1 INTRODUCTION

The purpose of the Crime Laboratory Division (CLD) Quality Operations Manual is to provide CLD staff with written policies and procedures that:

- Promote an efficient and effective operation within the CLD;
- Assist CLD staff in performing assigned duties and tasks;
- Ensure that the work product of the laboratory is of the highest quality possible;
- Help to demonstrate that the CLD operates a quality management system, is technically competent, and is able to generate technically valid results.

This manual also describes the Quality Assurance Program of the CLD and provides personnel with a description of the Division’s policies for maintaining an effective quality assurance program. This program applies to all work done in all areas within the CLD and the policies and procedures are binding on all personnel of the CLD and shall be adhered to.

The official version of this manual is the electronic version on the Forensic Laboratory Services Bureau (FLSB) Portal, the Bureau’s SharePoint site.
2 SCOPE

2.1 MISSION STATEMENT

The Washington State Patrol (WSP) CLD will provide forensic science services and training for Washington’s criminal justice agencies. The CLD is committed to providing the highest quality forensic services which ultimately enhances public safety for the citizens of Washington. The WSP CLD adheres to the WSP Mission Statement and CLD Mission Statement.

The Washington State Patrol makes a difference every day, enhancing the safety and security of our state by providing the best in public safety services.

The Crime Laboratory Division enhances public safety by providing high-quality forensic science services and forensic evidence training for all criminal justice partner agencies within the State of Washington.

2.2 VISION

Our vision is to be recognized as an international leader in forensic science in quality, service, and innovation.

2.3 VALUES

Every employee is a critical member of a team committed to professional excellence through:

- Unbiased and fair expertise
- Integrity
- Accountability
- Commitment to excellence
- Selfless service
- Continuous Improvement

2.4 GOALS AND OBJECTIVES

The goals and objectives of the CLD will be reviewed continually and are based upon the needs of the Criminal Justice System and the needs of the customers served by the respective laboratories.

2.5 LEGAL MANDATE

The Washington State Patrol Crime Laboratory Division is a publicly funded, legal entity and is responsible for its legislatively mandated actions. The CLD provides scientific and technical examinations for all criminal justice agencies as mandated
by the Revised Code of Washington (RCW) 43.43.670. The CLD also provides training assistance to the law enforcement agencies of Washington State.

3 DEFINITIONS

Additional definitions for more specific terms related mostly to a particular chapter are in the respective chapters. The following words when used in this document indicate addressing in writing: agreed, appoint, authorize, define, instructions, method, plan, procedure, program, record, schedule, and specify.

1. **Accuracy**
The ability of a measurement result to report the true or target value of the property being measured; the degree of conformity or nearness of a measurement to a standard or a true value.

2. **Accreditation Cycle**
The period of time between full on-site assessments by the accrediting body, generally a period of approximately four (4) years.

3. **Administrative Documentation**
Documentation either received or generated by the laboratory that does not constitute data or information resulting from testing. Administrative documentation includes records such as case related conversations, evidence receipts, description of evidence packaging and seals, printouts of thumbnail images, investigative reports and other pertinent information.

4. **Administrative Review**
Final review for non-technical matters of the case file and final report prior to release of the report to the customer.

5. **Amended Report**
A report generated to correct errors, make additions to, or improve wording in a previous laboratory report.

6. **ANAB**
American National Standards Institute (ANSI) {A} National Accreditation Board {NAB}

7. **Annual**
Annual in this manual refers to the calendar year unless otherwise specified.

8. **Calibration**
The process used to establish that equipment is capable of achieving the discipline’s and the manufacturer’s specifications for the test and by which known traceable standards having unbiased reference values are introduced into an item of equipment. The
equipment is then adjusted (either by software, hardware, electronics, etc.) to report the known reference value.

9. Casework
Analytical work performed by a forensic scientist or technician. Generally this involves the examination and analysis of physical evidence, but is extended to include the processing and analysis of convicted offender DNA samples by scientists in the CODIS laboratory, and the entry and correlation of ammunition components in the National Integrated Ballistic Information Network (NIBIN).

10. Case File
Administrative and examination documentation pertaining to a case that is received or generated by the laboratory.

11. Case Record
A case record, or test record, is all administrative and examination documentation pertaining to a case that is received or generated by the laboratory. Information in the case record may be in the case file or in other locations in the laboratory which are designated as extensions of the case file.

This may include, but is not limited to:

- Administrative and examination documentation maintained in the case file
- Electronically stored data
- Chain of custody documentation (including Laboratory Information Management System (LIMS))
- Digital images
- Instrument maintenance and verification documentation
- Reagent and standard quality control documentation
- Submitting agency/customer information, including name and address

Information in the case record may be in the case file or in other locations in the laboratory which are designated as extensions of the case file.

12. Certified Reference Material (CRM)
A material or substance, accompanied by a certificate, one or more of whose property values are certified by a procedure that establishes traceability to an accurate realization of the unit in which the property values are expressed. Each certified value is accompanied by an uncertainty at a stated level of confidence. An example of such a CRM would be a NIST traceable ruler.

13. Competency Test
A written, oral, and/or practical test or series of tests designed to evaluate an individual’s knowledge, skills, and/or ability to establish:
• that an individual has demonstrated achievement of technical skills, and;
• met minimum standards of knowledge necessary to perform forensic testing.

14. Consensus Standards
Consensus standards are voluntary standards that are accepted and agreed upon within an industry by bodies such as American Society for Testing and Materials (ASTM).

15. Correction
Immediate action taken to eliminate a detected nonconformity (a correction can be made in conjunction with a corrective action).

16. Corrective Action
The overall actions taken and plan or process used to address a nonconformity and to eliminate the cause of a detected nonconformity or other undesirable situation.

17. Corrective Action Plan (CAP)
A formal statement by the supervisor or designated authority prepared and entered using the QPIT, detailing the following:

• Description of the incident (what is the nature of the nonconformity?)
• Root cause analysis (including the chain of events leading to or causing the nonconformity)
• The immediate corrective actions taken (how was the problem handled?)
• Preventive action to avoid future occurrences
• A timeline for completion of the corrective and preventive actions
• A recommended schedule for follow-up to determine the effectiveness of the preventive measures to be taken (if required)

18. Corrective Action Report
A summary report in interoffice communication (IOC) format, or included in QPIT, prepared by the supervisor or designated authority upon the conclusion of a corrective action (if required).

19. Court Testimony Review
To oversee, evaluate or monitor testimony provided in a court of law under oath.

20. Division Operational Plan
The CLD Operational Plan is a document comprised of goals and objectives outlining the intended direction of the Crime Laboratory Division. This plan has the stated strategies (action items) detailing how the goals and objectives will be achieved.

21. Division Objectives
Objectives are general statements that address critical issues by breaking down goals into smaller, more specific pieces. Generally, there should be an objective assigned to
each critical issue within the Division. Objectives drive actions and represent the general end toward which the Division efforts are directed. An objective should provide a sense of the level of performance expected.

22. Division Strategies
Strategies state how the Division is going to accomplish the stated Division objectives. They include an action plan along with performance measures in order to determine to what level the objectives are achieved.

23. Draft Report
A preliminary version of a laboratory report prior to its issue under laboratory letterhead. The document is considered a work in progress until the technical review is completed. The final version is referred to as the Final Draft Report, which may be identified by the presence of both the author’s and technical reviewer’s initials, or signature, and date. (Note: The CODIS Laboratory does not issue reports.)

24. Equipment
Instruments or devices used to measure, record or identify any entity, and having a significant effect on the accuracy or validity of the result of the test, calibration or sampling, as determined by requirements in the functional area technical procedures: Also called test equipment.

25. Examination Documentation
Documentation of data or information resulting from testing, also referred to as case notes, and usually generated by the laboratory. Examples include reference to procedures followed, tests conducted, observations and results of examinations or tests, standards and controls used, diagrams, photographs, and printouts.; part of the technical record. (See 10.2.2 Examination Documentation)

26. External Assessment
A review conducted by personnel from outside the CLD which compares the various aspects of the laboratory’s performance against stated requirements, standards, policies and procedures.

27. Fully Documented
Documentation as to the source of the material and the date it was acquired (if known). Documentation may be made on the reference material itself, on its proximal packaging, or as part of a database record.

28. Internal Audit
A documented review process conducted by CLD personnel for obtaining objective evidence (records, statements of fact or other relevant information), and assessing them objectively to determine whether and the extent to which the laboratory fulfills the stated requirements, standards, policies and procedures.
29. **Internal Validation**
Validation that occurs within the CLD by qualified CLD personnel.

30. **Laboratory Developed Methods**
Methods developed in house as standard methods for a specific laboratory purpose.

31. **Laboratory Report**
The formal results of the requested analysis, issued under laboratory letterhead that is returned to the requestor.

32. **Management System Review (MSR)**
A Management System Review is an annual review by management of the laboratory management and quality systems, and its testing activities to ensure continuing suitability and effectiveness. The finding of this review will be used as a tool to introduce necessary changes or improvements by management.

