two years.

NOTE: The laboratory director (or a designee who meets CAP qualifications for laboratory director) must review and, at least every two years, approve the content and format of laboratory patient reports (whether paper or computer screen images) to ensure that they effectively communicate patient test results, and that they meet the needs of the medical staff. Further details on review of electronic reports are given in GEN.48500.

GEN.41077  Reporting Outside Results  Phase I

There is a policy for laboratory director input regarding whether outside laboratory results are reported through the primary reporting system (i.e. the laboratory information system or the institution's electronic medical record).

NOTE: At times patients may bring test results from outside laboratories to their physicians. Patients' physicians may request, or institutional policy may dictate, that such results (or other test results from outside laboratories) be integrated into the laboratory's primary reporting system (i.e. the LIS or the institution's electronic medical record). The laboratory director should be aware of whether results from outside laboratories are reported through the laboratory information system or the electronic medical record system. It is recognized that the laboratory director may not be in a position to prohibit entry of outside laboratory results into the electronic medical record system.

However, if such results are integrated, the name and address of the outside laboratory must be available in the primary reporting system, and there must be an indication available to the person viewing such results that the results originated from an outside laboratory. Criteria for inclusion of such results might include whether the quality of the outside laboratory has been evaluated by
the laboratory director; CLIA licensure or equivalent; whether reference ranges and/or units of measurement differ from in-house tests; whether units of measurement and reference range are included; and possession of an official report.

If the laboratory director believes that certain test results should not be integrated into the primary reporting system, one option is to include such results in a section of the electronic medical record other than the laboratory database.

**REVISED** 07/28/2015
GEN.41096 Report Elements Phase II

The paper or electronic report includes the following elements.

1. Name and address of testing laboratory (see note below)
2. Patient name and identification number, or unique patient identifier
3. Name of physician of record, or legally authorized person ordering test, as appropriate
4. Date of specimen collection, and if appropriate, time of collection
5. Date of release of report (if not on the report, this information should be readily accessible)
6. Time of release of report, if applicable (if not on the report, this information should be readily accessible)
7. Specimen source, when applicable
8. Test result(s) (and units of measurement, when applicable)
9. Reference intervals, as applicable
10. Conditions of specimen that may limit adequacy of testing

NOTE: All of the above data elements, as applicable, must be available in the laboratory information system or in paper records, and must be in the report that is available/sent to the clinician, whether electronic or paper, including electronic reports in systems interfaced to the laboratory information system directly or through middleware or an interface engine. (For electronic reports, data elements need not all be present on one screen, but must be readily available.)

The paper or electronic report must include the name and address of referral laboratories where patient testing was performed. For laboratories subject to US regulations, a “referral laboratory” includes outside referral laboratories as well as any affiliated or special function laboratory that is separately accredited and has a different CLIA registration number than the referring laboratory. For electronic reports, the name and address of referral laboratories need not all be present on the same screen(s) as the results but must be available to the clinician in the information system.

Under some circumstances it may be appropriate to distribute lists or tables of reference intervals to all users and sites where reports are received. This system is usually fraught with difficulties, but if in place and rigidly controlled, it is acceptable.

Patient reports must state the name of the physician (or other legally authorized person) ordering the test(s) or a physician of record. In those institutions where there are multiple ordering physicians and/or frequent changing of attending physicians, the ordering physician should be easily identifiable through a computer audit trail or other records of the test order.

Referral laboratories accredited by the CAP must provide a copy of the results to the referring laboratory (Exceptions to this requirement may be made under special circumstances or for special categories of testing, such as drugs of abuse or employee drug testing. The laboratory director may make these exceptions.). Results may be reported to the ordering physician of record (or other legally authorized person) by either the referral laboratory or the referring laboratory.

REFERENCES
GEN.41300 Report Retention and Retrieval

Copies or files of reports are legible and retained by the laboratory in a manner that permits prompt retrieval of the information.

NOTE: The length of time that reported data are retained in the laboratory may vary; however, the reported results must be retained for that period encompassing a high frequency of requests for the data. In all circumstances, a hospital laboratory must have access to the patient’s chart where the information is permanently retained.

REFERENCES

**REVISED** 07/28/2015

GEN.41303 Patient Confidentiality

The laboratory ensures that internal and external storage and transfer of data maintains patient confidentiality and security.

NOTE: Written procedures must address patient confidentiality during transfer of data to external referral laboratories or other service providers. This must include cloud based computing (e.g. for storage of confidential data), as appropriate

The laboratory must audit compliance with the procedures at least annually.

Evidence of Compliance:
✓ Records of patient privacy audit for compliance with the Health Insurance Portability and Accountability Act (HIPAA)

REFERENCES

**REVISED** 07/28/2015

GEN.41304 Patient Data Accessibility

There is a written policy to ensure that patient data are accessible in a timely manner only to those individuals who are authorized to review test results.

NOTE: Only those healthcare personnel authorized to review a patient's test results should have access to those results. Laboratories subject to US regulations must provide final test results to the patient or the patient's personal representative upon request. For completed tests, these results must generally be provided no later than 30 days after such a request.

Under the HIPAA Privacy Rule, only the patient or a personal representative, defined as an individual who has authority under applicable law to make health care decisions for the patient, can be given access to a patient's personal health data. Laboratories must take reasonable steps to verify the identity of the patient and the authority of a personal representative to have access to an individual's protected health information. The Rule also allows for the release of test reports to authorized persons responsible for using the test reports and to the laboratory that initially requested the test, if applicable.
For additional information see Department of Health and Human Services, Medicare and Medicaid Services, "CLIA Program and HIPAA Privacy Rule; Patients' Access to Test Reports; Final Rule." Fed Reg 79:7290 (2014); 45CFR164.502(g); 45CFR164.514.

REFERENCES

**REVISED** 08/17/2016

GEN.41306 Analyst Tracking ID  Phase II

There is a system whereby the identity of the analyst performing or completing the test and the date of the test can always be established.

NOTE: If results are released using autoverification, the system must be capable of identifying those test results that have been autoverified. In addition, the laboratory should be able to identify the technologist responsible for the instrument producing the result, such as through daily bench assignment charts, instrument set-up logs, or electronic audit trail.

REFERENCES

GEN.41307 Report Errors  Phase II

When errors are detected in patient test reports, the laboratory promptly notifies responsible clinical personnel or referring laboratory as applicable and issues a corrected report.

NOTE: Notification should include the department of health or other legal entity as required by local regulations.

Evidence of Compliance:
✓ Records of report error notification and corrected report

REFERENCES

**REVISED** 08/17/2016

GEN.41310 Corrected Report  Phase II

All corrected reports of previously reported, incorrect patient results are identified as corrected, and both the corrected and original data are clearly identified as such.

NOTE: 1. As clinical decisions or actions may have been based on the previous report, it is important to replicate previous information (test results, interpretations, reference intervals) for comparison with the corrected information. The previous information and the corrected information must be identified as such, and the original data must be present in the corrected report (for paper reports), or linked electronically or logically to the corrected information (in electronic reports).

2. This requirement applies to electronic reports in the laboratory information system and to the data systems interfaced to the laboratory information system either directly or through middleware or an interface engine (but not to systems that are further downstream in the interface chain).

3. Displays in an electronic medical record (EMR) downstream from the laboratory should include the original report as well as the corrected report. The report elements listed in GEN.41096 should be included in the EMR.
4. The correction should add explanatory language if an explanation would be helpful to the user. For example, a comment about transport or sample storage conditions uncovered post-analysis can help frame an original, invalid result.

5. For changes to anatomic pathology and cytopathology reports, refer to ANP.12185 and CYP.06475.

REFERENCES

GEN.41312 Multiple Corrections

Phase II

When there are multiple sequential corrections of a single test result, all corrections are referenced in sequential order on subsequent reports.

NOTE: When there are multiple sequential corrections of a previously reported result, it is considered inappropriate to note only the last correction made, as the clinician may have made a clinical decision based upon erroneous data rather than the "true" result. All corrections should be referenced in the patient report.

GEN.41316 Infectious Disease Reporting

Phase II

There is a policy regarding the timely communication, and documentation thereof, of diagnoses of infectious diseases of particular significance (e.g. human immunodeficiency virus and tuberculosis).

NOTE: The laboratory should have a policy to ensure that diagnoses of human immunodeficiency virus infection and other serious infections (for example, tuberculosis) are communicated to the responsible clinician in a timely manner.

The intent of this checklist item is NOT to require that these diagnoses be treated as critical results (this decision is up to the laboratory director); rather, the intent is that the laboratory assures that its reporting system is effective.

GEN.41325 Newborn Screening Results

Phase II

There must be a procedure for handling invalid and positive newborn screening results for samples submitted to other laboratories for testing.

NOTE: This requirement applies to the testing of whole blood heel stick samples from the newborn after birth on filter paper collection devices for the routine screening of congenital disorders. Positive results include those results that are outside of the expected range of testing results established for a particular condition. Invalid results include situations where the laboratory is unable to complete the screening process due to an unsuitable specimen, test, or incomplete information. Due to the urgent nature of newborn screening test results, procedures must include a process to track requests for repeat testing so that repeat specimens are submitted within the follow-up/recollection timeframe specified by the testing laboratory.

REFERENCES
1) Clinical and Laboratory Standards Institute. Newborn Screening Follow-up. CLSI document NBS02-A2. Clinical and Laboratory Standards Institute, 940 West Valley Road, Suite 1400, Wayne, Pennsylvania 19087-1898 USA

GEN.41345 Turnaround Time

Phase II

The laboratory has defined turnaround times (i.e. the interval between specimen receipt by laboratory personnel and results reporting) for each of its tests, and it has a policy for notifying the requester when testing is delayed.
NOTE: This does NOT imply that all instances of delayed reporting for all tests must lead to formal notification of clinical personnel. Rather, clinicians and laboratory must have a jointly agreed upon policy for when such notification is important for patient care.

Evidence of Compliance:
✓ Written policy defining test reporting turnaround time and process for communication of delays in turnaround time

REFERENCES

**REVISED** 08/17/2016

GEN.41350 Referral Laboratory Selection  Phase II

The laboratory has a written procedure for the selection and evaluation of laboratories to which it refers specimens or materials for testing.

NOTE:
1. The laboratory director, in consultation with the institutional medical staff or physician clients (where appropriate), is responsible for selecting referral laboratories
2. Selection of referral laboratories must be based primarily upon the quality of performance of such laboratories
3. Specimens or materials for testing include intermediate processing such as histologic and cytologic processing, preliminary analysis such as flow cytometry, and the use of distributive testing in next-generation sequencing. It also includes the referral of images or data files to an off-site location for interpretation.
4. For laboratories subject to US regulations: for tests in disciplines covered by CLIA, specimens and materials for testing must be referred only to a CLIA-certified laboratory or a laboratory meeting equivalent requirements as determined by CMS; this includes off-site locations where images or data files are frequently referred for review and interpretation. Laboratories that are part of the Department of Defense must meet the referral policies of the Clinical Laboratory Improvement Program (CLIP). With respect to patients on research protocols, whose tests are referred to a research laboratory: if those test results are used for patient management decisions, the research laboratory must be CLIA-certified, or meet equivalent requirements as determined by CMS.
5. For disciplines not covered by CLIA (e.g. histology), laboratories subject to US regulations must refer specimens to a laboratory accredited by CAP or a CAP-accepted organization.*
6. For non-US laboratories, whenever possible, specimens and materials for testing should be referred to a laboratory accredited by CAP; accredited to an established international standard from a recognized organization; or certified by an appropriate government agency. The inspector may need to exercise judgment with respect to determining if a referral laboratory is acceptable.
7. It is the responsibility of the laboratory director or designee to monitor the turnaround time and quality of test results received from referral laboratories.

*For overseas US military laboratories only, an exception to this requirement is acceptable if both of the following conditions are met:

1. Rapid turnaround time (TAT) is required to prevent either a delay in patient treatment/diagnosis or specimen degradation, and an acceptable TAT cannot be provided by a CAP-accredited or CLIA-certified laboratory.
2. The laboratory director has determined that the alternative testing site meets requirements that are equivalent to those of a CLIP or CLIA-certified laboratory as stipulated in the CLIP/CLIA Manual (11-32(8)c). This assessment must be recorded.

Evidence of Compliance:
✓ Records of the monitoring of referral laboratory services (e.g. problem log, review of reports)

REFERENCES
4) Brooks B. Cost considerations of esoteric testing. Advance/Lab. 1999;8(6):59-68
5) Carter JE, Bennett B. Laboratory "send out" test review by pathology house staff: cost-cutting strategy. Am J Clin Pathol. 1999;112:572

GEN.41430 Referral Laboratory Report Retention Phase II

For samples referred to another laboratory, the original or an exact copy of the testing laboratory’s report is retained by the referring laboratory.

NOTE: The report may be retained on paper or in electronic format. Exceptions to this requirement may be made under special circumstances or for special categories, such as drugs of abuse or employee drug testing. The laboratory director may make these exceptions.

Evidence of Compliance:
✓ Retained original referral laboratory reports OR direct access to referral laboratory reports via electronic transmission from the referral laboratory

REFERENCES

GEN.41440 Referral Laboratory Results Reporting Phase II

The essential elements of referred test results are reported by the referring laboratory as received from the referral laboratory, without alterations that could affect clinical interpretation.

NOTE: If the laboratory transcribes results from the referral laboratory report, the test result(s), interpretation, and information directly related to the interpretation must be copied as reported by the referral laboratory. This does not mandate that the referring laboratory report every word nor retain the exact format of the referral laboratory report. There is no requirement to fully replicate the complete content of the referral laboratory report beyond the results and interpretation. Suggestions for follow-up testing may, for example, be omitted at the discretion of the laboratory director.

Evidence of Compliance:
✓ Patient results from the referral laboratory consistent with laboratory-issued patient reports

REFERENCES
DIRECT-TO-CONSUMER TESTING

NOTE: Direct-to-consumer (DTC) tests are defined as tests that are requested or ordered by the consumer. All applicable requirements in other areas of the checklists apply to direct-to-consumer testing. This checklist section applies only to laboratories subject to US regulations. This checklist section does not apply to health fairs.

Inspector Instructions:

- Direct-to-consumer testing policies and procedures
- Sampling of direct-to-consumer laboratory reports

GEN.41460  DTC Jurisdiction

The laboratory performs DTC testing and reports results of DTC tests only in jurisdictions where such testing is lawful.

NOTE: No less than every two years, the laboratory must verify which jurisdictions permit DTC testing if it provides direct-to-consumer testing.

Evidence of Compliance:

✓  Record that the laboratory has reviewed applicable laws/regulations

GEN.41475  DTC Report

The test report includes test results, reference range, interpretation as applicable, and limitations of the test, as applicable, in language readily understandable by a lay person.

GEN.41485  DTC Report

The test report includes information that enables the consumer to contact a licensed health care professional about the clinical significance of the test result.

NOTE: This information may consist of the name, phone number, and email address of a health care professional. Alternatively, it may be the phone number of an office at the laboratory or medical center that can provide contact information to the consumer.

The practitioner or designee should be reasonably available during normal business hours.

GEN.41497  DTC Result Retention

The laboratory retains the results of DTC tests and reference ranges for at least 10 years after testing.

NOTE: This requirement applies only to DTC tests performed after June 15, 2009.
QUALITY OF WATER AND GLASSWARE WASHING

Inspector Instructions:

- Water quality policies and procedures
- Water quality test records
- How does your laboratory clean glassware?

GEN.41500 Defined Water Types

The laboratory defines the specific type of water required for each of its testing procedures and water quality is tested at least annually.

NOTE: The laboratory should define the type of water necessary for each of its procedures, and should have an adequate supply of same. The current edition of CLSI Guideline GP40-A4-AMD defines the following grades of water: Clinical Laboratory Reagent Water (CLRW), suitable for most laboratory procedures; Special Reagent Water (SRW), defined by a laboratory for procedures that need different specifications than CLRW; Instrument Feed Water, specified by IVD manufacturers as suitable for use with their measurement systems; and Commercially Bottled Purified Water that may be suitable for certain laboratory procedures.

CLRW is not required if the laboratory is able to record reliable results with an alternate grade of water.

The following specification for CLRW is adapted from this guideline and should be met at time of in-house production:

Bacteria may inactivate reagents, contribute to total organic contamination, or alter optical properties of test solutions. Resistivity provides a nonspecific measure of the ion content. Particulate matter includes organic carbon from biofilms and inorganic aggregates that can vary over time both in nature of the contamination and the effect on the laboratory use.

The CLSI Guideline provides testing information for microbial content, and resistivity, as well as total organic carbon; earlier specifications for silicates have been removed. It gives instructions for the preparation of the various types of water. It also addresses the use of purchased water, the effects of storing water, and the monitoring of stored water.

The quality (specifications) of the laboratory's water, whether prepared in-house or purchased, must be checked and recorded at least annually. The frequency and extent of checking may vary, according to the quality of source water and specific laboratory needs. Corrective action must be recorded if water does not meet acceptability criteria.

For CLRW, minimum monitoring includes resistivity and microbiology cultures. Other criteria, such as pH, endotoxin/pyrogens, silicates and organic contaminants are at the discretion of the laboratory, testing for these substances must be recorded only if the laboratory finds that they adversely affect specific test methods.

The laboratory must determine the level of testing necessary for other grades of water in use.
Typically, "sterile (pharmaceutical) water" is not manufactured to meet the specifications of CLRW, and should not be used as its equivalent.

For commercial instrument-reagent systems, the laboratory must use a specific type of water recommended by the manufacturer. Although routine commercial methods are typically designed to work with laboratory reagent grade water, higher-quality water systems exist and may be required for specific methods or if analytical imprecision or inaccuracy has been traced to the quality of in-lab water.

<table>
<thead>
<tr>
<th></th>
<th>CLRW</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maximum microbial content (CFU/mL)</td>
<td>10</td>
</tr>
<tr>
<td>Minimum resistivity (megohm-cm)</td>
<td>10 (in-line)</td>
</tr>
<tr>
<td>Particulate matter</td>
<td>0.22 um filter</td>
</tr>
</tbody>
</table>

Evidence of Compliance:
✓ Record of corrective action when water quality does not meet specifications

REFERENCES
8) Stewart BM. The production of high-purity water in the clinical laboratory. Lab Med. 2000;31:605-611

GEN.41770 Glassware Cleaning

Phase II

There are written procedures for handling and cleaning glassware, including methods for testing for detergent removal.

NOTE: Special instructions for micropipettes, cuvets, acid washing, etc. must be included.

A simple procedure to check for detergent residue uses brom cresol purple (0.1 g brom cresol purple in 50 mL ethyl alcohol). Pipette approximately 5 cm (2 inches) distilled water into a representative, washed, glassware item. Add two to three drops brom cresol solution. A purple color reveals residual detergent. A yellow color indicates satisfactory rinsing.

Evidence of Compliance:
✓ Records of detergent residue testing

REFERENCES

LABORATORY COMPUTER SERVICES

Multiple solutions for laboratory information systems (LIS) exist. Traditional systems have a local “host” database (i.e. the computer hardware and software) serving the information needs of the laboratory; the laboratory is the only “user.” In the current environment, the host is often physically remote from the laboratory and in fact the host may have multiple user laboratories. Many of the Computer Services requirements may apply to host, user, or both, depending on how information services are organized in the laboratory. The laboratory is responsible for ensuring that the provider of host functions meets CAP requirements (see GEN.42195, below).
The requirements in this section do NOT apply to the following:

1. Desktop calculators
2. Small programmable technical computers
3. Purchased services such as the Quality Assurance Service or Laboratory Management Index Service of the College of American Pathologists
4. Micro computers used solely for word processing, spreadsheets, or similar single user functions
5. Dedicated microprocessors or workstations that are an integral part of an analytic instrument

Inspector Instructions:

- CAP accreditation certificate of remote site or records that the host site is in compliance with this section of the checklist

**GEN.42195 Remote LIS**

If components of the LIS are located at a facility other than the one under this CAP accreditation number, there is evidence that the remote facility complies with CAP requirements for host LIS functions.

**NOTE:** This requirement does not apply if all components of the LIS are under the laboratory's CAP number. This requirement may be addressed by a copy of the CAP accreditation certificate from other sites, or evidence that the computer facility has been provided a copy of this Checklist, and has satisfactorily addressed the contents of the Computer Facility section, and all other pertinent requirements, with records provided to the laboratory director and the CAP inspector.