33. **Method**
Any technical procedure detailing the use of reagents and/or instrumentation for scientific analyses, synonymous with “test procedure”.

34. **Modified Method**
Standard scientifically validated and forensically adopted method that is used outside the intended scope, or has been amplified or modified.

35. **Non-Standard Method**
A scientifically validated method or procedure that is not typically applied or used for forensic analysis.

36. **National/International Standard**
A standard recognized by national or international agreement to serve as the basis for assigning values to other standards of the quantity concerned. The standards which generally apply are the metric system of measures expressed in SI units, the units of the International System of Units.

37. **National Institute for Standards and Technology**
This federal agency, also known as NIST, is located within the Department of Commerce and represents the final authority for metrology in the United States. Ideally, all measurement results should be documented and shown to be traceable to NIST.

38. **Nonconformity of Test/Work (nonconformance)**
Non-fulfillment of a test/work requirement; failure to appropriately follow or apply accepted protocols in case work; any aspect of testing that does not agree with established laboratory, technical or quality system procedures or requirements.
Inefficiency in work or behavioral issues in work performance are not necessarily a nonconformity.

39. **Objective Evidence**  
Data supporting the existence or verity of something: may be obtained through observation, measurement, test or other means.

40. **Opinions and Interpretations**  
A formal expression of judgment based upon the analyst’s interpretations of observations or data.

41. **Performance Verification**  
A set of operations to determine if a piece of equipment or instrumentation is working correctly within manufacturer’s specifications or CLD specified parameters, or to determine if a validated method is fit for purpose and performing as expected.

42. **Precision**  
The degree of agreement or repeatability among replicate measurements performed at the same time, on the same instrument, by the same operator and under the same conditions. Precision is quantified by the standard deviation.

43. **Properly Controlled**  
Access to reference materials under the control of the laboratory is restricted to those persons authorized by the laboratory manager.

44. **Quality**  
Adherence to generally recognized standards of good laboratory practice.

45. **Quality Assurance (QA)**  
Those processes and systematic actions necessary to provide confidence that the laboratory’s work product and services will satisfy given requirements for quality.

46. **Quality Assurance Audit**  
A systematic examination and review to determine whether quality processes and related results comply with the CLD protocols, policies, and procedures, and whether these practices are suitable and effective in achieving the quality objectives.

47. **Quality Operations Manual**  
A collection of the CLD quality system policies and objectives and how these policies and objectives will be implemented.
48. **Quality Assurance Program (QAP)**
A planned system of activities describing requirements for forensic analyses and reporting, the purpose of which is to provide confidence that the work product and services provided by the CLD are scientifically sound and valid.

49. **Quality Control (QC)**
Internal activities or activities conducted according to externally established standards used by the CLD laboratories to consistently ensure accurate analytical results.

50. **Quality Management System**
The total organizational structure, responsibilities, policies, procedures, and resources for implementing quality management. This includes all activities which directly or indirectly contribute to quality.

51. **Quality Assurance Records**
Records, logs, worksheets and electronic files that provide documented support of conformity to the quality management system. These records include, but are not limited to, method and equipment validation documents, equipment verification records, reagent and chemical QC logs, training records, proficiency and competency test records, courtroom testimony monitoring records and audit records.

52. **Quality Process Improvement Tracker (QPIT)**
The electronic data system used to track quality process improvements, nonconforming work and corrective actions. (Formerly entitled: Remedy Nonconformance Tracking Program)

53. **Reference Material**
A stable material, substance, object or artifact of known properties or origin which has been established to be fit for its intended use in a measurement process, may be used in the determination of the properties or origins of unknown items, and which is maintained for identification, comparison or interpretation purposes.

54. **Reference Standard**
A measurement standard designated for the calibration of other measurement standards and for calibrating equipment, verifying performance and/or for use as a control in experiments.

55. **Request**
The list of services or analyses which the submitting agency has asked the laboratory to perform on evidence received in the laboratory, formally made by completing and submitting the Request for Laboratory Examination, Form 3000-210-005 (RFLE).
56. **Substantive Nonconformance**
An incident where the nature or cause of the nonconformity would call into question the quality of the testing, raises immediate concern and has a fundamental, significant impact on the work product of the laboratory or the integrity of evidence; or where there is concern that if the nonconformance continues for an extended period, the work product of the laboratory or integrity of evidence could be negatively affected. Examples include, but are not limited to, an error that may result in immediate suspension of the laboratory activity or staff from casework or applicable task by the appointing authority; or an error in an issued lab report that has a substantial likelihood of adversely affecting the customer or interested party.

57. **Supervised Casework**
Casework conducted by an analyst that is monitored by the trainer or other proficient and experienced scientist to ensure casework approach, documentation, procedures and methods are appropriately applied.

58. **Technical Procedures**
Scientific methodologies used in forensic analyses. Written procedures will be prepared for routine tests performed in the CLD Laboratories. The procedures used may be those developed and validated in–house or by an outside laboratory.

59. **Technical Record**
Examination documentation, laboratory reports, and amended reports.

60. **Technical Review**
Review of examination documentation, draft reports and testimony to ensure the validity of test results, opinions and interpretations and that proper procedures were used and documented.

61. **Testimony**
A statement made under oath as a witness.

62. **Traceability**
The property of a measurement result whereby it can be related to standard references, usually national or international, through an unbroken chain of comparisons all having stated uncertainties.

63. **Trainee**
A trainee is any employee of the CLD who is training in a new discipline, functional area or job classification. Trainees can be permanent, non-permanent, probationary or trial service. Please refer to the Collective Bargaining Agreement and HRD for clarification of employee status.
64. Training Procedures
The foundational training program required for all qualified forensic scientists in their discipline prior to assuming forensic analysis.

65. Uniquely Identified
An item is uniquely identified when each item or group of similar items has a unique name or identifier. This may be a laboratory generated number, database generated number or simply the name of the item if it is unique.

66. Validation
Confirmation, through the provision of objective evidence, that the particular requirements for a specific intended use or application have been fulfilled and is fit for purpose.

67. Verbal/Email Reports
Oral or emailed communication of technically reviewed analytical results given prior to completion of a formal written case report.

68. Verification
The procedure used to evaluate and confirm the validity of a test result/opinion reached by re-performing the comparison between the unknown and the known by a different person. Also: provision of objective evidence that a given item fulfills specified requirements.

69. Yearly
Within the calendar year, and no sooner than four months from the previous action.
4 QUALITY ASSURANCE PROGRAM

4.1 QUALITY MANAGEMENT SYSTEM AND ASSURANCE PROGRAM

4.1.1 POLICY
The CLD will establish, implement and maintain a quality management system appropriate to the scope of its activities. The CLD will document its policies, systems, programs, procedures and instructions to the extent necessary to assure the quality of the test results. The system’s documentation will be communicated to, understood by, available to, and implemented by the appropriate personnel. The CLD Quality Management System policies, procedures and objectives are defined in this Manual.

The laboratory management is committed to complying with ISO/IEC 17025 and any supplemental standards and the policies and procedures described herein and will proactively strive to continually improve the effectiveness of the Quality Management System.

The documentation upon which the Quality Management System is built includes this manual, all technical procedures and training manuals, safety manuals, DNA Quality Assurance and FBI Quality Assurance Standards manuals, and the Forensic Services Guide. The Quality Operations Manual has over-riding authority over all technical procedures manuals, including LIMS Policies and Procedures. The Washington State Patrol Regulation Manual has over-riding authority over all FLSB Manuals.

Laboratory managers and supervisors are responsible for ensuring that the policies and procedures adopted by the CLD are implemented and integrated into the daily operations of the laboratory. Laboratory managers are also responsible for overseeing and monitoring and ensuring conformance to the Quality Management System. In this respect, they function as quality assurance managers for their individual laboratories.

4.2 QUALITY POLICY STATEMENT

The CLD, its management, and its employees, are committed to professional excellence. All CLD employees will work to continually maintain the highest degree of quality and integrity of laboratory services and to ensure that forensic conclusions are scientifically sound and valid.

To this end, all laboratory analyses and related services performed by the laboratory system shall meet generally recognized standards of the forensic community and its accrediting organizations. Specifically, the CLD shall carry out all testing activities in accordance with stated methods, the requirements of the customer, the State of Washington and federal regulatory authorities and the ISO/IEC 17025:2017 standards (hereafter ISO) and any supplemental standards required by the accrediting
organization. National DNA Index System (NDIS) participating laboratories shall conform to requirements in the NDIS Operational Procedures Manual and in applicable FBI Quality Assurance Standards.

The CLD embraces and supports the principles presented in the ANAB Guiding Principles of Professional Responsibility for Forensic Service Providers and Forensic Personnel, which includes statements addressing Professionalism, Competency and Proficiency, and Clear Communications, and ensures appropriate actions are taken when necessary.

The CLD Quality Management System is designed to continually improve the quality and level of services provided to Washington’s Criminal Justice System and to assure the credibility of the CLD. This will be accomplished by providing quality service and audits through every area of the division. The CLD Quality Operations Manual is binding on all personnel of the Division and shall be adhered to.