**COMPUTER FACILITY**

This section applies to laboratories where the computer facilities are housed.

Inspector Instructions:

- Computer facility equipment and location (clean, ventilated, protected against power surges)
- Fire extinguishers/equipment

**GEN.42750 Computer Facility Maintenance**

The computer facility and equipment are clean, well-maintained and adequately ventilated with appropriate environmental control.

**NOTE:** The computer facilities should be clean, well maintained and in a location that is environmentally controlled, as required by the most restrictive vendor specifications.

**GEN.42800 LIS Fire Equipment**

Fire-fighting equipment (extinguishers) is appropriate for electrical components available.
NOTE: Acceptable fire-fighting equipment/extinguishers in areas with information technology equipment may include:

1. Automatic sprinkler systems that are valved separately from other systems
2. Gaseous clean agent extinguishers systems
3. Listed portable fire extinguishers of carbon dioxide or halogenated agent type
4. Listed extinguishers with a minimum rating of 2-A for ordinary combustible material (paper and/or plastics)
5. Gaseous agent inside units or total flooding systems when there is critical need, e.g. to protect data in process, reduce equipment damage and to facilitate a return to service

Dry chemical extinguishers are not recommended because of the corrosive damage they cause. In the instance where no other extinguisher is available and there is imminent danger to personnel or property however, a dry extinguisher may be used.

REFERENCES

GEN.42900 LIS Power Phase II

The computer system is adequately protected against electrical power interruptions and surges.

NOTE: Protection from electrical surges and interruptions must be adequate to prevent loss of data. An uninterruptible power system (UPS) or similar protective device (e.g. isolation transformer) must be considered. Periodic testing of this protective equipment to ensure protection of data and proper shutdown of computer equipment is considered best practice.

REFERENCES

HARDWARE AND SOFTWARE

Inspector Instructions:

- Sampling of computer training records
- How does your laboratory verify the LIS following a hardware or software failure?
- Who do you notify when there is a computer malfunction?

GEN.43022 LIS Testing Phase II

There are records that programs are adequately tested for proper functioning when first installed and after any modifications, and that the laboratory director or designee has approved the use of all new programs and modifications.
NOTE: Computer programs must be checked for proper performance when first installed and after any changes or modifications. Any changes or modifications to the system must be recorded, and the laboratory director or designee must approve all changes, additions and deletions in programs, the test library, and major computer functions before they are released. Records must be retained for at least two years beyond the service life of the system.

REFERENCES

**REVISED** 07/28/2015
GEN.43033 Custom LIS Phase I

Customized software, and modifications to that software, are appropriately documented and records allow for tracking to identify persons that have added or modified that software.

NOTE: The purpose of the computer program, the way it functions, and its interaction with other programs must be clearly stated. The level of detail should be adequate to support troubleshooting, system modifications, or additional programming.

REFERENCES

**NEW** 07/28/2015
GEN.43040 LIS Policy and Procedure Approval Phase II

The laboratory director or designee reviews and approves all new LIS policies and procedures, as well as substantial changes to existing documents before implementation.

NOTE: Procedures should be appropriate to the level of use of the system, and must encompass the day-to-day activities of the laboratory staff as well as the daily operations of the Information Technology staff.

**REVISED** 07/28/2015
GEN.43055 Computer System Training Phase II

There are records for training of all users of the computer system initially, after system modification, and after installation of a new system.

NOTE: Review of LIS policies and procedures relevant to the scope of duties must be incorporated into the training.

GEN.43066 Computer Malfunction Notification Phase II

There is a written procedure with instructions for contacting a responsible person (e.g. Computer System Manager) in case of computer malfunction.

REFERENCES
SYSTEM SECURITY

The following requirements concern unauthorized users. If a system is vulnerable, steps should be taken to prevent unauthorized access.

Inspector Instructions:

- Sampling of computer security policies and procedures

GEN.43150 Access Patient Data Phase II

There are explicit written policies that specify who may use the computer system to enter or access patient data, change results, change billing or alter programs.

NOTE: Policies must define those who may only access patient data and users who are authorized to enter patient results, change results, change billing, or alter computer tables or programs. If data in other computer systems can be accessed through the LIS (e.g. pharmacy or medical records), policies must prevent unauthorized access to the data through the LIS.

REFERENCES

GEN.43200 Computer Access Codes Phase II

Computer access codes (security codes, user codes) are in place to confine individuals’ access to those functions they are authorized to use, and the security of access codes is maintained (e.g. inactivated when employees leave, not posted on terminals).

NOTE: The laboratory should establish security (user) codes to permit only specifically authorized individuals to access patient data or alter programs. A system that allows different levels of user access to the system based on the user’s authorization is desirable and usually provides effective security. Examples of best practices include: periodic alteration of passwords by users; minimum character length for passwords; password complexity requirements (e.g. a combination of alphanumerics); recording of failed log-on attempts with user lock-out after a defined number of unsuccessful log-on attempts.

REFERENCES

GEN.43262 Unauthorized Software Installation Phase I

There are written policies and procedures that govern installation of software on any computer used by the laboratory.

NOTE: Laboratory computers often serve multiple functions. Many of these computers are connected in a network. The security of the system should be sufficient to prevent the casual
user from installing software. Such unauthorized installation may cause instability of the operating system or introduce other unwanted consequences. Many operating systems allow procedures to restrict certain users from installing software.

REFERENCES

**REVISED** 07/28/2015

PUBLIC NETWORK SECURITY

If the facility uses a public network, such as the Internet as a data exchange medium, there are network security measures in place to ensure confidentiality of patient data.

NOTE: Information sent over a public domain such as the Internet or stored in "the cloud," is considered in the public domain. Thus it is potentially accessible to all parties on that network. Systems must be in place to protect network traffic, such as "fire walls" and data encryption schemes.

Evidence of Compliance:
✓ Written policy defining mechanism for data protection

REFERENCES

PATIENT DATA

Inspector Instructions:

- Records of the review of patient results containing calculated data
- How are absurd values detected?
- How does the technologist electronically enter comments regarding specimen quality?
- How does your laboratory verify manual and automated result entry?

**REVISED** 07/28/2015

CALCULATED PATIENT DATA VERIFICATION

Calculated values reported with patient results are reviewed every two years or when a system change is made that may affect the calculations.

NOTE: This checklist requirement applies only to calculations based on formulas modifiable by the user.

Errors can be inadvertently introduced into established computer programs. Calculations involving reportable patient results must be rechecked to ensure accuracy and records retained. This requirement applies to laboratory information systems, middleware, and analyzers. More frequent checks may be required for certain specific calculations, as delineated elsewhere in the checklists (e.g. INR).
When calculations are performed by an LIS shared by multiple laboratories, this review only needs to be done at one location and each individual laboratory must have access to a copy of the review records. However, any calculations specific to an individual laboratory's methodology must be reviewed by that laboratory and the record of that review must be available.

Evidence of Compliance:
✓ Records of validation of calculated test results

REFERENCES

GEN.43750 Specimen Quality Comment Phase II

The system provides for comments on specimen quality that might compromise the accuracy of analytic results (e.g. hemolyzed, lipemic).

Evidence of Compliance:
✓ Patient reports

REFERENCES
1) Jones JB. The importance of integrating POCT data into an organized database. Advance/Laboratory. 1999;8(9):8-10

GEN.43800 Data Input ID Phase II

There is an adequate system to identify all individuals who have entered and/or modified patient data or control files.

NOTE: When individual tests from a single test order (e.g. multiple tests with same accession number) are performed by separate individuals and the test result is entered into the LIS, the system must provide an audit trail to record each person involved. For example, a single accession number having orders for electrolytes and a lipid panel may have testing done by two or more individuals. The laboratory should be able to identify the responsible personnel who performed each test and posted the data. This includes sequential corrections made to a single test result. If autoverification is used, then the audit trail should reflect that the result was verified automatically at a given time.

With point-of-care testing, if the individual performing the test is different than the individual entering test data into the LIS, both should be uniquely identified by the system and retrievable by audit trail.

REFERENCES
1) Jones JB. The importance of integrating POCT data into an organized database. Advance/Laboratory. 1999;8(9):8-10

GEN.43825 Result Verification Phase II

Manual and automated result entries are verified before final acceptance and reporting by the computer.

NOTE: Data entered into the computer system either manually or by automated methods must be reviewed by an authorized individual who verifies the accuracy of the input data before final acceptance and reporting by the computer. An example of best practices for this step is checking the result against the reportable range and critical results for the test. Depending on the local environment, this may or may not require a second person. Verification procedures must generate an audit trail.

This checklist requirement does not apply to autoverification procedures (see below).

REFERENCES

GEN.43837 Downtime Result Reporting Phase II
There are written procedures to ensure reporting of patient results in a prompt and useful fashion during partial or complete downtime and recovery of the system.

REFERENCES

AUTOVERIFICATION

Autoverification is the process by which patient results are generated from interfaced instruments and sent to the LIS, where they are compared against laboratory-defined acceptance parameters. If the results fall within these defined parameters, the results are automatically released to patient reporting formats without any additional laboratory staff intervention. Any data that fall outside the defined parameters are reviewed by laboratory staff prior to reporting.

Inspector Instructions:
- Autoverification policies and procedures
- Autoverification validation records

**REVISED** 07/28/2015
GEN.43875 Autoverification Validation Phase II

There is documentation that the autoverification process was validated initially, and is tested at least annually and whenever there is a change to the system that could affect the autoverification logic.

NOTE: The range of results for which autoverification is acceptable must be defined for all patient tests subject to autoverification.

Validation of autoverification must include a process to confirm that the autoverification algorithm decision rules are functioning properly, including the use of previously assayed specimens with results that challenge the algorithm. Examples of specimens that may be needed to validate the autoverification algorithm decision rules may include specimens with analyte concentrations within the normal reference limit, above or below the reference limits, above or below the analytic measurement range, and in the critical range. Specimens with known interferences and specimens that require calculations should also be used, when applicable.

When changes are made that might affect the autoverification decision algorithm, validation appropriate to the scope and nature of the change must be performed.

Evidence of Compliance:
✓ Records of autoverification validation studies, including laboratory director approval AND
✓ Records of ongoing retesting of the autoverification process at least annually and at changes to the system

REFERENCES
For all test results subject to autoverification, the laboratory ensures that applicable quality control samples have been run within an appropriate time period, with acceptable results.

NOTE: This requirement may be met by, 1) the computer system automatically checking quality control status prior to autoverification, or, 2) manually disabling autoverification after any unacceptable QC result, or when QC has not been run within the required time interval.

Evidence of Compliance:
✓ Procedure defining the QC process AND
✓ QC data to show that QC was performed at defined intervals

**REVISED** 07/28/2015

GEN.43881 Autoverification Results Phase II

Results are compared with an appropriate range of acceptable values and flags or warnings reviewed prior to autoverification.

NOTE: Appropriate comparisons include checking patient results against absurd and critical results requiring manual intervention (repeat testing, dilution, telephone notification of results, etc.)

The mere presence of a flag may not disqualify a result from autoverification, but any flag that is not specifically recognized by the autoverification program must cause the flagged result to be held for manual review.

Evidence of Compliance:
✓ Records of system rules including comparison of patient results against absurd and critical values

GEN.43887 Autoverification Audit Trail Phase I

The audit trail in the computer system identifies all test results that were autoverified, and the date/time of autoverification.

GEN.43890 Autoverification Delta Checks Phase I

The autoverification process includes all delta checks that the laboratory performs prior to manual release of test results.

NOTE: This requirement does not require delta-checking for all autoverified results, but the laboratory’s delta-checking procedures should be the same for manually released and autoverified test results.

Evidence of Compliance:
✓ Records of system rules including the use of delta checks when appropriate

REFERENCES

GEN.43893 Autoverification Suspension Phase II

The laboratory has a procedure for rapid suspension of autoverification.

NOTE: Laboratory personnel should be able to suspend autoverification in the event of a problem with a test method, analytic instrument or the autoverification program.
DATA RETRIEVAL AND PRESERVATION

Inspector Instructions:

- Data preservation policies and procedures
- If there are indications that the computer system is inadequate to meet the patient needs of the organization, further evaluate laboratory/LIS leadership’s responses, corrective actions, and resolutions

GEN.43900 Archived Test Result

A complete copy of archived patient test results can be retrieved, including original reference ranges and interpretive comments, including any flags or footnotes that were present in the original report, and the date of the original report.

NOTE: Stored patient result data and archival information must be easily and readily retrievable within a time frame consistent with patient care needs.

REFERENCES

GEN.43920 Multiple Analyzer ID

When multiple identical analyzers are used, they are uniquely identified such that a test result may be appropriately traced back to the instrument performing the test.

NOTE: Best practice is to store these data in the LIS.

GEN.43946 Data Preservation/Destructive Event

There are written procedures for the preservation of data and equipment in case of an unexpected destructive event (e.g., fire, flood), software failure and/or hardware failure, and these procedures allow for the timely restoration of service, including data integrity check.

NOTE: Procedures must 1) be adequate to address scheduled and unscheduled interruptions of power or function; 2) be tested periodically for effectiveness; and 3) include systems to backup programs and data.

These procedures can include, but are not limited to, 1) steps to limit the extent of the destructive event, 2) periodic backing up and storing of information, 3) off-site storage of backup data, and 4) restoring information from backed up media. The procedures should specifically address the recoverability of patient information. Changes to hardware and software commonly require review and reevaluation of these written procedures. These procedures must specifically address the physical environment and equipment and are often addressed by the organization’s disaster plan.
REFERENCES

INTERFACES

Inspector Instructions:

READ

• Interface systems policies and procedures
• Sampling of reports transmitted to each interfaced system (laboratory data entry of results match patient reports, including reference ranges and comments)

ASK

• How does your laboratory verify the accuracy of data transmission from the LIS to interfaced systems?

GEN.46000 Reference Range/Units Transmission

As applicable, reference ranges and units of measure for every test are transmitted with the patient result across the interface.

NOTE: The reference range, including units of measure, may be specific for a given patient result and should be attached to that result such that it will be displayed along with the patient result.

GEN.48500 Interface Result Integrity

There is a procedure to verify that patient results are accurately transmitted from the point of data entry (interfaced instruments and manual input) to patient reports (whether paper or electronic).

NOTE: Verification must be performed prior to implementation of an interface (i.e. pre go-live), and every 2 years thereafter. This includes evaluation of data transmitted from the LIS to other computer systems and their output devices. Reference ranges and comments, as well as actual patient results and report formats, must be evaluated.

Verification of accurate data transmission from the LIS to other systems must be performed by reviewing data in the first downstream (or interfaced) system in which the ordering clinician/client (e.g. referring laboratory) may be expected to routinely access patient data. This requirement can be met by printing screen shots or by other methods that record that a verification procedure has been performed. If the LIS has separate interfaces to multiple receiving systems in which patient data can be accessed by clinicians, then reports from each receiving system must be validated. However, where multiple sites use the same recipient system (e.g. the same installed instance of an electronic medical record system), validation need only occur for the interface (i.e. at one of the sites) and not for each individual site that is served by that single installed system.

At implementation of a new interface, or change to an existing interface, validation of at least 2 examples of reports from each of the following disciplines, where applicable, satisfies the intent of this checklist requirement. Subsequently, at least 2 examples of reports from at least
4 of these disciplines should be validated every 2 years. Not all of these report types will be applicable to every laboratory:

1. Surgical pathology reports
2. Cytopathology reports (preferably gynecologic and non-gynecologic)
3. Clinical laboratory textual reports (e.g. molecular, protein electrophoresis, coagulation panel interpretation)
4. Quantitative results (e.g. chemistry, hematology, or coagulation)
5. Qualitative or categorical results (e.g. serology)
6. Microbiology reports (e.g. culture and antimicrobial sensitivity)
7. Blood bank reports (e.g. type and screen)

Interface validation should include examples of individual results, test packages or batteries, abnormal flags, and results with comments/footnotes. Initial interface validation should include verification that corrected results for clinical laboratory and anatomic pathology results are handled accurately in the receiving system.

Evidence of Compliance:
✓ Records of verification

REFERENCES

GEN.48750 LIS Interface Shutdown/Recovery Phase II
There are procedures for changes in laboratory functions necessary during partial or complete shutdown and recovery of systems that interface with the laboratory information system.

NOTE: These procedures must ensure integrity of patient test data. Procedures must include verifying recovery of interfaced systems, and replacement or updating of data files, as necessary.

REFERENCES

TELEPATHOLOGY AND REMOTE DATA ASSESSMENT
This section applies to telepathology, including the practice of pathology and cytology, in which a pathologist examines digitized or analog video, still image(s), or other data files (e.g. flow cytometry files, Sanger sequencing data) at an off-site or remote location and an interpretation is rendered that is included in a formal diagnostic report or in the patient record. It also includes the review of images by a cytopathologist when a judgment of adequacy is recorded in the patient record. This section may be applied to, but is not limited to the disciplines of anatomic pathology, cytopathology, hematopathology, cytogenetics, flow cytometry, histocompatibility, and molecular pathology.

Telepathology modes include:
- Static telepathology – interpretation based on pre-selected still image(s)
- Dynamic telepathology - viewing real-time images (includes robotic microscopy, video streaming, and desktop sharing)
- Virtual slides/whole slide imaging
This checklist section applies to:

- Primary diagnoses made by telepathology
- Frozen section diagnoses
- Formal second-opinion consultations
- Ancillary techniques in which the pathologist participates in interpretation of images
- Real-time evaluation of FNA specimens for triaging and preliminary diagnosis

This checklist section is NOT applicable to:

- Informal reviews without formal reporting
- Educational or research use of these systems

References:

Inspector Instructions:

- Sampling of telepathology policies and procedures
- Sampling of reports generated from reviews of images/slides and data files performed by telepathology

**REVISED** 08/17/2016
GEN.50057  Slide/Image ID  Phase II

There is a method for the individual reviewing cases to ensure that correct patient identification and slides/images and data files are submitted for review.

NOTE: There are multiple ways to accomplish positive patient identification, including verbal communications, images of slide identifier, etc.

Evidence of Compliance:

✓ Written procedure defining mechanism to positively identify slides/images and data files

**REVISED** 08/17/2016
GEN.50614  Clinical Information Access  Phase I

The individual reviewing cases has access to pertinent clinical information at the time of slide/image(s) or data file review.

NOTE: Typically this information includes at least the information on the surgical pathology or cytology requisition form.

**NEW** 08/17/2016
GEN.50630  Telepathology System Validation  Phase I

The laboratory validates telepathology systems used for clinical diagnostic purposes by performing its own validation studies, including approval for use by the laboratory director (or designee who meets CAP director qualifications) before the technology is used for the intended diagnostic purpose(s).

NOTE: The specific components of the validation study are left to the discretion of the laboratory. As general guiding principles, the validation process should:

- Closely emulate the real-world clinical environment and involve specimen preparation types and clinical settings relevant to the intended use(s).
Laboratory General Checklist

- Be carried out by a pathologist(s) adequately trained to use the system.

Refer to GEN.52920 for requirements on validation of whole slide imaging.

Evidence of Compliance:
✓ Records of completed validation study with review and approval

**REVISED** 08/17/2016
**REVISED** 08/17/2016
**REVISED** 08/17/2016
**REVISED** 07/28/2015

**TELEPATHOLOGY TRAINING**

**GEN.51728** Telepathology Training

The lab has a procedure addressing training requirements for all users of the telepathology system.

NOTE: The training procedure should be role-specific, as defined in the approved laboratory procedures. Retraining may be required when significant system changes are made.

Evidence of Compliance:
✓ Records for telepathology training in personnel files

**GEN.52842** Telepathology and Patient Confidentiality

There are procedures in place to ensure that sites engaging in telepathology provide reasonable confidentiality and security.

NOTE: Procedures might include message security, system and user authentication, activity logs, encryption, and access restrictions. These security considerations must be particularly adhered to when using mobile devices in public places.

For laboratories subject to US regulations, the procedures must be in conformance with HIPAA requirements.

**GEN.52850** Telepathology Result Records

The telepathology records include diagnoses made and statements of adequacy assessment, preliminary diagnosis, or recommendations for additional studies provided at the time of evaluation.

NOTE: Such records are not required to be included on the patient report.

Evidence of Compliance:
✓ Reports generated from reviews of images/slides and data files performed by telepathology

**GEN.52860** Quality Management Program

Telepathology services are included in the laboratory’s quality management program.

NOTE: For example, the laboratory might monitor the frequency of deferral cases, comparison to on-site evaluation, or consultation using traditional glass slide microscopy.

**WHOLE SLIDE IMAGING**

This section applies to laboratories using whole slide imaging systems for diagnostic purposes (primary and/or consultation).
Inspector Instructions:

- Sampling of training records
- System validation records

**REVISED** 08/17/2016

GEN.52900 Whole Slide Imaging User Training Phase I

There are records showing that all users of the whole slide imaging system have been trained.

NOTE: Users of the whole slide imaging system include individuals responsible for slide scanning and digital slide quality assessment, as well as pathologists. The training procedure should include role-specific training, as defined in the approved laboratory procedures. Retraining may be required when significant system changes are made.

Evidence of Compliance:
✓ Records for whole slide image training in personnel files

GEN.52920 Whole Slide Imaging System Validation Phase I

The laboratory validates whole slide imaging systems used for clinical diagnostic purposes by performing its own validation studies, including approval for use by the laboratory director (or designee who meets CAP director qualifications) before the technology is used for the intended diagnostic purpose(s).

NOTE: The specific components of the validation study are left to the discretion of the laboratory.

As general guiding principles, the validation process should:
- Closely emulate the real-world clinical environment and involve specimen preparation types and clinical settings relevant to the intended use(s);
- Be carried out by a pathologist(s) adequately trained to use the system;
- Assess intraobserver concordance between digital and glass slides;
- Encompass the entire whole slide imaging system, with reevaluation if a significant change is made to a previously validated system.

Evidence of Compliance:
✓ Records of completed validation study with review and approval

REFERENCES
1) Pantanowitz et al, Validating whole slide imaging for diagnostic purposes in pathology: Guideline from the College of American Pathologists Pathology and Laboratory Quality Center. ARPA, 2013


PERSONNEL

The laboratory must have personnel policies, and job descriptions that define qualifications and duties for all positions. Personnel files must contain records of educational qualifications (e.g. copies of diplomas, transcripts, primary source verification reports), laboratory personnel licenses (where required), training and continuing education for each employee. Ideally, these files should be located in the laboratory. If they are retained outside
of the laboratory, they must be readily available to the inspector on the day of inspection. The inspector reviews the personnel files using the Laboratory Personnel Evaluation Roster.

**Inspector Instructions:**

- Sampling of personnel policies and procedures
- Organizational chart or narrative description
- Sampling of competency assessments for assessment of all six elements of competency for each nonwaived test system, six month competency for new employees, and assessments done by qualified individuals based on complexity of testing performed
- Sampling of technical personnel files for educational qualifications, (e.g. diplomas, transcripts, primary source verification reports), laboratory personnel licenses (where required), training, and continuing education records for testing personnel, section directors/technical supervisors, supervisors/general supervisors, and consultants using the table below
- Records of diploma/transcript equivalency evaluation for non-US trained personnel by a foreign credentialing agency (for laboratories subject to US regulations)

Sampling to include a mix of the following: 1) laboratory personnel (including MD, DO, PhD, technicians and technologists) and non-laboratory personnel (e.g. POC, PPT, Radiology, Respiratory); 2) full and part-time employees on all shifts and throughout all departments; 3) supervisory staff and testing personnel.

All newly hired personnel for the last two years must be reviewed (if applicable), both laboratory and non-laboratory.

If any documents are missing from the personnel files, record the appropriate deficiency on the Inspector’s Summation Report, noting the specific instances of non-compliance.

DO NOT allow laboratory staff to select which personnel records to review. Randomly select specific individuals from the Laboratory Personnel Evaluation Roster following sampling instructions described above. Use the following criteria to determine the number of personnel records to be reviewed:

<table>
<thead>
<tr>
<th>Total Number of Personnel (both laboratory and non-laboratory)</th>
<th>Number of Personnel Records to Sample</th>
</tr>
</thead>
<tbody>
<tr>
<td>Up to 10</td>
<td>Review all personnel</td>
</tr>
<tr>
<td>11 - 100</td>
<td>8 - 10</td>
</tr>
<tr>
<td>101 - 250</td>
<td>10 - 12</td>
</tr>
<tr>
<td>251 - 400</td>
<td>13 - 15</td>
</tr>
<tr>
<td>401 - 500</td>
<td>16 - 18</td>
</tr>
<tr>
<td>More than 500</td>
<td>18 - 20</td>
</tr>
</tbody>
</table>

**Records of all technical personnel hired within the last two years must be reviewed**

- Do you have a specific example of an employee who demonstrated unacceptable competency assessments? What were the corrective actions?
- What continuing education classes are available to employees?
- If primary source verification reports are used, what is your process to ensure that the reports contain the required elements?
**SECTION DIRECTORS (TECHNICAL SUPERVISORS)/GENERAL SUPERVISORS**

This section applies to laboratories performing one or more highly complex tests. The individuals fulfilling these roles must be identified on the CAP's Laboratory Personnel Evaluation Roster form.

The term "section director" may be considered synonymous to the technical supervisor in the checklist requirements. The term "supervisor" may be considered synonymous to the general supervisor in the checklist requirements. Within the laboratory’s organizational structure, the actual position titles may be different. A qualified laboratory director may serve as the section director and general supervisor, and may set position requirements more stringent than defined in the checklist.

**REVISED** 08/17/2016

GEN.53400  Section Director (Technical Supervisor) Qualifications/Responsibilities  Phase II

**Section Directors/Technical Supervisors of high complexity testing meet defined qualifications for the specialties supervised and fulfill the expected responsibilities.**

**NOTE:** For high complexity testing, one or more individuals qualified as a technical supervisor must be identified on the CAP’s Laboratory Personnel Evaluation Roster form.

Requirements for the section directors of clinical cytogenetics, histocompatibility and transfusion medicine services are more stringent and are found in the Cytogenetics, Histocompatibility and Transfusion Medicine Checklists, respectively.

The technical supervisor must meet the following requirements:

1. **MD or DO licensed to practice (if required) in the jurisdiction where the laboratory is located with certification in anatomic pathology or clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology or possess qualifications equivalent to those required for certification.**
   - If responsible for anatomic pathology or cytopathology must be board certified in anatomic pathology or possess equivalent qualifications
   - If responsible for clinical pathology must be board certified in clinical pathology or possess equivalent qualifications
   - If responsible for anatomic pathology and/or cytopathology, and clinical pathology, must be board certified in both anatomic and clinical pathology or possess equivalent qualifications OR

2. For specialties other than Anatomic Pathology and Cytopathology, an individual will meet the qualifications of a technical supervisor providing the following qualifications are met:
   - MD or DO licensed to practice (if required) in the jurisdiction where the laboratory is located with at least one year of training and/or experience in high-complexity testing*; or
   - Doctoral degree in chemical, physical, biological or clinical laboratory science from an accredited institution with at least one year of laboratory training and/or experience in high complexity testing*; or
   - Master's degree in a chemical, physical, biological, or clinical laboratory science or medical technology from an accredited institution with at least two years of laboratory training and/or experience in high complexity testing*; or
Bachelor’s degree in a chemical, physical, or biological science or medical technology from an accredited institution with at least four years of laboratory training and/or experience in high complexity testing*.

*The technical supervisor's training and experience must be in the designated specialty or subspecialty area of service for which the individual is responsible.

For laboratories subject to US regulations, alternate qualifications for the following specialty areas can be found in Fed Register. 1992 (Feb 28): 7177-7180 [42CFR493.1449]: bacteriology, mycobacteriology, mycology, parasitology, virology, cytology, ophthalmic pathology, dermatopathology, oral pathology, and radiobioassay.

If more stringent state or local regulations are in place for supervisory qualifications, including requirements for state licensure, they must be followed.

For laboratories subject to US regulations, credentials for all personnel trained outside of the US must be reviewed and recorded to ensure that their training and qualifications are equivalent to CLIA requirements. The equivalency evaluations should be performed by a nationally recognized organization.

The section director, as designated by the laboratory director, must be accessible to the laboratory as needed for on-site, telephone, or electronic consultation and is responsible for the technical and scientific oversight of the laboratory. The section director is responsible for performing and recording competency assessment for high complexity testing. The duties for performing the competency assessment may be delegated, in writing, to individuals meeting general supervisor qualifications for high complexity testing. Other responsibilities of the technical supervisor include:

- Selection of test methodology
- Establishment or verification of laboratory test performance specifications
- Enrollment and participation in proficiency testing
- Establishment of a quality control program to monitor ongoing test performance
- Resolution of technical problems and ensuring that remedial actions are taken
- Ensuring that patient results are not reported until corrective actions are taken and test systems are functioning properly
- Identification of training needs

For functions that are delegated, such as review of quality control data, assessment of competency, or review of proficiency testing performance, delegation must be in writing and the technical supervisor is responsible to ensure that those functions are properly carried out by a qualified individual.

Evidence of Compliance:

- Records of qualifications including degree, transcript, equivalency evaluation, or current license (if required) AND
- Certification/registration (if required) and work history in related field AND
- Description of current duties and responsibilities AND
- Record of delegation of duties

REFERENCES

**REVISED** 08/17/2016

General Supervisor Qualifications/Responsibilities  Phase II

Supervisors/general supervisors meet defined qualifications and fulfill expected responsibilities.

NOTE: For high complexity testing, one or more individuals qualified as a general supervisor must be defined on the CAP’s Laboratory Personnel Evaluation Roster form.
Supervisors who do not qualify as a laboratory director or section director/technical supervisor must qualify as testing personnel and possess the minimum of a:

1. Bachelor's degree in a chemical, physical, biological or clinical laboratory science or medical technology with at least one year of training and/or experience in high complexity testing; or
2. Associate degree in a laboratory science or medical technology with at least two years of training and/or experience in high complexity testing; or
3. Have previously qualified or could have qualified as a general supervisor prior to 2/28/1992

Requirements for the supervisors/general supervisors of cytopathology and histocompatibility are more stringent and are found in the Cytopathology and Histocompatibility Checklists.

If more stringent state or local regulations are in place for supervisory qualifications, including requirements for state licensure, they must be followed.

For laboratories subject to US regulations, credentials for all personnel trained outside of the US must be reviewed and recorded to ensure that their training and qualifications are equivalent to CLIA requirements. The equivalency evaluations should be performed by a nationally recognized organization.

The supervisor of high-complexity testing must be accessible to the laboratory as needed for on-site, telephone, or electronic consultation and is responsible for day-to-day supervision or oversight of the laboratory operation and personnel performing testing and reporting test results. Individuals meeting the qualifications of a general supervisor for high complexity testing may assess the competency of high complexity testing personnel, if this duty is delegated, in writing, by the section director. Other responsibilities of the general supervisor include:

- Resolution of technical problems in accordance with policies and procedures established by the laboratory director or technical supervisor
- Monitoring of test performance
- Ensuring that remedial actions are taken when test systems deviate from the laboratory's established performance specifications
- Providing orientation of testing personnel

Evidence of Compliance:

- Records of qualifications including degree, transcript, equivalency evaluation, or current laboratory personnel license (if required) AND
- Certification/registration (if required) and work history in related field AND
- Description of current duties and responsibilities

REFERENCES

TECHNICAL AND CLINICAL CONSULTANT

The individuals fulfilling these roles must be identified on the CAP's Laboratory Personnel Evaluation Roster form.

Within the laboratory's organizational structure, the actual position titles may be different. A qualified laboratory director may also serve as the technical and clinical consultant, and may set position requirements more stringent than defined in the checklist.

**REVISED** 08/17/2016
GEN.53625  Technical Consultant Qualifications/Responsibilities   Phase II
Technical consultants meet defined qualifications and fulfill expected responsibilities.

NOTE: This requirement applies to laboratories performing moderate complexity testing, but not high complexity testing. For moderate complexity testing, one or more individuals qualified as a technical consultant must be identified on the CAP’s Laboratory Personnel Evaluation Roster form (for high complexity testing, refer to GEN.53400).

The technical consultant (including the laboratory director who serves as a technical consultant) must be qualified by education and experience by one of the following combinations:

- MD or DO, licensed to practice medicine in the jurisdiction where the laboratory is located (if required), with certification in anatomic and/or clinical pathology by the American Board of Pathology or the American Osteopathic Board of Pathology, or possess qualifications equivalent to those required for certification; or
- MD, DO, or DPM, licensed to practice in the jurisdiction where the laboratory is located (if required), with at least one year of training and/or experience in nonwaived testing*; or
- Doctoral or master's degree in a chemical, physical, biological or clinical laboratory science from an accredited institution with at least one year of training and/or experience in nonwaived testing*; or
- Bachelor's degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution with at least two years of training and/or experience in nonwaived testing*.

*The technical consultant’s training and experience must be in the designated specialty or subspecialty area of service for which the individual is responsible.

If more stringent state or local regulations are in place for supervisory qualifications, including requirements for state licensure, they must be followed.

The technical consultant is responsible for the technical and scientific oversight of the laboratory. The technical consultant must be available to the laboratory as needed for telephone, electronic and on-site consultation. Individuals meeting the qualifications of a technical consultant may assess the competency of personnel performing moderate complexity testing, if this duty is delegated, in writing, by the laboratory director. Other responsibilities of the technical consultant include:

- Establishment or verification of laboratory test performance specifications
- Selection of test methodology
- Enrollment and participation in proficiency testing
- Establishment of a quality control program to monitor ongoing test performance
- Resolution of technical problems and ensuring that remedial actions are taken
- Ensuring that patient results are not reported until corrective actions are taken and test systems are functioning properly
- Identification of training needs

Evidence of Compliance:

✓ Records of technical qualifications including degree, transcript, equivalency evaluation, or current license (if required) AND
✓ Certification/registration (if required) and work history in related field AND
✓ Description of current duties and responsibilities

REFERENCES

NOTE: This requirement applies to laboratories performing moderate complexity testing and/or high complexity testing. One or more individuals qualified as a clinical consultant must be identified on the CAP's Laboratory Personnel Evaluation Roster form.

Clinical consultants must be an MD, DO, DPM licensed to practice medicine in the jurisdiction where the laboratory is located (if required) or doctoral scientist certified by an HHS-approved board.

The clinical consultant must be available to provide and ensure that consultation is available on test ordering, and interpretation of results relating to specific patient conditions, and for matters relating to the quality of test results reported. The clinical consultant must also ensure that patient reports include pertinent information required for interpretation. See TLC.10440, TLC.10500, and TLC.10700.

Evidence of Compliance:
✓ Records of clinical consultant qualifications (i.e. a valid medical license AND
✓ Written job description or contract AND
✓ Records of activities performed by the consultant during visits consistent with the job description (e.g. meeting minutes, activity logs, signed summaries or data with evidence of review)

REFERENCES

ALL PERSONNEL

**REVISED** 08/17/2016
GEN.54000 Organizational Chart

There is an organizational chart for the laboratory, or a narrative description that describes the reporting relationships among the laboratory's owner or management, the laboratory director, section director(s)/technical supervisor(s), technical consultant(s), clinical consultant(s), and supervisor(s)/general supervisor(s), as appropriate.

**NEW** 08/17/2016
GEN.54025 Laboratory Personnel Evaluation Roster

The Laboratory Personnel Evaluation Roster is current and accurate and is audited by the laboratory director or designee at least annually for nonwaived testing personnel and personnel fulfilling supervisor roles.

NOTE: The laboratory’s audit of the laboratory personnel evaluation roster must include a review of a mixture of the following types of personnel:
• All nonwaived testing personnel hired within the last 12 months (laboratory and non-laboratory)
• Laboratory and non-laboratory (POC, PPT, Radiology, Respiratory, etc.) personnel
• Full and part-time nonwaived testing personnel on all shifts and throughout all departments
• Personnel fulfilling supervisory roles (e.g. laboratory director, technical supervisor, staff pathologist)

Personnel performing any CLIA-defined duty must be listed on the roster. Personnel performing waived testing only or whose duties are limited to phlebotomy, clerical work or specimen processing are not required to be listed on the Laboratory Personnel Evaluation Roster. Histology personnel that do not perform high-complexity testing are also excluded. All grossing performed in histology is considered high-complexity testing.
Evidence of Compliance:
✓ Records of completed rosters accurately reflecting personnel AND
✓ Records of annual audits performed by the laboratory director or designee

GEN.54200  Continuing Education  Phase I

There is a functional continuing laboratory education program adequate to meet the needs of all personnel.

Evidence of Compliance:
✓ Written policy for continuing laboratory education

REFERENCES

**REVISED** 08/17/2016

GEN.54400  Personnel Records  Phase II

Personnel files are maintained on all current technical personnel and personnel records include all of the following, as applicable:

1. For nonwaived testing personnel, copy of academic diploma, transcript, or primary source verification reports confirming credentials (Refer to NOTE 2 for use of primary source verification reports)
2. Laboratory personnel license, if required by state, province, or country
3. Summary of training and experience
4. Certification, if required by state or employer
5. Description of current duties and responsibilities as specified by the laboratory director: a) Procedures the individual is authorized to perform, b) Whether supervision is required for specimen processing, test performance or result reporting, c) Whether supervisory or section director review is required to report patient test results
6. Records of continuing education
7. Records of radiation exposure where applicable (such as with in vivo radiation testing), but not required for low exposure levels such as certain in-vitro testing
8. Work-related incident and/or accident records
9. Dates of employment

NOTE 1: All records, in either electronic or paper form, must be readily available for review by the inspector at the time of the CAP inspection.

NOTE 2: For laboratories subject to US regulations:
- The file must include records demonstrating that each individual meets the required educational qualifications for the position(s) held, such as a copy of the academic diploma, transcript, or primary source verification (PSV) report.
- If PSV reports are used, the laboratory must have a defined system for reviewing the reports, with written criteria for acceptance. PSV is typically performed by a third-party agent or company that directly contacts institutions and former employers to verify training and experience, such as diplomas, board certification, licensure, and reported work history. PSV reports confirming the required qualifications may be retained in lieu of obtaining paper copies of these records. If there are required elements for the qualifications that the PSV report does not adequately verify (e.g. transcripts, educational equivalency for personnel trained outside of the US, or reports lacking the type of degree earned), there must be records showing that qualifications are met using other means.
Laboratory General Checklist

The credentialing systems used by the Department of Veterans Affairs (i.e. VetPro Credentialing System) and Department of Defense may be used to document educational qualifications. Records must be available upon request.

If the laboratory is located in a state that requires laboratory personnel licensure, the license may be used instead of the diploma or transcript to show that educational qualifications were met. Licensure records for any other discipline, such as nursing, respiratory therapy, or radiology is not required, and cannot be used to meet educational qualifications for non-waived laboratory testing. These individuals must have all required educational and training records in their files.

While certification of technical personnel by a professional organization, such as ASCP or AMT, is highly desirable, records of the certification alone are not considered adequate to demonstrate that educational qualifications have been met.

The training and qualifications of all personnel trained outside of the US must be evaluated to determine equivalency of their education to an education obtained in the United States, with records of the evaluation available in the personnel file. The equivalency evaluations should be performed by a nationally recognized organization. For Department of Defense laboratories, the equivalency must be performed using a process approved by the Center for Laboratory Medicine Services.

NOTE 3: Laboratories not subject to US regulations may authenticate educational achievement according to prevailing governmental rules.

Evidence of Compliance:
✓ Copies of diplomas, transcripts, equivalency evaluation, or current laboratory personnel licensure (if required) accessible at the laboratory OR
✓ Policy for use of primary source verification reports, with criteria for acceptance, if used AND
✓ Primary source verification reports with required elements

REFERENCES

**REVISED** 08/17/2016

**GEN.54750** Testing Personnel Qualifications Phase II

All testing personnel meet the following requirements.

1. Personnel performing high complexity testing must have a minimum of one of the following:
   - Bachelor’s degree in a chemical, physical, biological or clinical laboratory science or medical technology from an accredited institution; or
   - Associate degree in a laboratory science (chemical or biological science) or medical laboratory technology from an accredited institution, or equivalent laboratory training and experience meeting the requirements defined in the CLIA regulation 42CFR493.1489 (see NOTE 2).