All employees are required to familiarize themselves with the appropriate manuals and implement the CLD quality assurance policies and procedures in their work. In doing so, the CLD will maintain the highest level of staff expertise and analytical abilities, promote staff confidence, and conform to the ISO accreditation standards and any supplemental standards. DNA forensic scientists have additional quality requirements which can be found in the DNA Quality Manual and the FBI DNA Quality Assurance Standards document.

Laboratory management is committed to complying with ISO and any supplemental standards and the policies and procedures described herein and will proactively strive to continually improve the effectiveness of the Quality Management System.

The commitment to implement a successful Quality Management System begins with the CLD Commander and is supported by a commitment from the Standards and Accountability Section. The CLD Commander and the Standards and Accountability Section (SAS) Manager affirm their commitment to this quality policy statement, a signed copy of which is available on the FLSB Portal.

### 4.3 Quality Assurance Objectives

The objectives of the CLD overall quality assurance program are:

#### 4.3.1 Service

To provide quality service to Washington’s Criminal Justice System by competent personnel using procedures that are valid, reliable and sufficient for the intended purpose, and by meeting customer, statutory and regulatory requirements.
4.3.2 Standardization
To bring uniformity to the technical policies and procedures across the Division with the intent of producing high quality work that satisfies the customer’s requirements for service and meets the accrediting body’s standards.

4.3.3 Training and Education
- To provide training to CLD personnel in their area of expertise,
- To provide training to CLD personnel in the quality management system,
- To provide growth opportunities for CLD personnel,
- To facilitate involvement of CLD personnel at the national and international level,
- To provide continuing education to our customers regarding CLD services, policies and procedures.

4.3.4 Accreditation
To maintain laboratory quality, excellence and reliability by conforming to accreditation standards and maintaining accreditation requirements.

4.3.5 Review
To proactively review and monitor the Quality Management System to identify potential nonconformities to standards.

4.3.6 Audit
To assess and document quality assurance activities through an audit process in order to demonstrate conformance to accreditation standards and maintain trust in the work product.

4.3.7 Survey and Communication
- To facilitate and enhance communication within the organization.
- To conduct surveys and maintain open communication with customers.
- To review and analyze customer requirements and satisfaction with the CLD services.
- To communicate to all CLD personnel the importance of meeting customer requirements as well as statutory and regulatory requirements.

4.4 Adherence to the Quality Assurance Program
In order to adhere to the Quality Assurance Program and meet quality assurance objectives, the CLD will:
- Use established technical procedures in laboratory analyses that are reliable, reproducible, accepted in the forensic science community and adequate for the intended purpose.
• Participate in a proficiency testing program to monitor the routine operational performance of the laboratory.

• Provide sufficient training to all staff. Continuing education and professional career development training will be available to all staff as necessary to provide the best possible work product.

• Conduct regular court testimony monitoring of staff members.

• Have an employee performance evaluation program in which the tasks, responsibilities, safety and career development needs of the employee are reviewed each year.

• Have a system of technical and administrative review for case files and reports.

• Conduct annual internal quality assurance audits. The audits will be overseen and evaluated by the Standards and Accountability Section (SAS) to ensure that the CLD stated policies and procedures are being followed.

• Have an annual management system review to include a review of the quality assurance program and conduct periodic management meetings to discuss CLD policies and objectives. Information from these meetings will be communicated to all CLD staff through the FLSB Portal.
5 STRUCTURE, SERVICES AND FUNCTIONS

5.1 ORGANIZATION

The CLD is a part of the WSP FLSB. The CLD Headquarters, Combined DNA Index System (CODIS) Lab and Seattle Crime Lab are co-located in Seattle. The Division has laboratories located in Kennewick, Marysville, Olympia, Seattle, Spokane, Tacoma, and Vancouver, with laboratory managers reporting to the CLD Commander. The Standards and Accountability Section (SAS) is co-located in Seattle and reports to the CLD Commander. FSLB Headquarters is located in Olympia.

The CLD will provide services in scientific examination of physical evidence, collection and preservation of evidence, and expert testimony regarding the scientific examinations according to the legal mandate listed above. Training will be provided to law enforcement agencies within the state in the collection and preservation of evidence, and crime laboratory capabilities.

5.2 SERVICES

The CLD laboratories provide examination and casework services in the functional areas listed below, each responsible for testing in at least one accredited forensic discipline:

5.2.1 Materials Analysis

Scientists in this functional area identify seized drugs, pharmaceuticals, materials from clandestine drug laboratories, ignitable liquids and ignitable liquid residues in fire debris, explosives and post-blast explosive residues, selected poisons, chemical unknowns; perform quantitation of THC to identify marijuana and marijuana concentrates; examine and compare trace materials such as hairs, fibers, glass, paint, impression evidence, and other miscellaneous materials; and may provide consultation or assistance with clandestine laboratory investigations.

5.2.2 Firearms and Toolmarks

Scientists in this functional area examine and compare firearms, ammunition components, gunshot residues for distance determinations, and toolmarks. These scientists also reconstruct shooting scenes, restore obliterated serial numbers, and image fired ammunition components for inclusion in the Integrated Ballistics Identification System (IBIS)/ National Integrated Ballistics Information Network (NIBIN).

5.2.3 DNA

Scientists in this functional area characterize biological stains, extract and type human DNA from biological evidence, compare the DNA typing profiles to the DNA of known individuals and enter DNA profile into the CODIS database.
5.2.4 Combined DNA Index System (CODIS)
Scientists in the CODIS Laboratory type convicted offender samples for the CODIS database and manage the statewide CODIS database.

5.2.5 Questioned Documents
Scientists in this functional area examine and compare handwriting, hand printing, altered documents, indented writing, machine-generated documents, paper and ink.

5.2.6 Latent Prints
Scientists in this functional area examine and process evidence for friction ridge impressions, analyze and compare friction ridge impressions, and enter impressions into a variety of finger and palm print databases.

5.2.7 Crime Scene Response
Participating scientists are members of the Crime Scene Response Team (CSRT) and provide crime scene assistance to law enforcement agencies investigating major crimes. These scientists assist the agencies with evidence recognition and collection, scene documentation (including 3D scanning), bloodstain pattern interpretation, shooting incident reconstruction, and crime scene reconstruction.

CSRT full-time and part-time responders share the responsibility to produce timely reports and technical reviews and to testify in court. Part-time responders must also balance these needs with their regular case work and other assignments.

Part-time crime scene responders are expected to:
- Be on call for a week at a time on a predetermined rotation schedule.
- Dedicate their time during their on-call week, to
  - callouts
  - report writing
  - technical review
  - training and continuing education
  - vehicle/supply/equipment maintenance and ordering
  - any other crime scene related responsibilities or issues
- Revert to the duties in their regular discipline if the CSRT member has no pending CSRT related duties.

Supervisors must allow part-time CSRT members to focus on CSRT duties during their on-call week. Exceptions may be necessary for court appearances or to complete rush cases from the CSRT member’s regular discipline. When these exceptions occur, the supervisor will notify the CSRT Manager or designee. In the reverse circumstance, an extraordinary CSRT callout or rush CSRT report or review may take the CSRT member away from regular discipline duties during non-callout times. This will be coordinated between the member’s regular supervisor and the
CSRT Manager or designee. The CSRT Manager or designee and the supervisor of the part-time CSRT responder should work to resolve any issues that arise due to conflicting priorities.

While responding to a crime scene, all responders, full-time and part-time, are working under the supervision of the Crime Scene Response Team Manager or designee. Technical questions that arise at the scene will be referred to the CSRT Manager or designee.

5.2.8 Administrative

Administrative staff is non-technical and includes property and evidence custodians, office assistants, and office managers. The administrative section is responsible for proper receipt, documentation and storage of evidence, and general office and clerical duties in support of laboratory operations.

5.3 Budget

The CLD budget is part of the FLSB overall budget. Overall responsibility of this budget is under the direction of the FLSB Director. The CLD Commander is expected to manage, direct, and develop the CLD budget with oversight from the Bureau Director. The CLD Commander will keep the local Laboratory Managers informed of the status of the budget through the periodic Strategic Advancement Forum (SAF) report. The local Laboratory Managers have no individual laboratory budgets but are expected to manage operational expenditures at their level.

5.4 Communications

5.4.1 Policy

The CLD Commander, Laboratory Managers and Supervisors will establish a proper flow of communication internally throughout the CLD and externally with our customers. Management will ensure that within each laboratory all employees are well informed and employees at each level have input into the system. In addition, management will ensure that communication with our customers is effective and responsive to our customers’ needs.

Examples of various forms of communication to be used internally by the CLD include but are not limited to the following:

- Managers meetings
- Supervisors meetings
- Local Manager/Supervisors meetings
- Laboratory meetings
- Section meetings
- Conference calls
- Functional area meetings
• Interoffice Communications (IOC)
• E-mail
• FLSB Portal

Some examples of external communication are as follows:
• Personal contact by telephone, e-mail, letter, or in person
• Attendance at meetings of local law enforcement, prosecutors or medical examiners
• Forensic Investigation Council meetings
• Customer newsletters
• FLSB Forensic Services Guide
• Training provided to law enforcement, prosecutors or SANE nurses
• Membership and participation in WSP or State committees
• Customer surveys

The extent of database (e.g., CODIS, AFIS, NIBIN) searches shall be communicated to customers in the Forensic Services Guide and updated as needed, or in the laboratory report where applicable, and updated as needed.