2. Personnel performing moderate complexity testing, including non-laboratory personnel, must have a minimum of one of the following:
   - Associate degree in a chemical, physical, or biological science or medical laboratory technology from an accredited institution; or
   - High school graduate or equivalent and have successfully completed an official military medical laboratory procedures course and have held the military enlisted occupational specialty of Medical Laboratory Specialist; or
   - High school diploma or equivalent and have a record of training defined in the CLIA regulation 42CFR493.1423 (see NOTE 4)
NOTE 1: Laboratory and non-laboratory (e.g. nurses, respiratory therapists, radiologic technologists, and medical assistants) testing personnel must meet the qualifications appropriate to the complexity of testing performed. GEN.54400 contains the specific requirements for the types of records that must be maintained in the personnel file to demonstrate compliance. Additional information for assessing personnel qualifications is available at the following link: CAP Personnel Requirements by Testing Complexity.

NOTE 2: For high complexity testing, equivalent laboratory training and experience includes the following:

- 60 semester hours or equivalent from an accredited institution that, at a minimum, includes either 24 semester hours of medical laboratory technology courses, OR 24 semester hours of science courses that include six semester hours of chemistry, six semester hours of biology, and 12 semester hours of chemistry, biology or medical laboratory technology in any combination; AND
- Laboratory training including either completion of a clinical laboratory training program approved or accredited by the ABHES, NAACLS, or other organization approved by HHS (note that this training may be included in the 60 semester hours listed above), OR at least three months documented laboratory training in each specialty in which the individual performs high complexity testing.

NOTE 3: For US Department of Defense laboratories, effective May 29, 2014, newly hired high complexity testing personnel must have either:

- A minimum of an associate degree in a biological or chemical science or medical laboratory technology from an accredited institution AND be certified by the ASCP, AMT or other organization deemed comparable by OASD(HA) or their designee (CCLM) as an MLT or MT/MLS; OR
- Successfully completed an official U.S. military medical laboratory procedures training course of at least 50 weeks duration and currently hold the military enlisted occupational specialty of medical laboratory specialist (laboratory technician).

NOTE 4: For moderate complexity testing personnel qualifying with a high school diploma or equivalent qualifications only, training records must demonstrate skills for the following:

- Specimen collection, including patient preparation, labeling, handling, preservation, processing, transportation, and storage of specimens, as applicable;
- Implementation of all laboratory procedures;
- Performance of each test method and for proper instrument use;
- Preventive maintenance, troubleshooting and calibration procedures for each test performed;
- Working knowledge of reagent stability and storage;
- Implementation of quality control policies and procedures;
- An awareness of interferences and other factors that influence test results; and
- Assessment and verification of the validity of patient test results, including the performance of quality control prior to reporting patient results.

NOTE 5: Students gaining experience in the field must work under the direct supervision of a qualified individual.

NOTE 6: If more stringent state or local regulations are in place for personnel qualifications, including requirements for state licensure, they must be followed.

Evidence of Compliance:

✓ Records of qualifications including diploma, transcript, equivalency evaluation, or current laboratory personnel license (if required) AND
✓ Work history in related field

REFERENCES


**GEN.55400**  Visual Color Discrimination  

**Phase I**

**Personnel are tested for visual color discrimination.**

*NOTE:* Personnel performing testing or other tasks that require color discrimination should be evaluated for difficulty with visual color discrimination. Evaluation is not required for personnel who do not perform such functions. Evaluation limited to discrimination of those colored items pertinent to the job is sufficient.

**Evidence of Compliance:**

✓ Record of color discrimination testing OR functional assessment, if indicated

**REVISED** 08/17/2016

**GEN.55450**  Initial Training  

**Phase II**

**There are records that all laboratory personnel have satisfactorily completed initial training on all instruments/methods applicable to their designated job.**

*NOTE:* The records must cover all testing performed by each individual. Training records must be maintained for a minimum of two years (five years for transfusion medicine). After the initial two-year (or five-year) period, records of successful ongoing competency assessment may be used to demonstrate compliance with this requirement.

Retraining must occur when problems are identified with personnel performance.

**REFERENCES**


**REVISED** 08/17/2016

**GEN.55500**  Competency Assessment of Testing Personnel  

**Phase II**

The competency of each person performing patient testing to perform his/her assigned duties is assessed.

*NOTE:* Prior to starting patient testing and prior to reporting patient results for new methods or instruments, each individual must have training and be evaluated for proper test performance as required in GEN.55450. Retraining and reassessment of employee competency must occur when problems are identified with employee performance.

For waived testing:

- Competency assessment must be performed at least annually (semiannual assessment not required)
- The laboratory may select which elements to assess for each test system
- The qualifications of individuals assessing competency of waived testing personnel shall be determined by the laboratory director
- If more stringent state or local regulations are in place for competency assessment of waived testing, they must be followed

For nonwaived testing:

- During the first year of an individual's duties, competency must be assessed at least semiannually
- After an individual has performed his/her duties for one year, competency must be assessed at least annually
- Competency assessment must include all six elements described below for each individual on each test system during each assessment period, unless an element is not applicable to the test system

Elements of competency assessment include but are not limited to:
1. Direct observations of routine patient test performance, including, as applicable, patient identification and preparation; and specimen collection, handling, processing and testing
2. Monitoring the recording and reporting of test results, including, as applicable, reporting critical results
3. Review of intermediate test results or worksheets, quality control records, proficiency testing results, and preventive maintenance records
4. Direct observation of performance of instrument maintenance and function checks
5. Assessment of test performance through testing previously analyzed specimens, internal blind testing samples or external proficiency testing samples; and
6. Evaluation of problem-solving skills

The laboratory must identify the test systems that an employee uses to generate patient test results. Competency must be evaluated and recorded for all testing personnel for each test system. A TEST SYSTEM is the process that includes pre-analytic, analytic, and post-analytic steps used to produce a test result or set of results. A test system may be manual, automated, multi-channel or single use and can include reagents, components, equipment or instruments required to produce results. A test system may encompass multiple identical analyzers or devices. Different test systems may be used for the same analyte. In many situations, tests performed on the same analyzer may be considered one test system; however, if there are any tests with unique aspects, problems or procedures within the same testing platform (e.g. pretreatment of samples prior to analysis), competency must be assessed as a separate test system to ensure staff are performing those aspects correctly.

Many of the elements of competency assessment are performed during routine review of an employee throughout the year. Records of these elements, including adherence to laboratory policies and procedures, observation of test performance, results reporting, instrument maintenance, review of worksheets, recording QC, performance of PT, and demonstration of taking appropriate corrective actions are examples of daily activities that can be used to demonstrate competency. If elements of competency are assessed by routine review, the competency procedure must outline how this routine review is used to evaluate competency. Competency assessment during routine review may be recorded using a checklist.

The laboratory director must ensure that the individuals performing competency assessments are qualified through education and experience to meet the defined regulatory requirements associated with the complexity of the testing.

- Testing personnel performing high complexity testing must be assessed by the section director, or individual meeting general supervisor requirements if delegated in writing by the section director.
- Testing personnel performing moderate complexity testing must be assessed by an individual meeting the qualifications of a technical consultant.

**Evidence of Compliance:**

✓ Records of competency assessment for new and existing testing personnel reflecting the specific skills assessed, the method of evaluation AND

✓ Written procedure defining the method and frequency for assessing competency

**REFERENCES**


**REVISED** 07/28/2015

**GEN.55525** Performance Assessment of Supervisors/Consultants  

**Phase II**

The performance of section directors/technical supervisors, general supervisors, and technical consultants is assessed and satisfactory.

**NOTE:** All responsibilities of section directors (as technical supervisors in laboratories performing high complexity testing) and technical consultants (in laboratories performing moderate complexity testing, but not high complexity testing) must be delegated by the laboratory director in writing. Unsatisfactory performance must be addressed in a corrective action plan.

The assessment may take the form of a checklist or other written record of performance of responsibilities, as defined by the individual's job description. If assessment of these individuals is not performed or there are inadequate or inconsistent records, a deficiency should also be cited for TLC.11425 (Director Responsibility - Delegation of Functions) in the Team Leader Assessment of Director and Quality Checklist.

If the individuals in these roles are also performing nonwaived patient testing, competency assessment requirements for testing personnel (GEN.55500) also apply, including all six elements of competency.

**Evidence of Compliance:**

- Job descriptions that list regulatory responsibilities AND
- Records of performance assessment

**REFERENCES**


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**GEN.57000** Competency Corrective Action  

**Phase II**

If testing personnel fail to demonstrate satisfactory performance on the competency assessment, the laboratory follows a plan of corrective action to retrain and reassess competency.

**NOTE:** If it is determined that there are gaps in the individual's knowledge, the employee should be re-educated and allowed to retake the portions of the assessment that fell below the laboratory's guidelines. If, after re-education and training, the employee is unable to satisfactorily pass the assessment, then further action should be taken which may include, supervisory review of work, reassignment of duties, or other actions deemed appropriate by the laboratory director.

**Evidence of Compliance:**

- Records of corrective action to include evidence of retraining and reassessment of competency
- Written procedure for competency assessment corrective action
PHYSICAL FACILITIES

Inspector Instructions:

- All Sections, including technical work areas, administrative space, storage areas, patient phlebotomy areas, etc. (adequate space, acceptable temperature/humidity, areas clean, adequate storage areas, adequate emergency power)

- Is the work area sufficient for you to perform your duties safely and accurately?

SPACE

Deficiencies in space should be recorded so there is incentive to improve. Deficiencies in space are regarded as minor unless they are so severe as to interfere with the quality of work or quality control activities or safety, in which case they become a Phase II deficiency. As laboratory operations expand over time, Phase I space deficiencies may become Phase II deficiencies by the time of the next inspection.

GEN.60000 Adequate Space Phase II

The general laboratory has adequate, conveniently located space so the quality of work, safety of personnel, and patient care services are not compromised.

REFERENCES
1) Koening AS. Medical laboratory planning and design. Northfield, IL: College of American Pathologists, 1992

**REVISED** 07/28/2015

GEN.60100 Adequate Space Phase I

All of the following areas have sufficient space and are located so there is no hindrance to the work.

1. Laboratory director
2. Staff pathologists and residents
3. Clerical staff
4. Section supervisors
5. Outpatient/ambulatory waiting and reception
6. Lavatories
7. Library, conference and meeting room
8. Personnel lounge and lockers

GEN.60150  Adequate Space

There is adequate space for:

1. Technical (bench) work
2. Instruments and equipment
3. Storage (records, slides, tissue, etc.)
4. Refrigerator/freezer storage
5. Media preparation, as applicable
6. Accessioning of potentially biohazardous specimens, as applicable
7. Radionuclide storage, as applicable
8. Microscopy and imaging, as applicable

ENVIRONMENT

Ambient or room temperature and humidity must be controlled to minimize evaporation of specimens and reagents, to provide proper growth conditions for room temperature incubation of cultures, and not to interfere with the performance of electronic instruments.

GEN.60250  Working Environment

The following are adequate for the facility.

1. Lighting
2. Water taps, sinks, drains
3. Electrical outlets
4. Ventilation
5. Gas and suction, when applicable

GEN.61300  Climate Control

The room temperature and humidity are adequately controlled in all seasons.

Evidence of Compliance:
✓ Temperature and humidity records, if specific ranges are required for instrument and/or reagent use

GEN.61350  Direct Sunlight

Exposure to direct sunlight is minimized.

NOTE: Direct sunlight should be avoided because of its extreme variability and the need for low light levels necessary to observe various computer consoles, etc. Lighting control should be sectionalized so general levels of illumination can be controlled in areas of the room, if desired.

GEN.61400  Hallway Obstructions

Passageways are unobstructed.

GEN.61500  Environment Maintenance

Floors, walls and ceilings are clean and well-maintained.
Laboratory General Checklist

GEN.61600  Environment Maintenance  Phase I

Bench tops, cupboards, drawers and sinks are clean and well-maintained.

COMMUNICATIONS

Communications within the laboratory should be appropriate for the size and scope of the laboratory. Messages should be transferred efficiently to all sections.

**REVISED** 07/28/2015
GEN.61750  Hand-Off Communication  Phase I

The laboratory implements a procedure for effective “hand-off” communication.

NOTE: The laboratory must have a procedure for communicating information about pending specimens, tests and patient care issues when responsibility is “handed off” from one person to another, such as at a change in shift, or when the responsibility for a case is transferred from one pathologist to another. The procedure should include provision for asking and responding to questions.

Evidence of Compliance:
✓ Logs or message boards showing communication between shifts

GEN.61800  Telephone/Computer Locations  Phase I

Telephones and computer terminals are conveniently located.

INVENTORY AND STORAGE OF SUPPLIES

GEN.61900  Inventory Control  Phase I

There is an effective supply inventory control system in operation.

NOTE: An effective inventory control system minimizes emergency requisitions and shortages of supplies.

REFERENCES
1) Chapman J. Saving money with computerized materials management. Advance/Lab. 1999;8(9):16-18

GEN.62000  Intralaboratory Storage  Phase I

The intralaboratory storage area is sufficient and free of clutter.

POWER

**REVISED** 07/28/2015
GEN.66100  Emergency Power  Phase I

Emergency power is adequate for the functioning of the laboratory.
NOTE: Emergency power supply must be adequate for refrigerators, freezers, incubators, etc., to ensure preservation of patient specimens. Depending on the type of testing performed in the laboratory, emergency power may also be required for the preservation of reagents, the operation of laboratory instruments, and the functioning of the data processing system.

REFERENCES

LABORATORY SAFETY

Requirements in this section cover the general safety program for the entire laboratory and must be answered for all laboratory sections. Non-compliance with any of these requirements in any one section of the laboratory represents a deficiency for the entire laboratory. Requirements related to safety features specific to an individual section will be found in the checklist for that section.

With respect to fire safety, if a checklist requirement conflicts with regulations of the Authority Having Jurisdiction (i.e. state and local fire codes), the regulations of the Authority Having Jurisdiction take precedence.

SAFETY POLICIES, PROCEDURES, AND RECORDS

Inspector Instructions:

- Sampling of safety policies and procedures
- Adequate emergency lighting
- How are your laboratory’s safe work practices reviewed?
- Is there a specific example of an occupational injury or illness that required medical treatment? What steps were taken to address the incident?

GEN.73200 Safety Policy and Procedure Approval

The laboratory director or designee reviews and approves all changes to the safety policies and procedures before implementation.
Safety Policy and Procedure Training

**REVISED** 07/28/2015
GEN.73300 Safety Policy and Procedure Training Phase II

There are records for the training of all laboratory workers in safety policies and procedures.

NOTE: A system to ensure that all personnel have read the policies and procedures is required and must form a portion of the orientation program for new personnel. Posting of specific warnings or hazards as appropriate is urged.

Evidence of Compliance:
✓ Records of personnel review of safety policies and procedures

REFERENCES

Safe Work Practices Review

**REVISED** 07/28/2015
GEN.73400 Safe Work Practices Review Phase II

There are records of periodic review (at least annually) of safe work practices to reduce hazards.

NOTE: Review must include bloodborne hazard control and chemical hygiene. If the review identifies a problem, the laboratory must investigate the cause and consider if modifications are needed to the safety policies and procedures to prevent reoccurrence of the problem or mitigate potential risk.

Evidence of Compliance:
✓ Safety committee minutes OR records of regular safety inspections OR incident reports and statistics OR another method defined by the laboratory director

Lab Accidents

GEN.73500 Lab Accidents Phase II

There are written policies and procedures for the reporting and recording of all laboratory accidents resulting in property damage or involving spillage of hazardous substances.

Occupational Injuries

**REVISED** 07/28/2015
GEN.73600 Occupational Injuries Phase II

There are written policies and procedures for the reporting of all occupational injuries or illnesses that require medical treatment (except first aid).

NOTE: For US laboratories subject to OSHA regulations, all workplace fatalities must be reported to the Occupational Safety and Health Administration (OSHA) within eight hours and work-related in-patient hospitalizations, amputations, or losses of an eye within 24 hours.

REFERENCES

Occupational Injury Evaluation

GEN.73700 Occupational Injury Evaluation Phase II

An evaluation of laboratory accident and occupational injury/illness reports is incorporated into the laboratory’s quality management program to avoid recurrence.

Evidence of Compliance:
✓ Records of report evaluation OR committee minutes with records of discussion
**REVISED** 07/28/2015
GEN.73800 Disaster Preparedness  Phase II

There are written policies and procedures defining the role and responsibilities of the laboratory in internal and external disaster preparedness.

REFERENCES

GEN.73900 Evacuation Plan  Phase II

There is a written comprehensive and workable evacuation plan specific for the laboratory.

NOTE: 1. This plan must cover all employees, patients and visitors, and must address the special needs of persons with disabilities. Evacuation routes must be clearly marked (Posting evacuation routes is optional). 2. Emergency lighting is adequate for safe evacuation of the laboratory.

REFERENCES
1) Occupational Safety and Health Administration. Exit routes, emergency action plans, and fire prevention plans: standard, 2002 [29CFR1910.38]

BLOODBORNE PATHOGENS

Inspector Instructions:

- Sampling of safety policies and procedures
- Sampling of records of hepatitis B vaccination or records declining the vaccination
- Sampling of personnel safety education records

- PPE usage

- What has your laboratory done to reduce or eliminate exposure to bloodborne pathogens during phlebotomy and laboratory testing?

GEN.74000 Bloodborne Pathogens  Phase II

The laboratory has written policies and procedures for infection control that comply with national, state, and local guidelines on occupational exposure to bloodborne pathogens and to the institution's exposure control plan.

NOTE: Universal or standard precautions must be used when handling all blood and body fluid specimens. The term "universal precautions" refers to a concept of bloodborne disease control requiring all human blood and other potentially infectious materials to be treated as if infectious for HIV, HBV, HCV or other bloodborne pathogens, regardless of the perceived "low risk" status of a patient or patient population. Alternative concepts in infection control are called
Body Substance Isolation (BSI) and Standard Precautions. These latter terms define all body fluids and substances as infectious. All health care workers must routinely use appropriate barrier precautions to prevent skin and mucous membrane exposure when contact with blood or other body fluids is anticipated. For laboratories subject to US regulations, policies and procedures must comply with the OSHA Standard on Bloodborne Pathogens.

Evidence of Compliance:
✓ Safety manual AND
✓ Records of universal precaution training for all personnel expected to have contact with body fluids

REFERENCES

**REVISED** 08/17/2016
GEN.74100  PPE Provision and Usage  Phase II

Appropriate personal protective equipment (gloves, gowns, masks and eye protectors, etc.) is provided and maintained in a sanitary and reliable condition in all technical work areas in which blood and body substances are handled and in circumstances during which exposure is likely to occur.

NOTE: 1. Appropriate personal protective equipment (PPE) are items that do not permit blood or other potentially infectious materials to pass through to the skin or reach the employee’s work clothes, skin, footwear, etc. In addition to fluid-resistant gowns, aprons may be required if exposure to large volumes of body fluids is anticipated. 2. OSHA requires gloves to be worn with each patient contact and changed after contact when performing vascular access, except when drawing voluntary blood donors. Hands must be cleaned after glove removal using an effective antimicrobial method.

REFERENCES

**REVISED** 08/17/2016
GEN.74200  PPE Instruction  Phase II

Personnel are instructed in the proper use of personal protective clothing/equipment (e.g. gloves, gowns, masks, eye protectors, footwear) and records are maintained.

NOTE: The required elements of training in the use of gloves include (a) Proper fitting of gloves; (b) Replacing gloves immediately when torn or contaminated; (c) Not washing or disinfecting gloves for reuse; (d) Using hypoallergenic gloves when indicated by patient or health care provider history; (e) Decontamination of hands after glove removal using an effective antimicrobial method.
Evidence of Compliance:
✓ Written policy for the use of PPE for specific tasks AND
✓ Records of personnel training for PPE

REFERENCES
1) Murray RL. Keep wearing your gloves. Med Lab Observ. 1994;Mar:80

**NEW** 08/17/2016
GEN.74250 Hand Cleaning Phase II
All personnel remove gloves and clean hands using an effective antimicrobial method after manipulating biological samples or after each patient contact.

REFERENCES

GEN.74300 Manual Manipulation of Needles Phase II
There is a written policy that prohibits the recapping, purposeful bending, breaking, removing from disposable syringes, or other manual manipulations of needles.

NOTE: Resheathing instruments or self-sheathing needles may be used to prevent recapping of needles by hand.

REFERENCES

GEN.74400 Eating/Mouth Pipetting Phase II
There is a written policy that prohibits smoking, eating, drinking, application of cosmetics and lip balm, manipulation of contact lenses, and mouth pipetting in all technical work areas.

REFERENCES

GEN.74500 Specimen Transport Procedures Phase II
There are written procedures for the procurement, transportation, and handling of patient specimens (e.g. blood, body fluids, tissue) to ensure that all specimens are submitted in an appropriately labeled and well-constructed container with a secure lid to prevent leakage during transport.

NOTE: Specimens sent through pneumatic tube systems must be sealed in fluid-tight bags. If pneumatic tube systems are used for transporting specimens, the laboratory must have procedures to respond to a spill within the tube, including appropriate decontamination measures.

REFERENCES

GEN.74600 Spill Handling Phase II
There are written procedures for handling spills of blood and other body fluids.

GEN.74700 Hepatitis B Vaccinations Phase II
Personnel reasonably expected to have direct contact with body fluids are identified and offered hepatitis B vaccinations free of charge.

Evidence of Compliance:
✓ Written policy offering the hepatitis B vaccination to employees AND
✓ Records of vaccination OR records signed by employees declining the vaccine

REFERENCES

GEN.74800 Viral Exposure Phase II
There is a policy for follow-up after possible and known percutaneous, mucous membrane or abraded skin exposure to HIV, HBV or HCV that includes the following elements.

1. HIV, HBV and HCV testing of the source patient after consent is obtained
2. Appropriate clinical and serologic evaluation of the healthcare worker
3. Consideration of appropriate prophylaxis for personnel acutely exposed to HIV, HBV or HCV, based upon medical indications, the serologic status and the informed consent of the health-care worker
4. Reporting of the exposure as required by law

Evidence of Compliance:
✓ Records of exposure follow-up consistent with policy

REFERENCES
OTHER INFECTIOUS HAZARDS

Inspector Instructions:
- Sampling of safety policies and procedures
- Sampling of sterilizing device monitoring records

GEN.74900  TB Exposure Plan

The laboratory has a written tuberculosis exposure control plan.

NOTE: This plan must include an exposure determination at defined intervals for all employees who may have occupational exposure to tuberculosis. Additional elements of the plan include engineering and work practice controls for hazardous activities that potentially may aerosolize Mycobacterium tuberculosis. Such activities include the handling of unfixed tissues in surgical pathology or autopsies, and processing of specimens in the microbiology section from patients with suspected or confirmed tuberculosis.

If respiratory protection is needed because of potential exposure to an infectious agent by aerosol or droplet, personnel must use either a properly fit-tested filter respirator (N-95 or higher) or a powered air-purifying respirator (PAPRS) equipped with high efficiency particulate air (HEPA) filters. Accurate fit testing is a key component of effective respirator use.

For laboratories subject to US requirement, the filter respirator must be NIOSH-approved.

REFERENCES

GEN.75000  Sterilizing Device Monitoring

All sterilizing devices are monitored periodically with a biologic indicator (or chemical equivalent) for effectiveness of sterility under conditions that simulate actual use.

NOTE: Each sterilizing device must be monitored periodically with a biologic indicator to measure the effectiveness of sterility. Chemical indicators that reflect sporicidal conditions may be used. The test must be performed under conditions that simulate actual use. One recommended method is to wrap the Bacillus stearothermophilus spore indicator strip in packaging identical to that used for a production run, and to include the test package with an actual sterilization procedure. Weekly monitoring is recommended.

Evidence of Compliance:
- ✓ Written procedure for monitoring sterilizing devices AND
- ✓ Records of monitoring at defined frequency
FIRE PREVENTION AND PROTECTION

Inspector Instructions:

- Sampling of safety policies and procedures
- Sampling of employee fire safety training records
- Sampling of employee fire extinguisher training records
- Automatic fire extinguisher systems, if required
- Two exit access doors, if required
- Audible automatic fire detection and alarm system
- Fire alarm station
- Portable fire extinguishers, where appropriate

GEN.75100 Fire Prevention Policies and Procedures Phase II

Policies and procedures are written and adequate for fire prevention and control.

NOTE: Fire safety plans must include the use of alarms, response to alarms, isolation of the fire, evacuation of the area, extinguishment of the fire, and the responsibilities of laboratory personnel for those elements.

REFERENCES

GEN.75200 Fire Separation Phase II

The laboratory is properly separated from inpatient areas and/or provided with automatic fire extinguishing (AFE) systems.

NOTE: For those facilities with no inpatients, no AFE is required.

For those facilities with inpatients, where the laboratory is separated by two-hour construction (rated at 1.5 hours) and Class B self-closing doors (SCD), no AFE system is required. An AFE system is required for those laboratories separated from inpatient areas by one-hour construction and class C SCD if flammable and combustible liquids are stored in bulk. An AFE system is always required if there are unattended laboratory operations employing flammable or combustible reagents. "Stored in bulk" means more than two gallons (7.5 L) of Class I, II, and IIIA liquids in safety cabinets and safety cans per 100 ft² (9.2 m²), or half that amount if not in safety containers. The following are the definitions of these Classes:

Class I flammable: any liquid that has a closed-cup flash point below 37.8°C and a Reid vapor pressure not exceeding 2068.6 mm Hg at 37.8°C as determined by ASTM D 323
Class II combustible: any liquid that has a flash point at or above 37.8°C and below 60°C
Class IIIA combustible: any liquid that has a flash point at or above 60°C but below 93°C

REFERENCES
GEN.75300 Fire Exit
Phase II

Each room larger than 1000 ft² (92.9 m²), or in which major fire hazards exist, has at least two exit access doors remote from each other, one of which opens directly into an exit route.

REFERENCES

GEN.75400 Fire Safety Training
Phase II

Fire safety training is performed for new employees, with a fire safety review conducted at least annually.

NOTE: Fire safety training must be recorded for all employees to show that they have been instructed on use and response to fire alarms and to execute duties as outlined in the fire safety plan. While fire exit drills are not required, physical evaluation of the escape routes must be performed annually, to ensure that fire exit corridors and stairwells are clear and that all fire exit doors open properly (i.e., not rusted shut, blocked or locked). Paper or computerized testing of an individual’s fire safety knowledge on the fire safety plan is acceptable; all personnel must participate at least once a year.

Evidence of Compliance:
✓ Records of participation for all employees in fire safety plan review at least annually (e.g. employee roster with dates of participation)

REFERENCES

**REVISED** 07/28/2015
GEN.75500 Fire Detection/Alarm
Phase II

There is an automatic fire detection and alarm system.

NOTE: 1. The system must connect to the facility’s overall system, where such a system exists. It must sound an immediate alarm in the event of smoke or fire. 2. The fire alarm is audible in all parts of the laboratory, including storage areas, lavatories, and darkrooms. 3. Laboratories employing hearing-impaired persons must have other means to alert these individuals, such as a visual alarm system.

REFERENCES

GEN.75600 Fire Alarm Station
Phase II

There is a fire alarm station in or near the laboratory.

NOTE: Alarm stations must be visible, unobstructed, and accessible.

REFERENCES

GEN.75700 Fire Extinguishers
Phase II
Appropriate portable fire extinguishers are provided for all areas in which flammable and combustible liquids are stored or handled.

NOTE: If gallon bottles of such materials are used, the minimum rating for Class B extinguishers is 10-B or higher. These are best located near or outside of doors leading to the area having solvent fire hazards.

REFERENCES

GEN.75800 Fire Extinguishers Phase II
If the fire safety plan includes laboratory staff use of fire extinguishers, personnel are instructed in the use of portable fire extinguishers.

NOTE: It is strongly recommended that instruction include actual operation of extinguishers that might be used in the event of a fire, unless prohibited by the local fire authority.

Evidence of Compliance:
✓ Records for fire extinguisher training

REFERENCES

ELECTRICAL SAFETY

Inspector Instructions:

- Sampling of electrical grounding records, if applicable

**REVISED** 07/28/2015
GEN.75900 Electrical Grounding Phase II
There are records that all laboratory instruments and appliances are adequately grounded and checked for current leakage before initial use, after repair or modification, and when a problem is suspected.

NOTE: Exceptions to these requirements are as follows:
1. Devices protected by an approved system of double insulation or its equivalent. Such devices must be distinctively marked
2. Devices connected to wall receptacles or circuit breakers with ground-fault circuit interrupter (GFCI) protection built-in need not be checked for current leakage
3. Equipment operating at 240 v must be checked for ground integrity only
Verification of electrical safety is required whenever the electrical/electronic systems of a powered device has been removed or altered. Hospital laboratories may follow ground checks and current leakage checks as performed in patient locations.

In addition, the US Occupational Safety and Health Administration (OSHA) requires that power cords of portable electrical equipment be visually inspected for external defects whenever relocated. Grounding configurations may not be bypassed by, for example, an adapter that interrupts the continuity of the grounding. If manufacturer's recommendations for grounding are available, they must be followed.

REFERENCES

CHEMICAL SAFETY

Inspector Instructions:

- Sampling of chemical safety policies and procedures
- Sampling of SDS (formerly MSDS) sheets
- Flammable and combustible liquids (properly stored)
- Acids and bases (properly stored)
- Sampling of hazardous chemicals (labeling)
- PPE usage

**REVISED** 07/28/2015

GEN.76000 Chemical Hygiene Plan Phase II

The laboratory has a Chemical Hygiene Plan (CHP) that defines the safety policies and procedures for all chemicals used in the laboratory.

NOTE 1: The laboratory director or designee must ensure that the laboratory has a written chemical hygiene plan (CHP) that defines the safety policies for all chemicals used in the laboratory. The plan must include evaluation of carcinogenic potential, reproductive toxicity, and acute toxicity. The plan must include specific handling requirements for all hazardous chemicals used in the laboratory.

The purpose of the CHP is to ensure that the hazards of all chemicals are evaluated, and that information concerning their hazards is transmitted to employers and employees. This transmittal of information is to be accomplished by means of comprehensive hazard communication programs, which are to include container labeling and other forms of warning, safety data sheets and employee training. An acceptable CHP contains the following elements:

1. Responsibilities of the laboratory director and supervisors
2. Designation of a chemical hygiene officer
3. Policies for all operations that involve chemicals
4. Criteria for the use of personal protective equipment and control devices
5. Criteria for exposure monitoring when permissible levels are exceeded
6. Provisions for medical consultations and examinations
7. Provision for training anyone working in the laboratory on the elements of the CHP
8. A copy of the OSHA Laboratory Standard, for laboratories subject to US regulations, or (for non-US laboratories) a copy of appropriate local standard

9. Evaluation of the carcinogenic potential, reproductive toxicity and acute toxicity for all chemicals used in the laboratory. The product label, safety data sheet (SDS), or for chemicals purchased prior to June 1, 2015 with no appropriate SDS, records of investigation by the safety officer may be used for this evaluation.

10. Specific handling requirements for all hazardous chemicals used in the laboratory

NOTE 2: For laboratories subject to US regulations, chemicals that must be handled as potential carcinogens include those defined by OSHA as “select carcinogens.” OSHA defines select carcinogens as any substance that is:

1. Regulated as a carcinogen by OSHA, has been classified as "known to be carcinogenic" by the NTP, or listed as a group I carcinogen by the IARC
2. Has been classified as "reasonably anticipated to be carcinogenic" by the NTP or listed as a group 2A or 2B carcinogen by the IARC if it meets the toxicological criteria listed in the January 31, 1990 Fed Register, pages 3319-3320

OSHA also requires special containment procedures for substances that are reproductive toxins or are acutely hazardous.

Authoritative sources include (but are not limited to) OSHA (Code of Federal Regulations. Title 29, Part 1910.1200 and 1450); NIOSH (Registry of Toxic Effects of Chemical Substances); the National Toxicology Program; the International Agency for Research on Cancer, and Safety Data Sheets.

Evidence of Compliance:
✓ Written evaluation of chemicals used in the laboratory for carcinogenic potential, reproductive toxicity and acute toxicity AND
✓ Written procedure for chemical fume hood function verification AND
✓ Records of testing

REFERENCES
Precautionary labels are present on the containers of all hazardous chemicals, indicating type of hazard and what to do if accidental contact occurs.

NOTE: The laboratory may use signs, placards, process sheets, batch tickets, operating procedures, or other such written materials in lieu of affixing labels to individual stationary process containers, as long as the alternative method identifies the containers to which it is applicable and conveys the information otherwise required to be on a label. The written materials shall be readily accessible to the employees in their work area throughout each work shift. It is not required to label portable containers into which hazardous chemicals are transferred from labeled containers, and which are intended only for the immediate use of the employee who performs the transfer. Existing labels on incoming containers of hazardous chemicals shall not be removed or defaced, unless the container is immediately marked with the required information.

Additional requirements for the labeling and expiration date of chemicals used for the preanalytic and analytic testing process, such as reagent preparation, are included in the Reagents section of the All Common Checklist. Deficiencies cited relating to the labeling and expiration of chemicals are cited in the checklist section where the chemicals are used.

REFERENCES

GEN.76300 PPE And Hazardous Materials Phase II

Personnel use the proper personal protective devices when handling corrosive, flammable, biohazardous, and carcinogenic substances.

NOTE: Such devices may include gloves of appropriate composition, aprons, and eye protection. Shoes or shoe covers must protect the entire foot in areas where splashing is expected.

REFERENCES

GEN.76400 Chemical Hazard Emergencies Phase II

Explicit instructions are posted, and appropriate supplies available, for the emergency treatment of chemical splashes and injuries and the control of chemical spills wherever major chemical hazards exist.

NOTE: Spill kits must be handled in accordance with manufacturer's instructions. If no expiration date is assigned, the spill kit must indicate the date it was put into service and the laboratory director or designee must periodically assess its usability.

REFERENCES

GEN.76500 Flammable Storage Phase II

Supplies of flammable and combustible liquids are reasonable for the laboratory's needs, and are properly stored.

NOTE: 1. In each laboratory area, up to one gallon (3.7 L) of Class I, II and IIIA liquids may be stored outside of fire-resistant cabinets for each 100 ft² (9.2 m²) of space defined by fire-resistant walls/doors. Up to two gallons (7.5 L) of Class I, II, and IIIA liquids may be stored in safety cans and safety cabinets for each 100 ft² (9.2 m²). These amounts may be doubled if there is an automatic fire suppression system (e.g. sprinklers). For example: a 1000 ft² (92.9
m³) laboratory defined by fire resistant walls/doors can store 10 gallons (37.8 L) outside a safety cabinet and 20 gallons (75.7 L) inside a safety cabinet and double those numbers if there is an automatic fire suppression system. 2. Safety cans should be used for bulk storage of flammable and combustible liquid (National Fire Protection Association classes I and II). Metal or DOT-approved plastic containers provide an intermediate level of hazard containment between glass and safety cans. One pint (0.4 L) of a highly volatile solvent such as isopentane, stored in glass has about the same ignitability risk as two gallons (7.5 L) stored in safety cans. Safety cans should be used instead of glass bottles if the purity required does not mandate glass storage.

REFERENCES

GEN.76600 Volatile Solvent Ventilation Phase II

Storage areas and/or rooms where volatile solvents are used are adequately ventilated.

NOTE: Areas where flammable liquids are used must be ventilated for protection of employee health, as well as fire prevention. Areas where flammable liquids are stored should be ventilated primarily for fire protection. Storage cabinets do not need to be vented, but if they are vented the duct system must be explosion proof.

REFERENCES

**REVISED** 07/28/2015

GEN.76700 Acid/Base Storage Phase II

Supplies of concentrated acids and bases are stored safely.

NOTE: 1) Storage must be below eye level. Storage near the floor is recommended. 2) Strong acids and bases must not be stored under sinks, where contamination by moisture may occur. 3) Storage containers of acids and bases should be adequately separated to prevent a chemical reaction in the event of an accident/spill/leak. 4) Bottle carriers are used to transport all glass containers larger than 500 mL that contain hazardous chemicals.

REFERENCES

COMPRESSED GASES

Inspector Instructions:

- Gas cylinders (properly stored and secured)

GEN.76800 Gas Cylinder Storage Phase II

Compressed gas cylinders are secured to prevent accidental falling and damage to the valve or regulator.

GEN.76900 Flammable Gas Cylinders Phase II
Flammable gas cylinders, if inside a health care facility, are stored properly.

NOTE: Proper storage practices include:
1. Storage in a separate, ventilated room or enclosure
2. Cylinders are positioned well away from open flame or other heat sources, not in corridors and not within exhaust canopies

REFERENCES

RADIATION SAFETY

Inspector Instructions:

- Sampling of radiation safety policies and procedures

GEN.77000 Radiation Safety Manual

There are written policies and procedures adequate for radiation safety.

NOTE TO INSPECTOR: The following requirement applies to laboratories that do not perform anatomic pathology on-site, and for whom the Anatomic Pathology checklist is not used.

GEN.77100 Radioactive Material Handling

There are specific policies and procedures for the safe handling of tissues that may contain radioactive material (e.g. sentinel lymph nodes, breast biopsies, prostate "seeds", etc.).

NOTE: These policies and procedures should be developed in conjunction with the institutional radiation safety officer, and must comply with any state regulations for the safe handling of tissues containing radionuclides. The policies and procedures should distinguish between low radioactivity specimens such as sentinel lymphadenectomy and implant devices with higher radiation levels.

REFERENCES
ENVIRONMENTAL SAFETY

Inspector Instructions:

READ

● Ergonomic evaluation

OBSERVE

● Emergency eyewash available and tested properly

ASK

● How does your laboratory prevent workplace-related musculoskeletal disorders?

GEN.77200  Ergonomics  Phase II

There is a written ergonomics program to prevent musculoskeletal disorders (MSDs) in the workplace through prevention and engineering controls.

NOTE: The program may include training of employees about risk factors, identifying physical work activities or conditions of the job commonly associated with work-related MSDs, and recommendations for eliminating MSD hazards. Laboratory activity, workplace and equipment (e.g. chairs, laboratory workstations, computer keyboards, and displays) should be designed to reduce the risks of ergonomic distress disorders and accidents.

Evidence of Compliance:
✓ Records of ergonomic evaluation including recommendations for eliminating MSD hazards and appropriate corrective action based on assessment findings

REFERENCES
2) U.S. Dept. of Labor, Occupational Safety and Health Administration. Ergonomic safety and health program management guideline. 54 Fed Register 3904 (1989), modified at 29CFR1910

GEN.77300  Excessive Noise  Phase II

The laboratory has a policy to protect personnel from excessive noise levels.

NOTE: The laboratory should provide protection against the effects of noise exposure when sound levels equal or exceed an 8-hour time-weighted average sound level of 85 decibels. The laboratory should monitor noise exposure if there is an indication that excessive noise levels are present (for example, when noise levels exceed 85 decibels, people have to shout to be heard).

REFERENCES

**REVISED** 07/28/2015
GEN.77400  Emergency Eyewash  Phase II

The laboratory has adequate plumbed or self-contained emergency eyewash facilities in every area where there are hazardous chemicals as defined by the laboratory’s chemical hygiene plan (e.g. chemicals that are irritating, corrosive, toxic by contact or absorption). Testing records are maintained.

NOTE 1: The eyewash facility includes the following:

1. No greater than 10 seconds travel distance from areas in the laboratory where hazardous chemicals are present
2. Capable of delivering 1.5 L per minute for 15 minutes
3. Flow is provided to both eyes simultaneously
4. Nozzles or covers to protect from airborne contaminants
5. Hands-free flow once activated
6. Signage for location of eyewash
7. Unobstructed path with unlocked doors opening in the direction of the eyewash
8. Plumbed systems are protected from unauthorized shut off
9. Tepid fluid temperature (Water temperature should be between 15°C and 37°C (60°F and 100°F). Actual temperature recording is not required.)
10. Plumbed systems are activated weekly
11. Self-contained units are visually examined weekly

NOTE 2: The manufacturer's specifications should be available.

REFERENCES

OTHER HAZARDS

Inspector Instructions:

- Sampling of safety policies and procedures
- UV light signage

GEN.77500  Liquid Nitrogen  Phase II

Adequate policies, procedures, and practices are in place for the use of liquid nitrogen.

NOTE: Practices for the safe handling of liquid nitrogen include:

1. The mandatory use of appropriate gloves, shielding of all skin and the use of a face shield when decanting or entering an open container of LN
2. Storage and use of all containers of LN only in well-ventilated areas
3. Availability of a Safety Data Sheet
UV Light Exposure

There are written policies and procedures to prevent or reduce ultraviolet light exposure from instrument sources.

NOTE: UV light may cause corneal or skin burns from direct or deflected light sources. Wherever UV light sources are used, suitable and adequate personal protective equipment must be provided, and appropriate approved signage displayed. Laboratories may obtain information on safety from manufacturers of devices that emit UV light.