5.4.2 Chain of Command

The chain of command is the hierarchical structure of authority and responsibility along which information is passed. The chain of command works in both directions. (See Organization Charts on FLSB Portal).

CLD employees will follow the chain of command for all internal communications as required by WSP Regulation 10.01.010. The chain of command, in ascending order, will normally be the employee’s Supervisor, the Laboratory Manager, the CLD Commander/Designee, the FLSB Director and the Chief of the Washington State Patrol.

The reporting authority is as follows:
• Each staff member is accountable to one and only one Laboratory Supervisor for each functional area.
• Laboratory Supervisors report to the Laboratory Managers.
• Laboratory Managers report to the CLD Commander.
• When a supervisor or manager is unavailable, the responsibility will fall to the level immediately below in the chain of command unless an alternate acting supervisor or manager has been designated.

The SAS Manager, Quality Process Manager, and Quality Assurance Manager may direct work to laboratory personnel and Technical Leads within the Quality Program with approval for use by the employee’s supervisor or manager.
5.5 **CUSTOMER FEEDBACK**

Customer feedback, both positive and negative, will be solicited. This may be accomplished through any of a number of methods:

- a periodic statewide customer survey
- a local focus group conducted by an individual laboratory manager and staff
- questionnaires submitted randomly to limited numbers of customers
- other direct interaction, both formal and informal, with specific customers.

The objective is to gather information to provide insight into the wants and needs of the customer agencies and how we can improve our service.

The decision to submit a statewide customer survey will be made jointly by the Laboratory Managers and the CLD Commander. Individual Lab Managers are responsible to document use of any other methods to collect feedback along with the feedback itself from their own service area.

Customer feedback will be reviewed, addressed at the annual management review, and used to improve the management system, laboratory activities and customer service.

Laboratory staff shall be willing to cooperate with customers or their representatives in clarifying the customer’s request and in monitoring the laboratory’s performance in relation to the work performed, provided that the laboratory appropriately ensures confidentiality to other customers.

5.6 **COMPLAINTS**

5.6.1 **Policy**

A complaint is an allegation of conduct or omission that is contrary to state statute, Washington Administrative or Civil Service Rules, Bargaining Unit Agreement, WSP Agency or CLD policies, regulations, rules, and procedures. They may include Quality Assurance policies and procedures, or an allegation of conduct or omission that could amount to misconduct, exercise of poor judgment, or failure to meet established standards. A complaint may be made against an individual CLD employee, a laboratory, a procedure or the Division.

Complaints regarding laboratory personnel, policies or procedures may come from internal or external sources (e.g., officers, prosecutors, defense attorneys, and the public). Complaints could be written or communicated orally. Personnel that become aware of a complaint either from an internal or external source have the responsibility to communicate the complaint through their chain of command. CLD management has the responsibility to ensure that complaints are resolved.
appropriately using one of the procedures outlined below and shall be responsible for all decisions at all levels of the handling process for complaints.

5.6.2 Procedure

Received complaints shall be documented and directed through the chain of command for review and action as appropriate. Non-Quality System complaints follow the WSP Agency Complaint Procedures (WSP Regulation Manual 12.00.020 Complaints), which includes a description of the process for receiving, tracking, recording, validating, and investigating the complaint, and deciding what actions are to be taken in response.

Complaints regarding any aspect of forensic testing and results of forensic analyses that do not conform to quality policies and/or procedures shall be directed to the SAS for evaluation. Procedures outlined in the section on Nonconforming Work and Corrective Actions shall be followed in these cases.

If management determines that the complaint originated due to a misunderstanding of laboratory policy, the manager may respond directly to the complainant and attempt to resolve the issue by discussing existing policies. Corrective or preventive actions may be initiated as a response as necessary.

Any changes or revisions to controlled documents resulting from complaints will follow the Document Control and Document Revision Policy and Procedure section of the Quality Operations Manual.

5.7 Impartiality of Laboratory Activities

5.7.1 Policy

The management of the CLD strives to ensure there is no influence on the professional judgments and actions of employees, including any undue internal and external commercial, financial or other pressures and influences that may adversely affect the quality of their work. Personnel shall not engage in activities that may diminish confidence in the laboratory’s competence, impartiality, judgment, or operational integrity. A conflict of interest, or an appearance of a conflict of interest, may arise when an employee has a personal relationship outside of work or a financial relationship with a suspect, victim, witness, police officer, attorney or judge involved in a case. This may include a relative or close friend personally or financially involved with any of the groups listed above. This list does not exhaust the possibilities of a conflict of interest. Even in cases where the employee doesn’t think that a conflict exists, a possible conflict of interest may exist in the eyes of an observer, which could lead to a diminished confidence in the laboratory and its work.
5.7.2 Procedure
Conflict of interest concerns and situations that could cause undue pressure that adversely affect the quality of the work shall be brought to the attention of management. Laboratory Managers have the responsibility and authority to take action on employee concerns within their labs and shall evaluate such concerns and take appropriate action.

Personnel with CLD funded memberships in the American Society of Crime Laboratory Directors, American Academy of Forensic Sciences, regional forensic associations, and/or certifications with professional certification bodies are bound by their codes of ethics.

Serious instances of undue influence or conflict of interest will be reported in accordance with the WSP Regulation Manual.

5.8 CONFIDENTIALITY
Every employee has the responsibility to safeguard all confidential information obtained in his or her official capacity from unauthorized distribution. In addition, employees will not access or disclose any confidential information except where authorized (see section on Disclosure and Release of Information). Subcontractors will sign a statement declaring that they will abide by this confidentiality policy. See also WSP Regulation Manual 8.00.240 CODE OF ETHICS – EMPLOYEES.

5.9 ETHICAL AND PROFESSIONAL RESPONSIBILITIES
Employee training and functional area training manuals shall include the application of ethical practices in forensic sciences. Many resources exist for review of ethical and professional responsibilities. These include, but are not limited to:

- American Academy of Forensic Sciences Code of Ethics and Conduct
- WSP Regulation Manual Chapter 8 Rules of Conduct (annual review required to be completed and documented)
- Guiding Principles of Professional Responsibility for Forensic Service Providers and Forensic Personnel

5.9.1 Guiding Principles
The Guiding Principles of Professional Responsibility for Forensic Service Providers and Forensic Personnel shall be reviewed annually by all laboratory personnel. The review is completed by viewing the ANAB Guiding Principles CLD on-line training presentation. Successful completion is documented on the employee training transcript through the agency eTrain system.
5.9.2 Cognitive Bias

Forensic Laboratories establish routine quality assurance and quality control procedures to ensure the accuracy of forensic analyses and the work of forensic practitioners. These quality control procedures should be designed to minimize cognitive bias. There are several different techniques that can be used to manage or recognize cognitive bias. Training manuals in the functional areas may incorporate these tactics in ways that are most appropriate for the category of testing. All forensic scientists will receive training in cognitive bias while administrative staff should be aware of the concept. Training materials are available on the FLSB Portal and include a “Cognitive Bias” PowerPoint presentation and selected articles. Additional training details can be found in the functional area training manuals.
6 CLD MANAGEMENT AND PERSONNEL

Top management for the CLD begins with the Bureau Director, who is responsible for all Bureau operations and management. Top management also includes the CLD Commander who sets direction for the Division laboratories.

Key management positions consist of the local laboratory managers and supervisors, who share the responsibility to ensure that policies, rules, procedures, directives, goals and guidelines adopted by the CLD are implemented, understood and practiced by all employees. In addition, the FLSB Standards and Accountability Section, which includes the Standards and Accountability Manager, Quality Process Manager, Quality Assurance Manager, DNA Operations Manager and the DNA Technical Leader, have key roles in Division operations, having quality oversight of Division work. The CODIS Administrator is also considered a key position. All technical leads (Forensic Scientist 4 positions) and the DNA Technical Leader are regarded as technical management and have key roles in our technical operations.

6.1 CRIME LAB DIVISION Commander

This position is responsible for managing all aspects of the Crime Laboratory Division with respect to its operation, organization, policy, and budget.

6.1.1 The CLD Commander:

- Prepares the legislative budget
- Coordinates operation of crime laboratories with other criminal justice agencies
- Define the areas of management authority and responsibility
- Is responsible to ensure that all policies, rules, procedures, directives, goals and guidelines are written in a clear manner, are consistent with Department Policy, state and federal Law, and are made available to division laboratories
- Translates policy into goals, objectives, and strategies, and projects a shared vision of the future to all employees.
- Ensures the Division’s operational objectives are achieved
- Ensures resources are utilized to their maximum effectiveness and all programs are providing the most effective and timely service
- Ensure that all employees recognize and support the Division’s Quality Assurance Program
- Supervises and trains Division management personnel
- As the CLD appointing authority, authorizes all hiring
6.2 **Laboratory Manager**

Laboratory Managers, who manage and operate the CLD Crime Laboratories and the Crime Scene Response Team, are responsible for overseeing, monitoring and ensuring conformance to the Quality Management System. In this respect, they function as quality assurance managers for their individual laboratories. The Crime Scene Response Team Manager manages the Crime Scene Response Team consisting of multiple employees across the CLD.