A suggested sign for display is: Warning: This device produces potentially harmful ultraviolet (UV) light. Protect eyes and skin from exposure.

Evidence of Compliance:
✓ Warning signage on source equipment AND
✓ Suitable PPE available, as required

REFERENCES

Latex Allergy

The laboratory has a written program to protect personnel and patients from allergic reactions from exposures to natural rubber latex in gloves and other products.

NOTE: The latex program should address at least the following elements:

1. Selection of products and implementation of work practices that reduce the risk of allergic reactions. If latex gloves are used, the employer should provide reduced protein, powder-free gloves to protect workers from infectious materials
2. Provision of education programs and training materials about latex allergy
3. Evaluation of current prevention and control strategies whenever a worker is diagnosed with latex allergy

Evidence of Compliance:
✓ Records of employee education/training on latex allergies AND
✓ Records of evaluation of the plan, when appropriate

REFERENCES
WASTE DISPOSAL

Inspector Instructions:

- Sampling of waste disposal policies and procedures
- How does your laboratory dispose of sharps?
- How does your laboratory dispose of hazardous chemicals?

GEN.77800  Hazardous Chemical Waste Disposal  Phase II

Written policies and procedures are adequate for hazardous chemical waste disposal.

NOTE: 1. The laboratory is responsible for all real or potential hazards of wastes at all stages of disposal including transportation and final disposition. 2. The method for the disposal of all solid and liquid wastes is in compliance with local, state and federal regulations. (Whether or not laboratory management is responsible for waste disposal, the laboratory should have documentation that the facility is in compliance with all applicable regulations. Prevailing local, state and federal (EPA) regulations should be reviewed by the laboratory director, safety officer or hospital engineer to ensure that the laboratory is in compliance with regulations.)

Evidence of Compliance:
✓ Records of review of regulations for compliance

REFERENCES

GEN.77900  Biohazard Disposal Containers  Phase II

All infectious wastes (e.g. glassware, blood collection tubes, microbiologic and tissue specimens) and other solid or liquid waste or refuse are discarded into "biohazard"-labeled containers that do not leak and have solid, tight-fitting covers that are applied before transport from the laboratory work area for storage and disposal.

NOTE: All infectious wastes must be incinerated or appropriately decontaminated before being sent to a sanitary landfill. Stool and urine waste may be discarded into the sanitary sewerage system.

REFERENCES
Sharps Disposal

Sterile syringes, needles, lancets, or other blood-letting devices (“sharps”) that are capable of transmitting infection are used once only, and all waste sharps are discarded in puncture-resistant containers that are easily accessible, located in areas where needles are commonly used, and properly labeled to warn handlers of the potential hazard.

NOTE: Shearing or breaking of contaminated sharps is prohibited. Bending, recapping, or removing contaminated needles is prohibited as a general practice. Needles are expected to be used and immediately discarded, un-recapped, into accessible sharps containers.

REFERENCES

BIOREPOSITORIES

INTRODUCTION

The General Checklist applies to all sections of the biorepository. An inspection of a biorepository section or department will include the Biorepository Checklist.

The requirements in this section only apply to biorepositories enrolled in the Biorepository Accreditation Program.

POLICIES AND PROCEDURES

Inspector Instructions:

- Representative sample of procedures for completeness and biorepository director review. Current practice must match contents of policies and procedures.
- Document control policy
- Privacy and confidentiality policies and procedures
- How do you access procedures?
- What procedure has most recently been implemented or modified?
- How do you ensure all copies of procedures are up to date?
- How are changes in procedures recorded and communicated to staff?
- How does the facility protect patient information?
Identify a newly implemented procedure in the prior two years and follow the steps through authoring, director review and staff training.

**NEW** 08/17/2016
GEN.80000 Procedure Manual Phase II

A complete procedure manual is available in a paper-based, electronic, or web-based format at the workbench or in the work area.

NOTE 1: The use of inserts provided by manufacturers is not acceptable in place of a procedure manual. However, such inserts may be used as part of a procedure description, if the insert accurately and precisely describes the procedure as performed in the biorepository. Any variation from this printed or electronic procedure must be detailed in the procedure manual. In all cases, appropriate reviews must occur.

NOTE 2: A manufacturer’s procedure manual for an instrument/reagent system may be acceptable as a component of the overall departmental procedures. Any modification to or deviation from the procedure manual must be clearly recorded and approved.

NOTE 3: Card files or similar systems that summarize key information are acceptable for use as quick reference at the workbench provided that:
- A complete manual is available for reference
- The card file or similar system corresponds to the complete manual and is subject to document control

NOTE 4: Electronic manuals accessed by computer are fully acceptable. There is no requirement for paper copies to be available for the routine operation of the biorepository as long as the electronic versions are readily available to all personnel and personnel have been trained on how to access them. However, procedure manuals must be available to biorepository personnel when the electronic versions are inaccessible (e.g. during biorepository information system or network downtime); thus, the biorepository must maintain paper copies, electronic copies on CD or other digital media, or have an approved alternative mechanism to access web-based files during network downtimes. All procedures, in either electronic or paper form, must be readily available for review by the inspector at the time of the CAP inspection.

Electronic procedure manuals and electronic copies of procedures are subject to proper document control (see GEN.80600), and there must be records of biennial review. Records of review of electronic procedures may include statements such as “reviewed by [name of reviewer] on [date of review] in the electronic record.” Alternatively, paper review sheets may be used to record review of electronic procedures. Record of review by a secure electronic signature is NOT required.

REFERENCES
There are records of review of all technical policies and procedures by the current director or designee at least every two years.

NOTE: Only technical policies and procedures are addressed in this requirement. Biennial review is not required for other controlled documents.

The director must ensure that the collection of policies and procedures is complete, current, and has been thoroughly reviewed by a knowledgeable person. Technical approaches must be scientifically valid and clinically relevant. To minimize the burden on the biorepository and reviewer(s), it is suggested that a schedule be developed whereby roughly 1/24 of all policies and procedures are reviewed monthly. Paper/electronic signature review must be at the level of each procedure, or as multiple signatures on a listing of named procedures. A single signature on a Title Page or Index is not a sufficient record that each policy or procedure has been carefully reviewed. Signature or initials on each page of a policy or procedure is not required.

**NEW** 08/17/2016
GEN.80300 New Procedure Review Phase II

The director reviews and approves all new policies and procedures, as well as substantial changes to existing documents, before implementation.

NOTE: Current practice must match the policy and procedure documents.

**NEW** 08/17/2016
GEN.80400 New Director Procedure Review Phase II

If there is a change in directorship of the biorepository, the new director ensures (over a reasonable period of time) that biorepository procedures are well documented and undergo appropriate review.

**NEW** 08/17/2016
GEN.80500 Knowledge of Policies and Procedures Phase II

The biorepository has a defined process and records indicating that all personnel are knowledgeable about the contents of the policies and procedures (including changes) relevant to the scope of their biorepository activities.

NOTE: This does not specifically require annual procedure sign-off by testing personnel. The form of this system is at the discretion of the director.

Evidence of Compliance:
✓ Relevant quizzes and results OR record confirming competency AND
✓ Systems to record policy and procedure changes AND
✓ Records of receipt/training in either paper or electronic format

**NEW** 08/17/2016
GEN.80600 Document Control Phase II

The biorepository has a document control process to manage policies, procedures, and forms that are subject to CAP accreditation.

NOTE: The document control system must ensure that only current policies, procedures, and forms are in use.

It may be helpful for some biorepositories to maintain a control log listing all current policies, procedures, and forms with the locations of copies. The control log may contain other information as appropriate, such as dates when policies and procedures were placed in service, schedule
of review, identity of reviewer(s), and dates when policies and procedures were discontinued or superseded.

**Evidence of Compliance:**
✓ Electronic documents on a shared file OR commercial document system OR a biorepository developed organized system

**REFERENCES**

**NEW** 08/17/2016

GEN.80700 Discontinued Procedure

When a procedure is discontinued or replaced, a paper or electronic copy is maintained for at least two years, recording initial date of use, and retirement date.

**QUALITY MANAGEMENT**

The biorepository must have a written quality management program to systematically ensure the quality of services. In biorepositories that are part of a larger institution (e.g. a hospital), the biorepository quality management program may be integrated with the institutional program.

**Inspector Instructions:**
- Sampling of QM policies and procedures
- Procedure for communication of employee concerns
- Sampling of quality indicators with follow-up actions when targets are not achieved
- Annual appraisal of effectiveness of the QM Program
- Records of instructions/recommendations from IRBs and clients
- CAP sign regarding the reporting of quality concerns
- How are IRB and client concerns and recommendations addressed? What were the results and what actions were taken as a result of the findings?
- Is there a specific example when problems were identified that could have interfered with participant/client care or safety?
- If any problems are found during review of quality measurements, or when asking questions, further evaluate the biorepository’s investigation and resolution, including root cause analysis and associated risk-reduction activities when applicable

GEN.81000 Written QM Program

The biorepository has a written quality management (QM) program.
NOTE: There must be a document that describes the overall QM program. The document need not be detailed, but should spell out the objectives and essential elements of the QM program. If the biorepository is part of a larger organization, the biorepository QM program is coordinated with the organization’s QM plan.

REFERENCES

GEN.81100 QM Implementation
Phase II

The QM plan is implemented as designed and is reviewed annually by the director for effectiveness.

NOTE: 1) This requirement pertains to biorepositories that have been CAP accredited for more than 12 months. 2) Appraisal of program effectiveness may be evidenced by an annual written report, revisions to policies and procedures, or revisions to the QM plan, as appropriate.

Evidence of Compliance:
✓ Evidence that the plan has been implemented as designed requires all of the following:
  • quality measurements and assessments specified in the plan are being substantially carried out;
  • there is evidence of active review of quality measurements;
  • any interventions or changes to operations that are specified in the plan have been carried out as scheduled, or the reason for delay recorded; AND
  • any communication of information that is required by the plan have taken place

GEN.81200 QM Error and Incident Management
Phase II

The QM system includes a program to identify and evaluate errors, incidents, and other problems that may interfere with functions of the biorepository.

NOTE: There must be an organized program for recording of problems involving the biorepository that are identified internally, as well as those identified through outside sources such as complaints from other study collaborators or researchers. Any problem that could potentially interfere with research result integrity or safety must be addressed. Scientific impact, rather than business or management issues, should be emphasized.

The biorepository must:
1. Record investigation and resolution of these problems
2. Perform root cause analysis of any unexpected sentinel events
3. Be able to demonstrate any appropriate risk-reduction activities based on such root cause analyses

REFERENCES

GEN.81300 QM Indicators of Quality
Phase II

The QM program includes monitoring key indicators of quality.

NOTE: Key indicators are those that reflect activities critical to expected outcome or that have been problematic in the past. The biorepository must record comparison of performance of selected indicators against a benchmark, where available and applicable. New programs or services should be measured to evaluate their impact on service. The number of monitored
indicators should be consistent with the biorepository’s scope of service. Action plans should be developed for any indicator in which the biorepository falls outside a predetermined level.

**NEW** 08/17/2016

**GEN.81325** Correction of Biorepository Records  Phase II

The biorepository follows a written policy for the management and correction of biorepository records, including quality control data, temperature logs, and intermediate test results or worksheets.

**NOTE:** Biorepository records and changes to such records must be legible and indelible. Original (erroneous) entries must be visible (i.e. erasures, white and correction fluid are unacceptable) or accessible (e.g. audit trail for electronic records). Corrected data, including the identity of the person changing the record and when the record was changed, must be accessible to audit.

**Evidence of Compliance:**
✓ Records of corrections to biorepository records following the policy

**REVISED** 07/28/2015

**GEN.81350** Hand-Off Communication  Phase I

The biorepository implements a procedure for effective “hand-off” communication.

**NOTE:** The biorepository must have a procedure for communicating information about pending processes, quality or operational issues when responsibility is “handed off” from one person to another, such as at a change in shift, or when the responsibility for a case is transferred from one pathologist to another. The procedure should include provision for asking and responding to questions.

**Evidence of Compliance:**
✓ Logs or message boards showing communication between shifts or departments

**GEN.81400** Employee Quality Communication  Phase II

The biorepository has a procedure for employees, participants, and researchers to communicate concerns about research misconduct, quality, and safety to management.

**NOTE:** The biorepository must have a procedure that encourages employees to communicate any concerns or complaints with respect to the research misconduct, quality and safety. The investigation and analysis of employee complaints and suggestions, with corrective or preventive action as appropriate, should be a part of the quality management program and be specifically addressed in quality management records.

**Evidence of Compliance:**
✓ Records of employee complaints (if any) with appropriate follow up

**GEN.81500** CAP Sign  Phase II

The biorepository posts the official CAP sign regarding reporting of quality concerns.

**NOTE:** The biorepository must prominently post the official CAP sign regarding the reporting of quality concerns to CAP.

While personnel should report concerns to biorepository management, the biorepository must ensure that all personnel know that they may communicate with CAP directly if they have a concern not addressed by biorepository management, and that CAP holds such communications in strict confidence. In addition, the biorepository must have a policy prohibiting harassment or
punitive action against an employee in response to a complaint or concern made to CAP or other regulatory organization regarding biorepository quality or safety.

The dedicated, confidential CAP telephone line for quality or safety concerns is 866-236-7212 (US, toll-free) and 847-832-7533 (international).

Official CAP signs may be obtained by calling 800-323-4040 option 1#.

**REVISED** 07/28/2015

**GEN.81900** Terms of Accreditation

The biorepository has a policy that addresses compliance with the CAP terms of accreditation.

NOTE: The CAP terms of accreditation are listed in the biorepository's official notification of accreditation. The policy must include notification of CAP regarding the following:

1. Investigation of the biorepository by a government entity or other oversight agency, or adverse media attention related to biorepository performance; notification
must occur no later than two working days after the biorepository learns of an investigation or adverse media attention. This notification must include any complaint investigations conducted or warning letters issued by any oversight agency (i.e. FDA, OSHA, FAA).

2. Change in biorepository test menu (notification must occur prior to implementing scope of service changes)

3. Change in location, ownership or directorship of the biorepository; notification must occur no later than 30 days prior to the change(s); or, in the case of unexpected changes, no later than two working days afterwards

Evidence of Compliance:
✓ Records of notification, if applicable

**REVISED** 07/28/2015
GEN.81950 Selection and Evaluation of Services  Phase II

There is a written procedure for evaluating and selecting biospecimen source sites, contracted services, or referral laboratories, to ensure that specimens and test results are managed in a quality environment.

NOTE:
1. A written qualification process suitable for the process being performed is in place, e.g. vendor qualification, a system for the biorepository director to approve the service provider.

2. Specimens used for patient treatment decisions, including those from clinical trials, should be obtained or sent to a laboratory accredited by CAP, accredited to an established international standard from a recognized organization, or certified by an appropriate government agency.

3. It is the responsibility of the biorepository director or designee to monitor the quality of test results received from contracted services or referral laboratories.

Evidence of Compliance:
✓ Records of evaluation or qualification (e.g. certification, publications, audits or biorepository director-approved records of acceptable quality)

PERSONNEL

The biorepository should have an organizational chart, personnel policies, and job descriptions that define qualifications and duties for all positions. Personnel files should contain records of educational qualifications, references, training, competency assessments, health records and continuing education records for each employee. Ideally, these files should be located in the biorepository. However, they may be kept in the personnel office or health clinic if the biorepository has ready access to them (i.e. they are easily available to the inspector).

Inspector Instructions:

- Sampling of personnel policies and procedures
- Organizational chart or narrative description
- Sampling of all personnel files
● Do you have a specific example of an employee who demonstrated unacceptable competency assessments? What were the corrective actions?
● What continuing education classes are available to employees?

**DIRECTOR QUALIFICATIONS**

**GEN.82000  Director Qualifications**  
Phase II

The qualifications of director of the biorepository are appropriate for the scope of activities.

**NOTE:** The organization's leadership must have defined the appropriate training, experience, and/or educational credentials. The director's experience and qualifications must meet the institutional policy for the degree of responsibility acceptable to operate and manage the scope of the biorepository.

**REVISED** 08/17/2016

**GEN.82100  Delegation of Functions**  
Phase I

Delegation of the biorepository director's functions or responsibilities is in writing.

**NOTE:** Functions that may be delegated include duties, such as review of QC processes, ensuring that IRB protocols are followed, and implementation of the quality management plan. The biorepository director remains responsible that all persons performing delegated functions are qualified to do so and that delegated functions are properly carried out.

Functions that may not be delegated include provision of appropriately trained supervisory and technical staff and the identification of their responsibilities. The biorepository director must document personal, on-site assessment of physical and environmental conditions and the adequacy of staffing.

The responsibilities and duties of supervisors, consultants, and personnel involved in the biorepository services must be defined in writing, with records of authorization to perform the services and the level of supervision required, as applicable.

If there are multiple occasions when delegated duties are not being properly performed by the designee and there is a lack of consistency in performing corrective action, the team leader should cite this requirement as a deficiency, in addition to the specific checklist requirement(s) that relates to the duty not being performed (e.g. QC review). This may be overarching rather than a single issue.

**Evidence of Compliance:**
✓ Policy or statement signed by the biorepository director authorizing individuals by name or job title to perform tasks on behalf of the biorepository director AND
✓ Records showing that delegated tasks are performed by the designee, as required

**DIRECTOR OVERSIGHT RESPONSIBILITIES**

**GEN.82200  Director Responsibility/Authority**  
Phase II
The biorepository director has sufficient responsibility and authority to implement and maintain the standards of the College of American Pathologists.

NOTE: Examples of how the team leader may obtain information on the director's responsibility and authority include: interviews with the biorepository director, institution's administration, biorepository management and biorepository supervisory staff, review of the biorepository organizational chart, and review of minutes of quality management and other biorepository meetings.

GEN.82300 Effective QM Phase II

The biorepository director ensures an effective quality management program for the biorepository.

NOTE: The biorepository director must be involved in the design, implementation, and oversight of the biorepository's quality management program. The program must include monitoring of key indicators, investigation of problems, with corrective and preventive actions as appropriate; maintenance of safety; and ensuring the quality data.

Evidence of Compliance:
Written QM plan covering all areas of the biorepository and addressing all phases of testing

REFERENCES

GEN.82400 Policy and Procedure Development Phase II

The biorepository director is involved in development of all policies and procedures.

**REVISED** 07/28/2015

GEN.82500 Director's Responsibilities Phase II

The biorepository director must have policies to safeguard that:

1. IRB protocols and policies are upheld
2. HIPAA is not violated
3. Clinical care is not compromised in the process of procuring biospecimens
4. Basic ethical standards related to biospecimen collection and distribution are upheld (e.g. no selling tissues for a profit on the side)

GEN.82600 Director Responsibility - Education/R&D Phase II

The biorepository director ensures provision of educational programs, strategic planning, and research and development appropriate to the needs of the biorepository.

**REVISED** 07/28/2015

GEN.82700 Director Responsibility - Personnel Phase II

The biorepository director ensures that there are sufficient personnel with adequate training and experience to meet the needs of the biorepository, and that such training and experience is recorded.

REFERENCES
GEN.82800 Director Responsibility - Safe Environment

The biorepository director ensures implementation of a safe environment in compliance with good practice and applicable regulations.

NOTE: The biorepository director must ensure compliance with OSHA and state/local regulations, as well as other applicable safety regulations.

**DIRECTOR NOT ON-SITE FULL TIME**

NOTE TO THE TEAM LEADER: The following requirements apply to biorepository directors who are not present full-time at the biorepository.

**REVISED** 08/17/2016

GEN.82900 Director Off-Site

There is a written agreement defining the frequency of, and responsibilities for, activities to be performed by the biorepository director during on-site visits and remotely, with records of the director’s completed activities.

Evidence of Compliance:
✓ Records that show the frequency of on-site visits AND
✓ Meeting minutes showing director participation

**REVISED** 08/17/2016

GEN.83000 Director Visit

The involvement of the biorepository director in the biorepository's activities conducted during on-site visits or remote consultation follows the written policy or agreement and is considered adequate by the biorepository staff and the inspection team.

NOTE: The requirement is not met if the biorepository management and staff identify inadequate oversight by the biorepository director. If activities are conducted remotely, the biorepository director must ensure that there is an effective communication mechanism in place between the biorepository director and biorepository management and staff.

Evidence of Compliance:
✓ Minutes from meetings with staff OR
✓ Records of conformance with specified director responsibilities

**OPERATIONAL LEADERSHIP/MANAGEMENT SECTION**

GEN.83100 Leadership/Management Qualifications

Leadership/management have qualifications equal to the expertise of the level of service of the biorepository.

GEN.83200 Organizational Chart

There is an organizational chart for operational leadership, or a narrative description that describes the reporting relationships among the owner or management, the biorepository director, and management/leadership staff, as appropriate.
GEN.83300  Description of Duties  
Phase I

Duties for all staff are described in writing so that it is clear who is responsible for consent, banking, transport, inventory, triage, and release on any given day.

GEN.83400  Staff Qualifications  
Phase II

The biorepository director must define the minimum qualifications for each role in the biorepository based on the level of service of the biorepository.

Evidence of Compliance:
✓ Written description of minimum qualifications

GEN.83500  Continuing Education  
Phase I

There is a functional, continuing biorepository education program adequate to meet the needs of the biorepository’s mission and/or goals as outlined by the biorepository director.

NOTE: Continuing education may take place within the institution or at an offsite presentation.

Evidence of Compliance:
✓ Written policy for continuing education

REFERENCES

**REVISED** 08/17/2016

GEN.83600  Personnel Records  
Phase II

Personnel files are maintained on all current technical personnel and personnel records include all of the following items, as applicable.

1. Copy of academic diploma, transcript, or primary source verification (PSV) reports confirming credentials, if applicable (Refer to the NOTE for use of PSV reports)
2. Personnel license, if required by state
3. Summary of training and experience
4. Certification, if required by state or employer
5. Description of current duties and responsibilities as specified by the biorepository director: a) Procedures the individual is authorized to perform, b) Whether supervision is required for specimen processing, test performance or result reporting, c) Whether supervisory or director review is required to report participant results
6. Records of continuing education
7. Records of radiation exposure where applicable (such as with in vivo radiation testing), but not required for low exposure levels such as certain in-vitro testing
8. Work-related incident and/or accident records
9. Dates of employment

NOTE: All records in either electronic or paper form must be readily available for review by the inspector at the time of the CAP inspection.

If PSV reports are used, the biorepository must have a defined system for reviewing the reports, with written criteria for acceptance. PSV is typically performed by a third-party agent or company that directly contacts institutions and former employers to verify training and experience, such as diplomas, board certification, licensure, and reported work history. PSV reports confirming the required qualifications may be retained in lieu of obtaining paper copies of these records. If there
are required elements for the qualification that the PSV report does not adequately verify (e.g. transcripts, educational equivalency for personnel trained outside of the US, or reports lacking the type of degree earned), there must be records showing that qualifications are met using other means.

REFERENCES

GEN.83700 Initial Training

There are records of satisfactory completion of initial training of all staff on all instruments/methods applicable to their designated job.

NOTE: The records must show that training specifically applies to the duties performed by each individual.

Retraining must occur when problems are identified with employee performance.

GEN.83800 Competency Assessment

The competency of each person to perform his/her assigned duties is assessed.

NOTE: Prior to the initiation of job duties and the performance of new duties, each individual must have training and be evaluated for proper performance of duties as required in GEN.83700.

After an individual has performed his/her duties for one year, competency must be assessed annually. Retraining and reassessment of employee competency must occur when problems are identified with employee performance. Elements of competency assessment include but are not limited to:

1. Direct observations of routine process and procedure performance, including as applicable, participant identification and preparation; and specimen collection, handling, processing
2. Review of results or worksheets, quality control records, and preventive maintenance records
3. Direct observation of performance of instrument maintenance and function checks, as applicable, and
4. Evaluation of problem-solving skills

Many of the elements of competency assessment are performed during routine supervisory review of an employee throughout the year. Records of these elements, including adherence to biorepository policies and procedures, observation of test performance, results reporting, instrument maintenance, review of worksheets, recording QC, and demonstration of taking appropriate corrective actions are examples of daily activities that can be used to demonstrate competency. If elements of competency are assessed by routine supervisory review, the competency procedure must outline how this routine review is used to evaluate competency. Competency assessment by routine supervisory review may be recorded using a checklist.

Evidence of Compliance:
✓ Records of competency assessment for new and existing employees reflecting the specific skills assessed, the method of evaluation

REFERENCES

GEN.83900 Competency Corrective Action

Phase II
If an employee fails to demonstrate satisfactory performance on the competency assessment, the biorepository follows a plan of corrective action to retrain and reassess competency.

NOTE: If it is determined that there are gaps in the individual's knowledge, the employee should be re-educated and allowed to retake the portions of the assessment that fell below the biorepository's guidelines. If, after re-education and training, the employee is unable to satisfactorily pass the assessment, then further action should be taken which may include, supervisory review of work, reassignment of duties, or other actions deemed appropriate by the biorepository director.

Evidence of Compliance:
✓ Records of corrective action to include evidence of retraining and reassessment of competency
✓ Written procedure for competency assessment corrective action

PHYSICAL FACILITIES

Deficiencies in space should be recorded so there is incentive to improve. Deficiencies in space are regarded as minor unless they are so severe as to interfere with the quality of work or quality control activities and safety, in which case they become a Phase II deficiency. As biorepository operations expand over time, Phase I space deficiencies may become Phase II deficiencies by the time of the next inspection.

Ambient or room temperature and humidity must be controlled to minimize evaporation of specimens and reagents, to provide proper growth conditions for room temperature incubation of cultures, and not to interfere with the performance of electronic instruments.

Inspector Instructions:
- Floor plan and equipment locations
- Overview of Building Automation System (BAS), if available
- Sampling of electrical grounding records, if applicable
- Physical facility (adequate space, acceptable temperature/humidity, areas clean, adequate storage areas, adequate emergency power)
- Perimeter security and access security to specific specimen collections
- Is the work area sufficient for you to perform your duties safely and accurately?

GEN.84000 Restricted Access

Access to the biorepository is restricted to authorized individuals.

NOTE: This may be accomplished through the use of access codes (security codes, user codes) that limit individuals' access to those areas they are authorized to enter or use. Authorization is required for access to the:
1. Biorepository
2. Specimens, aliquots and any extracts thereof
3. Participant/client and study records

Access codes/user codes must be maintained and current (e.g. inactivated when employment of an authorized individual's employment ends).

**GEN.84100 Adequate Space Phase II**

The general biorepository has adequate, conveniently located space so the quality of work, safety of personnel, and patient care services are not compromised.

**REFERENCES**

**GEN.84200 Adequate Space Phase I**

All of the following areas have sufficient space and are located so there is no hindrance to the work.

1. Biorepository director
2. Staff pathologists and researchers
3. Biorepository technicians
4. Clerical staff
5. Chief technologist/biorepository manager
6. Section supervisors
7. Freezer storage area
8. Ambient temperature storage
9. Lavatories
10. Library, conference and meeting room
11. Personnel lounge and lockers

**GEN.84300 Climate Control Phase I**

The room temperature and humidity are adequately controlled in all seasons.

**Evidence of Compliance:**
✓ Temperature and humidity records, if specific ranges are required for instrument and/or reagent use

**GEN.84400 HVAC Phase I**

HVAC units, if present, are properly serviced and functioning to maintain appropriate compressor activity.

**Evidence of Compliance:**
✓ Records of maintenance

**GEN.84500 Hallway Obstructions Phase II**

Passageways are unobstructed.

**GEN.84600 Environment Maintenance Phase I**

Floors, walls and ceilings are clean and well-maintained.
GEN.84700  Environment Maintenance  Phase I

Bench tops, cupboards, drawers and sinks are clean and well-maintained.

GEN.84800  Environment Maintenance  Phase II

There are oxygen sensors or sufficient airflow to prevent asphyxiation in areas where liquid nitrogen is used.

GEN.85100  Inventory Control  Phase I

There is an effective supply inventory control system in operation.

NOTE: An effective inventory control system minimizes emergency requisitions and shortages of supplies.

Evidence of Compliance:
✓ A written procedure detailing relevant personnel, when to order supplies and levels of buffer stock required

REFERENCES
1) Chapman J. Saving money with computerized materials management. Advance/Lab. 1999;8(9):16-18

GEN.85200  Intrabiorepository Storage  Phase I

The intrabiorepository storage area is sufficient and free of clutter.

**REVISED** 07/28/2015

GEN.85300  Emergency Power  Phase II

Emergency power is adequate for the functioning of the biorepository.

NOTE: Emergency power supply must be adequate for refrigerators, freezers, incubators, etc., to ensure preservation of specimens.

**REVISED** 07/28/2015

GEN.85400  Emergency Power Load Testing  Phase II

Load testing is performed to ensure that emergency power is adequate for the functioning of the biorepository.

NOTE: Emergency power supply must be adequate for refrigerators, freezers, incubators, etc. to ensure preservation of specimens.

**REVISED** 07/28/2015

GEN.85420  Electrical Grounding  Phase II

There are records that all instruments and appliances are checked for adequate grounding and current leakage before initial use, after repair or modification, and when a problem is suspected.

NOTE: Exceptions to these requirements are as follows:

1. Devices protected by an approved system of double insulation or its equivalent. Such devices must be distinctively marked.
2. Devices connected to wall receptacles or circuit breakers with ground-fault circuit interrupter (GFCI) protection built-in need not be checked for current leakage.
3. **Equipment operating at 240 v must be checked for ground integrity only**

Verification of electrical safety is required whenever the electrical/electronic systems of a powered device has been removed or altered.

In addition, the US Occupational Safety and Health Administration (OSHA) requires that power cords of portable electrical equipment be visually inspected for external defects whenever relocated. Grounding configurations may not be bypassed by, for example, an adapter that interrupts the continuity of the grounding. If manufacturer's recommendations are available, they must be followed.

**REFERENCES**

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**GEN.85500 Contingency Plans**

Contingency plans are in place in the event that the back-up generator is not operational and if there is not enough fuel present to operate the generator.

**Evidence of Compliance:**
- Written contingency plan **AND**
- Schedule of fuel deliveries

**SAFETY**

Requirements in this section cover the general safety program for the entire biorepository.
GENERAL SAFETY

Inspector Instructions:

- Sampling of safety policies and procedure
- Ergonomic evaluation
- Sampling of personnel safety education records

- Adequate emergency lighting
- Flammable and combustible liquids (properly stored)
- Emergency eyewash available and tested properly

- How are your biorepository’s safe work practices reviewed?
- Is there a specific example of an occupational injury or illness that required medical treatment? What steps were taken to address the incident?

- For any occupational injury or illness that required medical treatment, further evaluate leadership's responses, corrective actions, follow-up procedures, and additional measures taken to ensure safety in the workplace

GEN.85600  Safety Policy and Procedure Approval

The biorepository director or designee reviews and approves all changes to the safety policies and procedures before implementation.

**REVISED** 07/28/2015

GEN.85700  Safety Policy and Procedure Availability

There are records for the training of all laboratory workers in safety.

NOTE: A system to ensure that all personnel have read the policies and procedures is required and must form a portion of the orientation program for new personnel. Posting of specific warnings or hazards as appropriate is urged.

Evidence of Compliance:
- Records of personnel review of safety policies and procedures

REFERENCES

**REVISED** 07/28/2015

GEN.85800  Safe Work Practices Review

There are records of periodic review (at least annually) of safe work practices to reduce hazards.
NOTE: Review must include bloodborne hazard control and chemical hygiene. If the review identifies a problem, the biorepository must investigate the cause and consider if modifications are needed to safety policies and procedures to prevent reoccurrence of the problem or mitigate the potential risk.

Evidence of Compliance:
✓ Safety committee minutes OR records of regular safety inspections OR incident reports and statistics OR another method defined by the biorepository director

GEN.85820  Ergonomics  Phase II

There is a written ergonomics program to prevent musculoskeletal disorders (MSDs) in the workplace through prevention and engineering controls.

NOTE: The program may include training of employees about risk factors, identifying physical work activities or conditions of the job commonly associated with work-related MSDs, and recommendations for eliminating MSD hazards. Biorepository activity, workplace and equipment (e.g. chairs, workstations, computer keyboards, and displays) should be designed to reduce the risks of ergonomic distress disorders and accidents.

Evidence of Compliance:
✓ Records of ergonomic evaluation including recommendations for eliminating MSD hazards and appropriate corrective action based on assessment findings

REFERENCES
1)  U.S. Dept. of Labor, Occupational Safety and Health Administration. Ergonomic safety and health program management guideline. 54 Fed Register 3904 (1989), modified at 29CFR1910

**REVISED** 07/28/2015

GEN.85900  Accidents  Phase II

There are written policies and procedures for the reporting and recording of all accidents resulting in property damage or involving spillage of hazardous substances.

GEN.85920  Gas Cylinder Storage  Phase II

Compressed gas cylinders are secured to prevent accidental falling and damage to the valve or regulator.

GEN.85940  Flammable Gas Cylinders  Phase II

Flammable gas cylinders are stored properly.

NOTE: Proper storage practices include:
1. Storage in a separate, ventilated room or enclosure
2. Cylinders are positioned well away from open flame or other heat sources, not in corridors and not within exhaust canopies

REFERENCES

GEN.85960  Liquid Nitrogen and Dry Ice  Phase II

Adequate policies, procedures, and practices are in place for the use of liquid nitrogen and dry ice.

NOTE: Practices for the safe handling of liquid nitrogen and dry ice include:
1. The mandatory use of appropriate gloves, shielding of all skin and the use of a face shield when decanting or entering an open container of LN2
2. The mandatory use of insulated loose-fitting gloves, dry ice tongs or scoop, and safety goggles/glasses when handling dry ice
3. Storage and use of all containers of LN2 and dry ice only in well-ventilated areas
4. Availability of a Safety Data Sheet

REFERENCES

**REVISED** 07/28/2015

GEN.86000 Occupational Injuries Phase II

There are written policies and procedures for the reporting of all occupational injuries or illnesses that require medical treatment (except first aid).

NOTE: For US facilities subject to OSHA regulations, all workplace fatalities must be reported to the Occupational Safety and Health Administration (OSHA) within eight hours and work-related in-patient hospitalizations or losses of an eye within 24 hours.

REFERENCES

GEN.86100 Occupational Injury Evaluation Phase II

An evaluation of these reports of biorepository accidents and occupational injury/illnesses is incorporated into the biorepository’s quality management program to avoid recurrence.

Evidence of Compliance:
✓ Records of report evaluation OR committee minutes with records of discussion

GEN.86120 Excessive Noise Phase II

The biorepository has a policy to protect personnel from excessive noise levels.

NOTE: The biorepository should provide protection against the effects of noise exposure when sound levels equal or exceed an eight-hour time-weighted average sound level of 85 decibels. The biorepository should monitor noise exposure if there is an indication that excessive noise levels are present (for example, when noise levels exceed 85 decibels, people have to shout to be heard).

REFERENCES

**REVISED** 07/28/2015

GEN.86130 Emergency Eyewash Phase II

The biorepository has adequate plumbed or self-contained emergency eyewash facilities in every area where there are hazardous chemicals as defined by the biorepository’s chemical hygiene plan (e.g. chemicals that are irritating, corrosive, toxic by contact or absorption). Testing records are maintained.

NOTE 1: The eyewash facility includes the following:
1. No greater than 10 seconds travel distance from areas in the biorepository where hazardous chemicals are present
2. Capable of delivering 1.5 L per minute for 15 minutes
3. Flow is provided to both eyes simultaneously
4. Nozzles or covers to protect from airborne contaminants
5. Hands-free flow once activated
6. Signage for location of eyewash
7. Unobstructed path with unlocked doors opening in the direction of the eyewash
8. Plumbed systems are protected from unauthorized shut off
9. Tepid fluid temperature (water temperature should be between 15°C and 37°C (60-100°F). Actual temperature recording is not required)
10. Plumbed systems are activated weekly
11. Self-contained units are visually examined weekly

NOTE 2: The manufacturer's specifications should be available.

REFERENCES

**REVISED** 07/28/2015
GEN.86140 UV Light Exposure  Phase II

There are written policies and procedures to prevent or reduce ultraviolet light exposure from instrument sources.

NOTE: UV light may cause corneal or skin burns from direct or deflected light sources. Wherever UV light sources are used, suitable and adequate personal protective equipment must be provided, and appropriate approved signage displayed. Laboratories may obtain information on safety from manufacturers of devices that emit UV light.

A suggested sign for display is: Warning: This device produces potentially harmful ultraviolet (UV) light. Protect eyes and skin from exposure.

Evidence of Compliance:
✓ Warning signage on source equipment AND
✓ Suitable PPE available, as required

**REVISED** 07/28/2015
GEN.86200 Disaster Preparedness  Phase II

There are written policies and procedures defining the role and responsibilities of the biorepository in internal and external disaster preparedness.

REFERENCES

**REVISED** 07/28/2015
GEN.86300 Evacuation Plan  Phase II

There is a written comprehensive and workable evacuation plan specific for the facility.

NOTE: 1) This plan must cover all employees and visitors, and must address the special needs of persons with disabilities. Evacuation routes must be clearly marked (Posting evacuation routes is optional). 2) Emergency lighting is adequate for safe evacuation of the biorepository.

REFERENCES
1) Occupational Safety and Health Administration. Exit routes, emergency action plans, and fire prevention plans: standard, 2002 [29CFR1910.38]
BIOLOGICAL SAFETY

Inspector Instructions:

- Sampling of safety and waste disposal policies and procedures
- Sampling of sterilizing device monitoring records
- Sampling of records of hepatitis B vaccination or records declining the vaccination

- PPE usage
- Biohazard disposal bins

- What has your facility done to reduce or eliminate exposure to bloodborne pathogens?
- How does your biobank dispose of sharps?

GEN.86400 Bloodborne Pathogens

The biorepository has written policies and procedures for infection control that comply with the OSHA Standard on occupational exposure to bloodborne pathogens and to the institution's exposure control plan.

NOTE: Universal or standard precautions must be used when handling all blood and body fluid specimens. The term "universal precautions" refers to a concept of bloodborne disease control requiring all human blood and other potentially infectious materials to be treated as if infectious for HIV, HBV, HCV or other bloodborne pathogens, regardless of the perceived "low risk" status of a participant or participant population. Alternative concepts in infection control are called Body Substance Isolation (BSI) and Standard Precautions. These latter terms define all body fluids and substances as infectious. All health care workers must routinely use appropriate barrier precautions to prevent skin and mucous membrane exposure when contact with blood or other body fluids is anticipated. Policies must comply with the OSHA Standard on Bloodborne Pathogens.

Evidence of Compliance:
✓ Safety manual AND
✓ Records of universal precaution training for all personnel expected to have contact with body fluids

REFERENCES

**REVISED** 08/17/2016
GEN.86500 PPE Provision and Usage

Appropriate personal protective equipment (gloves, gowns, masks and eye protectors, etc.) is provided and maintained in a sanitary and reliable condition in all technical work
areas in which blood and body substances are handled and in circumstances during which exposure is likely to occur.

NOTE: 1) Appropriate personal protective equipment (PPE) are items that do not permit blood or other potentially infectious materials to pass through to the skin or reach the employee’s work clothes, skin, footwear, etc. In addition to fluid-resistant gowns, aprons may be required if exposure to large volumes of body fluids is anticipated. 2) OSHA requires gloves to be worn with each participant or subject contact and changed after contact when performing vascular access procedures. Hands must be cleaned after glove removal using an effective antimicrobial method.

REFERENCES

**REVISED** 08/17/2016
GEN.86600  PPE Instruction  Phase II

Personnel are instructed in the proper use of personal protective clothing/equipment (e.g. gloves, gowns, masks, eye protectors, footwear).

NOTE: The required elements of training in the use of gloves include (a) Proper fitting of gloves; (b) Replacing gloves immediately when torn or contaminated; (c) Not washing or disinfecting gloves for reuse; (d) Using hypoallergenic gloves when indicated by patient or health care provider history; (e) Decontamination of hands after glove removal using an effective antimicrobial method.

Evidence of Compliance:
✓ Written policy for the use of PPE for specific tasks AND
✓ Records of personnel training for PPE

REFERENCES

GEN.86620  Latex Allergy  Phase II

The biorepository has a written program to protect personnel and participants/clients from allergic reactions from exposures to natural rubber latex in gloves and other products.

NOTE: The latex program should address at least the following elements.