### 6.2.1 The Laboratory Manager:

- Is accountable for overall forensic operations within a given laboratory
- Provides clear direction and expectations to employees in the laboratory
- Works collectively with other CLD Laboratory Managers and Supervisors, the CLD Commander, and the Standards and Accountability Section to ensure operations are coordinated on a statewide basis as a single system culture
- Assists the CLD Commander in developing and implementing division-wide policies and procedures
- Exercises control over discretionary funds for laboratory supplies, overtime, and training
- Works with employees under their supervision to resolve complaints
- Administers discipline to subordinates as appropriate
- Ensures the effective application of the Division’s Quality Assurance Program and is responsible for the quality of the operations for their laboratory
- Ensures the personnel under their supervision receive appropriate training

The CLD Commander will assign CLD Managers to act as Management Liaison for one or more of the following functional areas: Materials Analysis, DNA, Latent Prints, Firearms, Questioned Documents, Crime Scene Response, and Administration. In addition to duties associated with local crime laboratory management, Management Liaisons will have additional duties pertaining to the operation of the functional area throughout the division, to include:

- Develops and updates effective plans to address program direction, staffing, training, equipment, and quality assurance needs within the discipline
- Represents the needs of the functional area to division management, and acts as a conduit between the functional area forensic scientists and division management
- Ensures supervisory and functional area meetings are held and records of meetings are kept
- Works with the functional area technical lead(s) and SAS to determine root causes for systemic or laboratory-level technical non-conformities
• Manages training budget for the functional area; ensures training and travel requests are submitted appropriately according to CLD training needs and available resources
• In collaboration with the functional area technical lead(s), reviews technical and training manuals and revisions to verify conformance with the CLD Quality Operations Manual, and accreditation requirements
• Facilitates communication within the functional area so that all labs meet division goals and objectives as a single entity
• Reviews and approves functional area-specific research projects

The DNA Operations Manager is the Management Liaison for the DNA discipline.

6.3 CODIS MANAGER

The CODIS manager is the system administrator of the WSP Crime Laboratory Division’s Combined DNA Index System (CODIS) network and is responsible for the security of DNA profile data stored in CODIS.

6.3.1 The CODIS Manager:
• Manages and operates the CODIS Laboratory
• Is responsible for the overall quality and security of the DNA data that is entered into CODIS by the six laboratories within the WSP CLD that perform DNA typing
• Monitors the oversight of CODIS computer training and quality assurance data
• Has the authority to suspend or terminate the laboratory’s participation in CODIS in the event of a problem until the reliability of the computer data can be assured
• Monitors the Convicted Offender Collection Kit distribution program for the collection of reference samples for all individuals convicted of a qualifying offense
• Advises senior management on DNA database operations
• Participates in strategic planning
• Represents the Crime Laboratory Division at the state and national level

6.4 STANDARDS AND ACCOUNTABILITY SECTION (SAS)

The SAS is responsible for ensuring the overall quality of forensic service and for monitoring conformance with policies and procedures. The Section is responsible for the development, implementation, operation and continuous improvement of the Quality Assurance Program. Any quality issues will also be shared with the responsible laboratory manager. The laboratory managers report directly to the Standards and Accountability Section Manager (SAS Manager) on all issues.
regarding the Quality Assurance Program. The SAS Manager has direct access to the CLD Commander where decisions are made on laboratory policy and resources.

This SAS is composed of the SAS Manager, Quality Process Manager, Quality Assurance Manager, DNA Operations Manager, DNA Technical Leader and support staff.

6.4.1 Standards and Accountability Section Manager:

The Standards and Accountability Section Manager:

- Is also the Assistant Division Commander (ADC) and acts as CLD Commander in the Commander’s absence
- Provides management oversight of the Standards and Accountability Section and reports directly to the CLD Commander
- Supervises the Quality Assurance Manager, Quality Process Manager, and DNA Operations Manager
- Attends FIC and SAF meetings to represent Standards and Accountability as a statewide division function
- Provides advice and recommendations on the operations of CLD
- Assists in development and implementation of leadership professional development programs for managers and supervisors
- Assists in compiling and analyzing data to support resource allocation decisions
- Oversees the CLD Quality Assurance and accreditation programs
- Assists the CLD Commander in developing and implementing division-wide policies and procedures
- Provides oversight of the nonconformance and corrective action process
- Oversees internal audits: ensures annual internal audits and QAS audits are performed on a regular schedule and reviews and approves final internal audit reports
- Ensures internal auditors are trained
- Organizes, coordinates and facilitates CLD Management Team meetings
- Organizes, coordinates and conducts annual Management System Reviews (MSR) and provides MSR summaries to the CLD Commander and Bureau Director
- Ensures timely review, approval, and posting of manual revisions
- Oversees CLD public disclosure and ensures timely and responsive public disclosure
- Oversees grant management
- Periodically reviews nonconformances to identify patterns and trends
- Works to continuously improve the quality system
- Approves training plans
- Reviews and approves manuscripts for publications and research projects
• Reviews and approves new methodology, technologies and equipment validations/verifications

6.4.2 Quality Process Manager (QPM):
The Quality Process Manager:
• Supervises the grant specialist and forms and records analyst positions
• Works to maintain and improve the quality program of the WSP Crime Laboratory Division
• Administers and coordinates the proficiency testing program, including document retention and responses to inquiries from the accrediting body Proficiency Review Committee (PRC)
• Oversees the interlab technical review program
• Serves as liaison to the WSP Human Resources Division (HRD): oversees the hiring program for the division; leads recruiting and hiring program for CLD staff
• Serves as the Public Information Officer for the Division
• Coordinates and publishes the Customer Newsletter
• Serves on the Bureau Safety Committee as the headquarters representative; assists in the oversight and implementation of the safety program
• Coordinates and monitors content of the FLSB Portal
• Maintains and issues Quality System documents and records, including all manuals and forms
• May be involved in the review of root cause analysis and corrective actions for nonconformities and inconsistencies in all testing
• Works with Risk Management Division providing documentation of CLD compliance with the Commission on Accreditation for Law Enforcement Agencies (CALEA) criteria
• Oversees regular audits of CLD firearms and drug reference material collections and participates in other quality audit activities as needed

6.4.3 Quality Assurance Manager (QAM)
The Quality Assurance Manager (formerly Laboratory Accreditation Manager):
• Functions as quality assurance manager for maintaining accreditation
• Monitors conformance with and maintains documentation of accreditation standards
• Functions as the point person for interactions with ANAB and all issues pertaining to accreditation for the CLD
• Coordinates and submits ANAB yearly conformance documents for all laboratories
• Works with ANAB to plan for surveillance visits and assessments, including logistics for hotels and in-state travel for assessors
• Devises, in conjunction with laboratory managers, corrective action plans to address nonconformances from surveillance and assessment activities
• Prepares responses to other ANAB inquiries
• Oversees the development and evaluation of revisions of CLD wide manuals for adherence to accreditation standards
• Conducts an annual review of the CLD Manuals
• Ensures manuals and procedures reflect accreditation requirements
• Coordinates the internal audit program
• Oversees the corrective action process in all testing

6.4.4 DNA Operations Manager
The DNA Operations Manager:
• Supervises the DNA Technical Leader
• Advises senior management on DNA operations and acts as the management liaison for DNA
• Participates in strategic planning
• Participates in obtaining and managing federal DNA-related grants, to include periodic grant adjustments, mandatory grant reporting, and facilitating procurements which utilize grant funding.
• Manages and secures DNA-related contracts in coordination with Budget and Fiscal Services
• Monitors DNA-related funding and budgets
• Represents the DNA Functional Area/CLD at the state and national level
• Develops, reviews, and approves manual and form updates in coordination with the DNA Technical Leader, Quality Process Manager, and Quality Assurance Manager
• Works with the DNA Technical Leader to provide training opportunities and coordinates training events for the DNA Functional Area
• Organizes annual functional area meetings and Forensic Scientist 4/5 meetings

6.4.5 DNA Technical Leader (TL)
The DNA Technical Leader manages the technical operations of the DNA functional area throughout the Crime Laboratory Division. This individual will be directly responsible for quality issues involving the DNA functional area and works with the SAS Manager and/or QAM on all quality matters (all documents on reviews, proficiency tests, reanalysis, etc.). The DNA TL works provides copies of quality documents relating to quality assurance measures in DNA to the SAS Manager and/or QAM, and works with SAS on corrective actions and recommended changes for continuous improvement of DNA quality assurance.

The DNA Technical Leader:
• Ensures all ANAB accreditation requirements are met by the DNA functional area system-wide
• Assists with coordination of external DNA audits
• Ensures CLD conformance with the FBI Quality Assurance Standards
• Monitors continuing education for the DNA functional area
• Works with the Standards and Accountability Section to develop the Quality Assurance Program for DNA
• Has the authority to initiate, suspend and resume DNA work if needed when quality issues are identified
• Has oversight and evaluates the validation for new methods
• Has oversight of the CODIS program
• Advises senior management on DNA operations
• Participates in strategic planning
• Represents the Crime Laboratory Division at the state and national level

6.4.6 DNA QA Support Technical Lead Forensic Scientist

• Assists the DNA Technical Leader in various quality assurance duties
• Provides DNA functional area support in the quality assurance processes including the review of proficiency test results, tracking quality variances and participating in audits and assisting with DNA Technical Leader site visits
• Assists in the interpretation of unusual or complex casework including mixture interpretation, kinship and paternity
• Prepares and presents training materials including training in mixture interpretation, use of appropriate stats and for the implementation of new technology.
• Participates in new technology validations and committees.
• Provides support for DNA outsourcing projects including writing technical review protocols, providing training in the protocols, distribution of technical reviews, liaisons with the participants, and keeping all the associated quality and protocol documents up-to-date.
• Provides support for post-conviction DNA testing including the status of in-house and outsourced case requests, liaisons with the participants, provide DNA testing assessments on possible candidate cases and compose declarations as required.
• Reports and provides assistance to the DNA Technical Leader in technical oversight.

6.4.7 Grant Specialist

The Grant Specialist helps develop and apply budgets and timelines for grant application planning, and logistics for implementation of awards. Additionally, the incumbent prepares orders, competitive procurement drafts and vendor contracts
to carry out purchases for goods and services specified in grant awards, and provides assistance in preparing grant progress reports.

6.4.8 Public Disclosure Tracking Coordinator
The Public Disclosure Tracking Coordinator, typically a Forms and Records Analyst, is primarily responsible to coordinate responses to all public disclosure requests for the CLD, to include delivery and distribution of requested records, and tracking disclosure responses.

6.5 CRIME LAB DIVISION PERSONNEL

6.5.1 Forensic Scientists
The Forensic Scientist class series consists of Forensic Scientists 5, 4, 3, 2 and 1, detailed below. The primary functions of forensic scientists generally include:

- Examination and/or collection of evidence
- Analysis of the physical evidence using accepted and validated methods and analytical instrumentation
- Preserving evidence according to laboratory procedures
- Maintaining chain of custody, i.e., documentation establishing the receipt, handling, and disposition of evidence
- Interpreting observations and test results; preparing written opinion reports
- Testifying as an expert witness in courts of law
- Participating in proficiency testing
- Receiving on-going training and professional development

6.5.1.1 Forensic Scientist 5
This position supervises forensic scientists and support staff within a forensic laboratory.

The Forensic Scientist 5 (FS5 or Supervisor):

- Is responsible for the overall daily administration, operation and coordination of activities within an operational unit of the laboratory
- Ensures compliance with standards of quality established by the Quality Manual, accreditation requirements, and the courts of the State of Washington
- Monitors quality through review of staff case work
- Assigns and directs tasks to be performed by laboratory staff
- Has a working knowledge of all facets of an accredited laboratory
- Ensures the success of the work-unit through effective leadership, proper training of personnel, effective management of resources, as well as, providing a safe working environment
- Is responsible to address operational, personnel and customer issues
• Analyzes case evidence, prepares written reports and testifies in court as required
• Works to ensure the volume and priorities of work are appropriate for the needs of the assigned functional area of the laboratory
• Represents the needs of the local lab functional area to the Management Liaison and, for the DNA functional area specifically, to the DNA Technical Leader
• Supports communication within the functional area to help meet division goals and objectives as a single entity
• Oversees implementation of training plans, verifies that trainees adhere to established training timelines, and verifies that training is conducted to the highest standards
• Seeks feedback from the functional area technical lead(s) (and for DNA, the DNA Technical Leader) regarding technical issues
• Works with the functional area technical lead(s) (and for DNA, the DNA Technical Leader) to develop training plans
• Resolves conflicts between case analyst and technical reviewer when disagreements occur

6.5.1.2 Forensic Scientist 4 (Technical lead)

This position serves as a forensic technical lead in a specific discipline or functional area of forensic science in a crime laboratory. The Technical Leads in the various functional areas are a vital part of the quality assurance program. The CLD Commander will appoint Technical Leads to provide quality assurance program support through technical oversight and leadership in the functional areas. In the CODIS Laboratory, the FS5 (supervisor) assumes the duties of the Technical Lead.

The Forensic Scientist 4 (FS4):
• Performs complex analyses on physical evidence. This involves casework where applied research, method modification, or a unique approach may be necessary; or a single definite conclusion is not possible and a weighted conclusion is warranted; or casework requiring the reconstruction of an event or series of events based upon the interpretation of physical evidence
• Works with the Standards and Accountability Section in the performance of their assigned quality assurance duties
• Is accountable for the quality of the casework product, for conformance with all applicable accreditation and audit criteria, conformance documentation, validation of new technology and methods, and investigation of assigned casework nonconformances
• Has responsibility for uniform methodology implementation and use in all laboratories within the discipline and responsibility for evaluating all methods used
• Has the responsibility to oversee standard training for new employees, and re-training of existing employees as needed in conjunction with the employee’s supervisor, within the discipline
• Has the responsibility to see that quality practices are utilized in all scientific equipment maintenance, and ensure appropriate quality control is implemented within the discipline
• Ensures that procedures and training manuals for the discipline accurately reflect established standards and comply with accreditation requirements
• Evaluates new analytical procedures, equipment or technologies, oversees their validation and assists with implementation
• Reviews proposed research projects
• Assists in coordination, content, and execution of the Functional Area Meetings to include setting up training workshops in conjunction with the FAM
• Supports communication within the functional area that helps all labs meet division goals and objectives as a single entity
• Conducts regular site visits to laboratories in the division to provide one-on-one mentoring of functional area scientists
• Ensure methodologies are in conformance with health and safety requirements
• Assists supervisors with resolving disagreements between case analysts and technical reviewers, resolving other technical issues, and assists Standards and Accountability with root cause analysis involving technical nonconformities
• Has the responsibility to recommend the termination of testing in their discipline in the event of a technical problem with a technical procedure, instrumentation or equipment. Communication of such action must follow the appropriate chain of command.

6.5.1.3 Forensic Scientist 3

This is the senior level of the Forensic Scientist series.

The Forensic Scientist 3 (FS3):
• Scientifically analyzes evidence in routine, non-routine and complex casework in an area(s) in which they have been authorized and are proficient and productive. Complex analysis involves casework where applied research, method modification, or a unique approach may be necessary; or a single definite conclusion is not possible and weighted conclusion is warranted; or casework requiring the reconstruction of an event or series of events based on the interpretation of physical evidence. Databasing scientists develop DNA profiles from convicted offenders for inclusion in the Combined DNA Index System (CODIS).
- Formulates sound conclusions from data without exceeding the boundaries of the data. Data may result from complex analyses involving multiple methods and techniques, multiple items and examinations
- Reports scientific findings in the form of a written forensic laboratory report based on the interpretation of observations and analytical test results
- Completes analyses of high quality, performed using forensically accepted scientific methods, and in accordance with the CLD Quality Operations Manual and the Section Technical Procedures Manual
- Documents and protects evidence according to laboratory procedures, ensuring that the chain of custody is maintained
- Provides technical review and administrative review
- Follows the standards of accreditation, laboratory and agency policies
- Participates in the proficiency testing program by successfully completing all assigned proficiencies and competencies
- Provides training as needed
- Assures that all assigned instruments have undergone the appropriate performance checks and maintenance
- Participates in assessment preparation and the internal audit program
- Receives continuing education as it relates to their discipline, including functional area meetings, conferences, seminars, and associated training.
- Testifies as an expert in courts of law

6.5.1.4 Forensic Scientist 2

This is the journey level of the series.

The Forensic Scientist 2 (FS2):
- Will have completed the majority of their training in an assigned discipline and will focus on the routine analysis of physical evidence. Routine analysis involves laboratory examination in which the items to be tested require a single specific examination or a standard battery of examinations or analyses, the results of which lead to a conclusion acceptable to experts in the field. Databasing scientists develop DNA profiles from convicted offenders for inclusion in the Combined DNA Index System (CODIS)
- Formulates sound conclusions from data without exceeding the boundaries of the data
- Reports scientific findings in the form of a written forensic laboratory report based on the interpretation of observations and analytical test results
- Completes analyses of high quality, performed using forensically accepted scientific methods, and in accordance with the CLD Quality Operations Manual and the Section Technical Procedures Manual
- Documents and protects evidence according to laboratory procedures, ensuring that the chain of custody is maintained
- Provides technical review and administrative review
- Abides by accreditation criteria
- Testifies as an expert in courts of law

6.5.1.5 Forensic Scientist 1

This is the entry level of the series.