1. Selection of products and implementation of work practices that reduce the risk of allergic reactions. If latex gloves are used, the employer should provide reduced protein, powder-free gloves to protect workers from infectious materials.
2. Provision of education programs and training materials about latex allergy
3. Evaluation of current prevention and control strategies whenever a worker is diagnosed with latex allergy

Evidence of Compliance:
✓ Records of employee education/training on latex allergies AND
✓ Records of evaluation of the plan, when appropriate
Laboratory General Checklist

GEN.86630  Manual Manipulation of Needles  Phase II

There is a written policy that prohibits the recapping, purposeful bending, breaking, removing from disposable syringes, or other manual manipulations of needles.

NOTE: Resheathing instruments or self-sheathing needles may be used to prevent recapping of needles by hand.

REFERENCES

GEN.86640  Sharps Disposal  Phase II

Sterile syringes, needles, lancets, or other blood-letting devices (“sharps”) that are capable of transmitting infection are used once only, and all waste sharps are discarded in puncture-resistant containers that are easily accessible, located in areas where needles are commonly used, and properly labeled to warn handlers of the potential hazard.

NOTE: Under US law, shearing or breaking of contaminated sharps is prohibited. Bending, recapping, or removing contaminated needles is prohibited as a general practice. Needles are expected to be used and immediately discarded, un-recapped, into accessible sharps containers.

REFERENCES

GEN.86650  Eating/Mouth Pipetting  Phase II

There is a written policy that prohibits smoking, eating, drinking, application of cosmetics and lip balm, manipulation of contact lenses, and mouth pipetting in all technical work areas.

NOTE: The biorepository must define the technical work area in particular when there is space sharing.

REFERENCES

GEN.86700  Specimen Transport Procedures  Phase II

There are written procedures for the procurement, transportation, and handling of biospecimens (e.g. blood, body fluids, tissue) to ensure that all specimens are submitted in an appropriately labeled and well-constructed container with a secure lid to prevent leakage during transport.

NOTE: Specimens sent through pneumatic tube systems must be sealed in fluid-tight bags. If pneumatic tube systems are used for transporting specimens, the biorepository must
have procedures to respond to a spill within the tube, including appropriate decontamination measures.

REFERENCES

GEN.86800  Spill Handling  Phase II
There are written procedures for handling spills of blood and other body fluids.

GEN.86900  Hepatitis B Vaccinations  Phase II
Personnel reasonably expected to have direct contact with body fluids are identified and offered hepatitis B vaccinations free of charge.

Evidence of Compliance:
✓ Written policy offering the hepatitis B vaccination to employees AND
✓ Records of vaccination OR records signed by employees declining the vaccine

REFERENCES

**REVISED** 07/28/2015
GEN.87000  Viral Exposure  Phase II
There is a policy for post-exposure follow-up after possible and known percutaneous, mucous membrane or abraded skin exposure to HIV, HBV or HCV that includes the following elements:

1. HIV, HBV and HCV testing of the source subject after consent is obtained
2. Appropriate clinical and serologic evaluation of the health-care worker
3. Consideration of appropriate prophylaxis for personnel acutely exposed to HIV, HBV or HCV, based upon medical indications, the serologic status and the informed consent of the health-care worker
4. Reporting of the exposure as required by law

Evidence of Compliance:
✓ Records of exposure follow-up

REFERENCES

**REVISED** 08/17/2016
GEN.87020  Biohazard Disposal  Phase II
All infectious wastes (e.g. glassware, blood collection tubes, microbiologic and tissue specimens) and other contaminated materials are discarded into "biohazard"-labeled containers that do not leak and have solid, tight-fitting covers that are applied before transport from the work area for storage and disposal.

NOTE: Waste disposal must be in accord with all regulations and disposed of with minimum danger to professional, technical, and custodial personnel.
All infectious wastes must be incinerated or appropriately decontaminated before being sent to a sanitary landfill.

Evidence of Compliance:
✓ Written procedure for waste disposal in accordance with local regulations

REFERENCES

GEN.87100 TB Exposure Plan Phase II

The biorepository has a written tuberculosis exposure control plan.

NOTE: This plan must include an exposure determination at defined intervals for all employees who may have occupational exposure to tuberculosis. Additional elements of the plan include engineering and work practice controls for hazardous activities that potentially may aerosolize Mycobacterium tuberculosis. Such activities include the handling of unfixed tissues in surgical pathology or autopsies.

If respiratory protection is needed because of potential exposure to an infectious agent by aerosol or droplet, personnel must use either a properly fit-tested NIOSH-approved filter respirator (N-95 or higher) or a powered air-purifying respirator (PAPRS) equipped with high efficiency particulate air (HEPA) filters. Accurate fit testing is a key component of effective respirator use.

REFERENCES

GEN.87125 Sterilizing Device Monitoring Phase II

All sterilizing devices are monitored periodically with a biologic indicator (or chemical equivalent) for effectiveness of sterility under conditions that simulate actual use.

NOTE: Each sterilizing device must be monitored periodically with a biologic indicator to measure the effectiveness of sterility. Chemical indicators that reflect sporidical conditions may be used. The test must be performed under conditions that simulate actual use. One recommended method is to wrap the Bacillus stearothermophilus spore indicator strip in packaging identical to that used for a production run, and to include the test package with an actual sterilization activity. Weekly monitoring is recommended.

Evidence of Compliance:
✓ Written procedure for monitoring sterilizing devices AND
✓ Records of monitoring at defined frequency

FIRE SAFETY

With respect to fire safety, if a checklist requirement conflicts with regulations of the Authority Having Jurisdiction (i.e. state and local fire codes), the regulations of the Authority Having Jurisdiction take precedence.
Inspector Instructions:

- Sampling of safety policies and procedures
- Sampling of employee fire safety training records
- Automatic fire extinguisher systems, if required
- Two exit access doors, if required
- Audible automatic fire detection and alarm system
- Fire alarm station
- Portable fire extinguishers, where appropriate

GEN.87200 Fire Prevention Policies and Procedures

**Policies and procedures are written and adequate for fire prevention and control.**

**NOTE:** Fire safety plans must include the use of alarms, response to alarms, isolation of the fire, evacuation of the area, extinguishment of the fire, and the responsibilities of biorepository personnel for those elements.

**REFERENCES**


GEN.87420 Fire Separation

**If the biorepository stores flammable materials, it is properly separated from inpatient areas and/or provided with automatic fire extinguishing (AFE) systems.**

**NOTE:** For those facilities with no inpatients, no AFE is required.

Where the biorepository is separated by two-hour construction (rated at 1.5 hours) and Class B self-closing doors (SCD), no AFE system is required. This applies to biorepositories that reside within a hospital or patient-care facility. An AFE system is required for those biorepositories separated from inpatient areas by one-hour construction and Class C SCD if flammable and combustible liquids are stored in bulk. An AFE system is always required if there are unattended biorepository operations employing flammable or combustible reagents. "Stored in bulk" means more than two gallons (7.5 L) of Class I, II, and IIIA liquids in safety cabinets and safety cans per 100 ft² (9.2m²), or half that amount if not in safety containers. The following are the definitions of these Classes:

- **Class I flammable:** any liquid that has a closed-cup flash point below 37.8°C and a Reid vapor pressure not exceeding 2068.6 mm Hg at 37.8°C as determined by ASTM D 323
- **Class II combustible:** any liquid that has a flash point at or above 37.8°C and below 60°C
- **Class IIIA combustible:** any liquid that has a flash point at or above 60°C but below 93°C

**REFERENCES**


GEN.87430 Fire Exit

**Each room larger than 1000 ft² (92.9m²), or in which major fire hazards exist, has at least two exit access doors remote from each other, one of which opens directly into an exit route.**
REFERENCES

GEN.87440 Fire Safety Training

Fire safety training is performed for new employees, with fire safety review conducted at least annually.

NOTE: Fire safety training must be recorded for all employees to show that they have been instructed on use and response to fire alarms and to execute duties as outlined in the fire safety plan. While fire exit drills are not required, physical evaluation of the escape routes must be performed annually, to ensure that fire exit corridors and stairwells are clear and that all fire exit doors open properly (i.e., not rusted shut, blocked or locked). Paper or computerized testing of an individual’s fire safety knowledge on the fire safety plan is acceptable; all personnel must participate at least once a year.

Evidence of Compliance:
✓ Records of participation for all employees in fire safety plan review at least annually (e.g. employee roster with dates of participation, sign-in sheet, etc.)

GEN.87442 Fire Detection/Alarm

There is an automatic fire detection and alarm system.

NOTE: 1) The system must connect to the facility’s overall system, where such a system exists. It must sound an immediate alarm in the event of smoke or fire. 2) The fire alarm is audible in all parts of the biorepository, including storage areas and lavatories. 3) Facilities employing hearing-impaired persons must have other means to alert these individuals, such as a visual alarm system.

REFERENCES

GEN.87444 Fire Alarm Station

There is a fire alarm station in or near the biorepository.

NOTE: Alarm stations must be visible, unobstructed, and accessible.

REFERENCES

GEN.87450 Fire Extinguishers

Appropriate portable fire extinguishers are provided for all areas in which flammable and combustible liquids are stored or handled.

NOTE: If gallon bottles of such materials are used, the minimum rating for Class B extinguishers is 10-B or higher. These are best located near or outside of doors leading to the area having solvent fire hazards.

REFERENCES
Laboratory General Checklist 08.17.2016

CHEMICAL SAFETY

Inspector Instructions:

- Sampling of chemical safety policies and procedures
- Sampling of SDS (formerly MSDS) sheets
- Sampling of formaldehyde vapor monitoring records
- Sampling of chemical waste disposal policies and procedures

- Acids and bases (properly stored)
- Sampling of hazardous chemicals (labeling)
- PPE usage

- How does your biobank dispose of hazardous chemicals?

**REVISED** 07/28/2015

GEN.87600 Chemical Hygiene Plan Phase II

The biorepository has a Chemical Hygiene Plan (CHP) that defines the safety policies and procedures for all chemicals used in the biorepository.

NOTE 1: The biorepository director or designee must ensure that the biorepository has a written chemical hygiene plan (CHP) that defines the safety policies and procedures for all chemicals used in the biorepository. The plan must include evaluation of carcinogenic potential, reproductive toxicity, and acute toxicity. The plan must include specific handling requirements for all hazardous chemicals used in the biorepository.

The purpose of the OSHA regulations is to ensure that the hazards of all chemicals are evaluated, and that information concerning their hazards is transmitted to employers and employees. This transmittal of information is to be accomplished by means of comprehensive hazard communication programs, which are to include container labeling and other forms of warning, safety data sheets and employee training. An acceptable CHP contains the following elements.

1. Responsibilities of the biorepository director and supervisors
2. Designation of a chemical hygiene officer
3. Policies for all operations that involve chemicals
4. Criteria for the use of personal protective equipment and control devices
5. Criteria for exposure monitoring when permissible levels are exceeded
6. Provisions for medical consultations and examinations
7. Provision for training anyone working in the biorepository on the elements of the CHP

8. A copy of the OSHA Laboratory Standard
9. Evaluation of the carcinogenic potential, reproductive toxicity and acute toxicity for all chemicals used in the biorepository. The product label, safety data sheets (SDS), or for chemicals purchased prior to June 1, 2015 with no appropriate SDS, records of investigation by the safety officer may be used for this evaluation.
10. Specific handling requirements for all hazardous chemicals used in the biorepository

**NOTE 2:** Chemicals that must be handled as potential carcinogens include those defined by OSHA as "select carcinogens." OSHA defines select carcinogens as any substance that is:

1. Regulated as a carcinogen by OSHA, has been classified as "known to be carcinogenic" by the NTP, or listed as a group I carcinogen by the IARC
2. Has been classified as "reasonably anticipated to be carcinogenic" by the NTP or listed as a group 2A or 2B carcinogen by the IARC if it meets the toxicological criteria listed in the January 31, 1990 Fed Register, pages 3319-3320

OSHA also requires special containment procedures for substances that are reproductive toxins or are acutely hazardous.

Authoritative sources include (but are not limited to) OSHA (Code of Federal Regulations. Title 29, Part 1910.1200 and 1450); NIOSH (Registry of Toxic Effects of Chemical Substances); the National Toxicology Program; the International Agency for Research on Cancer, and Safety Data Sheets.

**Evidence of Compliance:**
- ✓ Written evaluation of chemicals used in the biorepository for carcinogenic potential, reproductive toxicity, and acute toxicity **AND**
- ✓ Written procedure for chemical fume hood function verification **AND**
- ✓ Records of testing

**REFERENCES**
3) Karcher RE. Is your chemical hygiene plan OSHA-proof? Med Lab Observ. 1993(Jul);29-36

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**GEN.87700 Chemical Safety Document Access**  
**Phase II**

For US biorepositories, employees have access to all of the following documents.

1. Current Safety Data Sheets (formerly MSDS) and other references that list the details of hazards and the precautions for safe handling and storage
2. Chemical Hygiene Plan of the biorepository

**NOTE:** It is acceptable for SDS information to be electronically available to users, rather than in book format; there is no requirement for paper-based information. Indeed, electronic manuals have the advantage of more accurately reflecting current requirements. The central point is immediate availability to all personnel at all times.

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**GEN.87800 Chemical Precautionary Labels**  
**Phase II**

Precautionary labels are present on the containers of all hazardous chemicals, indicating type of hazard and what to do if accidental contact occurs.

**NOTE:** The biorepository may use signs, placards, process sheets, batch tickets, operating procedures, or other such written materials in lieu of affixing labels to individual stationary process containers, as long as the alternative method identifies the containers to which it is applicable and conveys the information otherwise required to be on a label. The written materials shall be readily accessible to the employees in their work area throughout each work shift. It is not required to label portable containers into which hazardous chemicals are transferred from labeled containers, and which are intended only for the immediate use of the employee who
**PPE And Hazardous Materials**

Personnel use the proper personal protective devices when handling corrosive, flammable, biohazardous, and carcinogenic substances.

**NOTE:** Such devices may include gloves of appropriate composition, aprons, and eye protection. Shoes or shoe covers must protect the entire foot in areas where splashing is expected.

**REFERENCES**


**Chemical Hazard Emergencies**

Explicit instructions are posted, and appropriate supplies available, for the emergency treatment of chemical splashes and injuries and the control of chemical spills wherever major chemical hazards exist.

**NOTE:** Spill kits must be handled in accordance with manufacturer's instructions. If no expiration date is assigned, the spill kit must indicate the date it was put into service and the director must periodically assess its usability.

**REFERENCES**


**Hazardous Chemical Waste Disposal**

Written policies and procedures are adequate for hazardous chemical waste disposal.

**NOTE:** 1) The biorepository is responsible for all real or potential hazards of wastes at all stages of disposal including transportation and final disposition. 2) The method for the disposal of all solid and liquid wastes is in compliance with local, state and federal regulations. (Whether or not biorepository management is responsible for waste disposal, the biorepository should have documentation that the facility is in compliance with all applicable regulations. Prevailing local, state and federal (EPA) regulations should be reviewed by the biorepository director, safety officer or facilities manager to ensure that the biorepository is in compliance with regulations.)

**Evidence of Compliance:**

✓ Records of review of regulations for compliance

**REFERENCES**


**Formaldehyde and Xylene Safety**

Formaldehyde and xylene vapor concentrations are maintained below the following maxima, expressed as parts per million, in all areas of the biorepository where formaldehyde or xylene are used.
NOTE: Formaldehyde and xylene vapor concentrations must be monitored in all areas where these reagents are used: e.g. surgical pathology gross dissection room, histology laboratory, etc. Initial monitoring involves identifying all employees who may be exposed at or above the action level or at or above the STEL and accurately determining the exposure of each employee identified. Further formaldehyde monitoring is mandated at least every six months if results of the initial monitoring equal or exceed 0.5 ppm (8 hr time-weighted exposure, the “action level”) or at least once per year if the results exceed the short term exposure limit (STEL) 2.0 ppm. The laboratory may discontinue periodic formaldehyde monitoring if results from two consecutive sampling periods taken at least seven days apart show that employee exposure is below the action level and the short-term exposure limit, and 1) no change has occurred in production, equipment, process or personnel or control measures that may result in new or additional exposure to formaldehyde, and 2) there have been no reports of conditions that may be associated with formaldehyde exposure.

Formaldehyde monitoring must be repeated any time there is a change in production, equipment, process, personnel, or control measures which may result in new or additional exposure to formaldehyde for any employee involved in the activity. If any personnel report signs or symptoms of respiratory or dermal conditions associated with formaldehyde exposure, the laboratory must promptly monitor the affected person’s exposure.

Xylene must be monitored initially, but there is no requirement for periodic monitoring of xylene. Repeat monitoring should be considered when there is a change in production, equipment, process, personnel, or control measures likely to increase exposure levels.

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<th>15 min Short-Term Average Exposure Limit (STEL) in ppm</th>
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<tr>
<td>Xylene</td>
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<td>150</td>
</tr>
</tbody>
</table>

Evidence of Compliance:

✓ Written procedure for formalin and xylene safety including action limits, criteria for discontinuation of monitoring and criteria for resumption of monitoring AND
✓ Record of initial formalin and xylene monitoring and repeat monitoring when indicated AND
✓ Records of corrective action when exposure limits are exceeded

REFERENCES
4) Occupational Safety and Health Administration. 29CFR1910.1048 and 1450, revised July 1, 1998

GEN.88100 Flammable Storage Phase II

Supplies of flammable and combustible liquids are reasonable for the biorepository’s needs, and are properly stored.

NOTE: 1) In each biorepository area, up to one gallon (3.7 L) of Class I, II and IIIA liquids may be stored outside of fire-resistant cabinets for each 100 ft² (9.2m²) of space defined by fire-resistant walls/doors. Up to two gallons (7.5 L) of Class I, II, and IIIA liquids may be stored in safety cans and safety cabinets for each 100 ft² (9.2m²). These amounts may be doubled if there is an automatic fire suppression system (e.g. sprinklers). For example: a 1000 ft² (92.9m²) laboratory defined by fire resistant walls/doors can store 10 gallons (37.7 L) outside a safety cabinet and 20 gallons (75.7 L) inside a safety cabinet and double those numbers if there is an automatic fire suppression system. 2) Safety cans should be used for bulk storage of flammable and combustible liquid (National Fire Protection Association classes I and II). Metal or DOT-
approved plastic containers provide an intermediate level of hazard containment between glass and safety cans. One pint (0.4 L) of a highly volatile solvent such as isopentane, stored in glass has about the same ignitability risk as two gallons (7.5 L) stored in safety cans. Safety cans should be used instead of glass bottles if the purity required does not mandate glass storage.

REFERENCES

GEN.88200 Volatile Solvent Ventilation Phase II
Storage areas and/or rooms where volatile solvents are used are adequately ventilated.

NOTE: Areas where flammable liquids are used must be ventilated for protection of employee health, as well as fire prevention. Areas where flammable liquids are stored should be ventilated primarily for fire protection. Storage cabinets do not need to be vented, but if they are vented the duct system must be explosion proof.

REFERENCES

**REVISED** 07/28/2015
GEN.88300 Acid/Base Storage Phase II
Supplies of concentrated acids and bases are stored in cabinets near floor level.

NOTE: 1) Strong acids and bases must not be stored under sinks, where contamination by moisture may occur. 2) Storage containers of acids and bases should be adequately separated to prevent a chemical reaction in the event of an accident/spill/leak. 3) Bottle carriers are used to transport all glass containers larger than 500 mL that contain hazardous chemicals.

GEN.88310 Evacuation/Clean-up Plan Phase II
The biorepository has a plan for evacuation and clean-up in the event of an LN2 or liquid CO₂ spill from a bulk source.

GEN.88325 Emergency Treatment - Toxic Fumes Phase II
The biorepository has a plan for the immediate treatment of an individual overcome by toxic fumes.

RADIATION SAFETY

GEN.88340 Radiation Safety Manual Phase II
If the biorepository handles specimens that are known to be radioactive, there are written policies and procedures adequate for radiation safety.

GEN.88350 Radioactive Material Handling Phase II
If the biorepository handles specimens that are known to be radioactive, there are specific policies and procedures for the safe handling of tissues that may contain radioactive material (e.g. sentinel lymph nodes, breast biopsies, prostate "seeds", etc.).

NOTE: These policies and procedures should be developed in conjunction with the institutional radiation safety officer, and must comply with any state regulations for the safe handling of
tissues containing radionuclides. The policies and procedures should distinguish between low radioactivity specimens such as sentinel lymphadenectomy and implant devices with higher radiation levels.

REFERENCES