The Forensic Scientist 1 (FS1):
- Works in a training capacity and under close supervision
- Performs beginning level analyses of physical evidence in criminal cases submitted to the forensic laboratory. Databasing scientists develop DNA profiles from convicted offenders for inclusion in the Combined DNA Index System (CODIS)
- Interprets analytical results, prepares written opinion reports, and testifies as an expert witness in courts of law
- With on-the-job training, the incumbent learns entry-level casework in a limited area in order to become proficient in a discipline of forensic science
- Abides by accreditation criteria

6.6 Technical Support Staff

6.6.1 Laboratory Technician

The Lab Technician performs routine tasks following clearly defined laboratory procedures, performs quality control measures and operates routine laboratory equipment. The Lab Technician may or may not be proficiency tested depending upon their assignment.

6.7 Administrative Staff

6.7.1 Property and Evidence Custodians

The Property and Evidence Custodian (PEC) receives evidence into custody from law enforcement agencies and releases evidence back to the submitting agency upon completion of analysis. They maintain the appropriate records of evidence transactions using the Laboratory Information Management System (LIMS). They also conduct periodic inventories and audits of evidence depositories. In addition, in some laboratories they may be responsible for general office and clerical duties, including ordering supplies, preparing pay documents, report preparation and filing as required by their supervisor. PECs may be required to testify in court.

6.7.2 Office Manager

The Office Manager (OM) plans, organizes, assigns, and supervises varied and extensive processing and service units and related central office activities, including evidence handling.
6.7.3 **Administrative Assistant**

The Administrative Assistant (AA) performs a variety of complex clerical duties in support of office and CLD operations.

6.7.4 **Office Assistant**

The Office Assistant (OA) performs a variety of routine clerical duties in support of office or unit operations.

6.7.5 **Trooper Cadet**

Occasionally trooper cadets may be assigned to a laboratory to assist with administrative tasks and functions. These positions are temporary and depend on the availability of cadets and the needs of the Agency.

6.7.6 **Volunteers and Interns**

Volunteers and interns may participate in research projects, data and reference materials collections, or other projects suitable to their knowledge, skills and abilities, but may not work directly on casework. Prior to beginning work in the crime lab, all volunteers and interns are required to submit to and pass a polygraph test and background investigation.
7 PERSONNEL QUALIFICATIONS AND TRAINING

7.1 Policy

The Division will ensure that personnel performing specific tasks shall be qualified on the basis of appropriate education, training, experience and/or demonstrated skills.

7.2 Qualifications of Personnel

All personnel assigned to or contracted by the WSP CLD must be competent, trained and supervised to ensure that they conduct work according to the quality program of the CLD (ISO 5.2.3). It is the responsibility of the laboratory manager to demonstrate the competence of all staff, whether directly employed or contracted. There must be documented evidence of the training and qualifications for all staff.

A Position Description Form (form OFM 12-002 WGS, hereafter PDF) shall be completed for all CLD employees. The PDF shall be retained in the employee’s supervisory desk file and shall be updated as necessary. The PDF includes minimum education and experience requirements for laboratory management, technical management and staff according to their respective positions.

When an individual under contract is conducting casework for the CLD, the supervisor or laboratory manager will advise the customer in writing of the subcontracting arrangement and will gain approval from the customer, preferably in writing. Communication to the customer regarding the transfer or referral of casework to another WSP laboratory is not required. Convicted offender DNA samples may be examined by contract personnel without pre-approval from the collecting agency.

7.3 Educational Background

Educational requirements for CLD technical positions are found on the Washington State Office of Financial Management website. The PDF includes educational requirements for each position. Verification of educational requirements for staff is under the purview of the Washington State Department of Enterprise Services (DES) and WSP Human Resource Division (HRD). The QP Manager will ensure that college transcripts of all employees are reviewed before employment. Transcripts are typically retained at the employee’s laboratory or at HRD. Copies of transcripts for those scientists performing casework in the DNA unit and CODIS are located in the employee’s assigned laboratory.
7.4 **HIRING PERSONNEL**

All employees shall be hired according to rules that govern their position. The supervisor and the laboratory manager of the position to be filled shall be involved in the hiring process. The CLD Commander has appointing authority and makes the final decision on hiring new employees.

7.5 **PERSONNEL TRAINING AND DEVELOPMENT**

CLD Management will ensure that proper training occurs for all CLD staff. Laboratory managers will ensure that employees meet and maintain competency requirements. In addition, each employee will share in the responsibility of maintaining his/her functional area discipline expertise and competency.

7.5.1 **Training and Development Goals**

Training goals include:

- Basic functional area competency
- Maintenance of acquired skills, abilities and expertise
- Instruction in new and improved techniques
- Acquiring and maintaining professional accreditation or certification
- Application of ethical practices in forensic sciences, a general knowledge of forensic science, and applicable criminal and civil law and procedures
- Meeting agency requirements for mandatory training and policy awareness
- Where applicable, training in the presentation of evidence in court
- Training in ethics and professional behavior expectations

7.5.2 **Provisions for Training, Maintenance of Skills and Expertise, and Development**

Various opportunities are available for training, maintenance of skills and technical competence, and employee development, including, but not limited to:

- Training from CLD personnel experienced in a variety of forensic analyses and processes
- CLD functional area in-service training
- CLD sponsored forensic training courses utilizing visiting experts
- WSP sponsored training
- Workshops, seminars and webinars
- State educational and career development resources
- Continuing education opportunities available through universities and community colleges
- Agencies and institutions such as the FBI, National Forensic Science Technology Center, National Institute of Justice, California Criminalistics
Institute, Bureau of Alcohol, Tobacco and Firearms, Washington Criminal Justice Training Commission, and others

- Professional forensic science organizations such as the American Academy of Forensic Sciences, the Northwest Association of Forensic Scientists, the Association of Firearm and Tool Mark Examiners, and others
- Journals of professional forensic science organizations and other scientific literature

The CLD will periodically provide discipline in-service training opportunities at Functional Area Meetings for the purpose of maintaining skills and expertise, and exchanging technical information on novel discipline procedures, techniques, and/or research developments. Such in-service meetings will occur at least annually for Forensic Scientists and Property and Evidence Custodians and are aimed at scientific advancement, process improvement, solving technical problems, and identification of relevant training needs and opportunities. These goals support the continuing professional development and maintenance of competency of individual employees which in turn support the overall competency of the CLD programs. The CLD will support the functional area meetings and endeavor to act on recommendations when possible.

The CLD will provide support to employees who wish to pursue personal certification through a relevant professional organization. The laboratory manager may provide some work time for study and preparation. The cost of the initial application, testing fees, and subsequent maintenance fees are the responsibility of the individual employee.

Attendance at conferences and workshops sponsored by professional forensic organizations is an effective way for employees to stay current in their field. This venue provides a significant source of continuing education that directly supports their professional development and maintenance of competency. Serving as members or officers of these organizations facilitates employees staying in contact with their peers across the nation, a process vital to scientific advancement. In order to achieve these goals, the CLD will provide membership in two relevant professional forensic organizations for each forensic scientist. The CLD will endeavor to send at least one scientist to each of the annual conferences sponsored by the major professional forensic organizations with a level of financial support consistent with current resources.

### 7.6 **TRAINING PROGRAM**

#### 7.6.1 Policy

The CLD will have a documented training program to include new employee training, training in a new discipline, retraining and continuing education for maintaining skills and expertise. Prior to being authorized to perform assigned
duties, trainees will successfully complete the applicable training program. The effectiveness of the training program shall be evaluated.

All technical and technical support staff who perform testing or sampling, regardless of academic qualifications or past work experience, shall satisfactorily complete competency testing prior to assuming casework responsibility in the laboratory. Crime Scene personnel shall satisfactorily complete a competency test prior to assuming primary responsibility for the examination, documentation and processing of a crime scene.

Each laboratory shall maintain records of the relevant authorization(s), competence, educational and professional qualifications, training, skills and experience of all technical personnel, including contracted personnel. This responsibility lies primarily with supervisors, but individual scientists should maintain copies of their own training records. The records to be maintained include documentation of completion of specific training modules as appropriate, successful completion of written tests, summaries of oral tests and responses, competency tests and associated reports. This information shall be readily available for review by an assessor and shall include the date on which authorization and/or competence is confirmed.

Training records will be sufficiently detailed to provide evidence that employees have been properly trained and that their ability to perform the task of their specific discipline has been assessed.

Training needs of an employee will be identified through individualized training plans and goals, CLD strategic plans, management requests and needs of the customer agency.

7.6.2 Trainer/Trainee Method
The CLD employs the trainer/trainee method as one component in teaching for technical discipline training and training in a new job classification. The trainer and trainee will have management commitment and resources to provide a quality training experience. Training may take place at sites other than the trainee’s assigned laboratory.

7.6.3 Trainer
Trainers have the responsibility for ensuring the trainee successfully completes the training plan. The trainer will be selected by the supervisor, DNA Technical Leader, Technical Lead and/or lab manager and should have the following abilities:
- Have an understanding of WSP and CLD structure, policies, and procedures.
- Have an understanding and working knowledge of the current procedures, requirements, and expectations for the given functional area or discipline.
• Ability to instruct and train based on the training manual for the given discipline
• Ability to offer constructive criticism and positive reinforcement that is crucial to the trainee’s learning process
• Ability to interact routinely, frequently, and one-on-one with the trainee(s) to assess their understanding and mastery of the subject matter
• Ability to record the trainees’ progress with evaluations per CLD and/or functional area requirements and routinely report on progress through the chain of command.
• Good organizational, verbal and written skills
• It is expected that senior level staff be able to serve as trainers in their functional area. Journey level staff may additionally assist with training as appropriate.

Instructors interact with the trainee on a limited basis to instruct a small portion of the training. Instructors are typically used for teaching of one or more aspects of a technical discipline or administrative topic.

### 7.6.4 Competency Test

A competency test is an examination provided to a trainee at the end of training modules or at the end of the training plan for a specific functional area discipline. The competency test results are evaluated by the assigned trainer and lab supervisor of the discipline. For any laboratory personnel whose job responsibility includes casework testing, performing specific tasks that create items that could be used for testing, test report writing, technical review or testimony, competency testing will occur and include, at a minimum:

- Examination of sufficient unknown samples to cover the anticipated spectrum of assigned duties and evaluate the individual’s ability to perform proper testing methods;
- A written report to demonstrate the individual’s ability to properly convey results and/or conclusions and the significance of those results/conclusions; and
- A written or oral examination to assess the individual’s knowledge of the discipline, category of testing, or task being performed. For testimony competencies, a written examination is not sufficient to demonstrate competency: an oral presentation is required.
- If testimony is within the anticipated work of the individual, a written evaluation conducted by a technically competent reviewer of the individual’s ability to provide testimony must be completed.

### 7.6.5 Training Manual

A training manual outlines the necessary requirements to become competent in a forensic discipline or functional area. The training manual is designed to provide
employees with an understanding of theory and principles, application, methodology, technical limitations, and equipment involved in the functional area. The manual is further intended to provide the trainee with a sufficient understanding and skill level to satisfactorily conduct independent casework examination in the forensic discipline or category of testing. It includes modules or sections of reading assignments, study/discussion questions, and practical exercises. The technical procedures or training manuals will contain the approved methods, the scientific references and resources, and the requirements for successfully completing a training program for each discipline.

Training manuals for each functional area will provide a detailed training outline for new and permanent employees. These training manuals will also be used as a guide when designing job performance improvement plans.

7.6.6 Training Plans

Supervisors will work with each employee to develop a training plan. In developing the training plan, the supervisor must consider the needs of the individual employee, the discipline, the CLD and the customer agencies.

Training plans developed for each functional area will be used for developing new employees. Training shall include the application of ethical practices in forensic sciences, a general knowledge of forensic science, testimony and applicable criminal and civil law and procedures. All training modules included in an employee’s training plan will have clearly defined goals for successful completion which are measurable in order to document progress and achievement of intended results. Measurement tools will normally include, but are not limited to, scored examinations and competency tests where pass and fail is defined.

If supervised casework is required in the training plan (this is not a requirement in all disciplines and may not be required for all training plans within a discipline), the number and type of cases shall be specified in the training plan. Competency testing for the testing being supervised must have already been successfully completed prior to supervised casework.

The training plan will be updated annually during the employee’s performance evaluation and may be adjusted as needed throughout the year. Supervisors must be actively involved in the employees’ training, including documenting training events and training performance in the employee’s supervisory desk file.

An employee adding or transitioning from one technology to another within the same functional area will undergo appropriate training from a developed training plan, followed by a competency test to ensure that they have demonstrated proficiency in that technology. For a newly validated method in the DNA Functional Area, a competency test may be waived by the DNA Technical Leader for those who have demonstrated their necessary competency in validation study work.
7.7 **Re-Training**

Re-training in a given discipline may be required when:

- Employees were once qualified in the discipline or functional area but have not maintained the required competency in that discipline or functional area
- Qualified analysts who, because of leave or other circumstances, are unable to complete a proficiency test in their discipline in a calendar year
- Employees were previously qualified at another laboratory system (non-WSP) in the particular functional area
- A discipline’s procedure or training manual, in the judgment of the functional area supervisors and technical leads, has been revised to the extent that re-training is necessary.
- Directed by a corrective action plan (CAP), job performance improvement plan (JPIP) and/or remedial training
- Required by administrative rule

Re-training will include an evaluation of current knowledge and skills by the supervisor in consultation with technical lead(s) familiar with the sub-disciplines required, and the laboratory manager. A training plan will be prepared and approved by those involved in the evaluation and will include, as needed, any reading, remedial training modules, and written tests deemed necessary. Successful completion of competency testing and authorization by the CLD Commander will be required before resuming casework.

7.8 **Training Program Procedures**

The following steps will be followed to ensure successful completion of the training program:

- Assign Trainer: The trainee will work under the direction of a trainer, who is assigned by the Technical Leader, supervisor, and/or lab manager.
- Develop Training Plan: The supervisor and/or technical lead/technical leader will develop a comprehensive training plan for each new employee with estimated timelines. This comprehensive plan will include the functional area specific training plan(s) and be approved by the SAS Manager. Modified training plans will require documented justification for the modification and approval from the supervisor, laboratory manager, functional area Technical Lead, or for DNA and CODIS, the DNA Technical Leader. Training plans may also be developed for journey level employees as required.
- The trainee will work with the assigned trainer and instructors to successfully complete the training plan. Completion of the required training elements will be documented by both the trainee and the trainer. The trainee’s supervisor will be notified of successful completion of the training plan.
• In certain areas it is necessary for trainees to get experience looking at and working with real world casework samples in order for them to gain the necessary knowledge and skill to become competent in that area. Some examples are microscopic comparisons of bullets, comparison of latent print evidence, questioned document examinations and various microanalysis procedures. In these areas, the examination of casework evidence by a trainee is permitted if the examination is overseen by a qualified Forensic Scientist, the examination is non-destructive and will not alter the evidence, and the trainee has completed applicable competency testing for that task prior to examination of casework evidence. The examination of the casework samples is purely for training purposes. The original analyst on the case will review the trainee’s documentation and provide feedback on the examinations. Additionally, the original casework analyst will ensure all of the evidence has been properly sealed and is ready for return to the submitting agency.

• Training Evaluations: During the period of training, training evaluations will be completed and documented by the trainer. The results of these evaluations will be discussed with both the trainee and the supervisor of the trainee. Progress on training will be reported monthly to the laboratory manager until the training concludes. Copies will be forwarded to the Standards and Accountability Section as requested.

• Mock Case Samples: During training, the analyst may be provided mock case samples in which he/she can gain experience in case management, case file preparation, and report writing. The number and type of mock cases will be determined by the trainers and documented in the training plan.

• Competency Tests: The supervisor and/or technical lead will be responsible for administering competency tests. The competency tests must be successfully completed by the trainee prior to the start of casework. The results will be maintained as part of the training record.

• Moot Court: Trainees whose duties anticipate providing testimony will participate in moot court prior to performing testing or performing specific tasks that create items that could be used for testing. Evaluations and feedback from the participants will be provided to the trainee and the supervisor.

• Training Records: Training records of the trainee will be retained by the trainee and/or supervisor.

• Approval to Conduct Forensic Analysis (or to perform specific tasks that create items that could be used for testing): The trainee must demonstrate the successful completion of the training plan by passing all examinations and competency tests that are part of the training plan. Upon successful completion of either training modules or the final training plan, an IOC will be prepared by the trainer and submitted through the employee’s chain of command to the Division Commander for final approval before the trainee
can begin work in that defined area. The approval documentation must include the appropriate functional area Technical Lead or, for a scientist in DNA or CODIS, the DNA Technical Leader. A copy of the final approval will be given to the SAS Manager. The analyst will be authorized to perform work in only those areas in which he/she was approved. Personnel authorized to perform casework are also authorized to develop, modify, verify and validate methods, and report opinions and interpretations within their area of authorization.

- Once approved for analytical work, the analyst may be required to perform supervised casework in accordance with the applicable functional area training and/or technical procedures manual. Supervised casework reports completed by the analyst will be signed by the analyst. The scientist monitoring the casework will sign, or initial, and date the draft report. The casework would then go through the normal technical and administrative reviews, with the scientist monitoring the casework being excluded from the technical review.

- For analysts already authorized to conduct independent casework, but completing supplemental training plans for new instrumentation or analytical technology, an IOC will be prepared by the trainer and submitted through the employee’s supervisor, technical lead or for DNA and CODIS, the DNA Technical Leader, and then to the lab manager for final approval before the analyst can conduct casework using the new instrumentation or analytical technology.

- Approval to Perform Technical Review: Authorization to perform technical review of other analysts’ casework must be documented in an IOC. This may be included within the IOC documenting the successful completion of the analyst’s training plan, or a separate IOC. If a separate IOC is issued, it will be submitted through the chain of command, with final approval by the Laboratory Manager. The approval documentation for a scientist in DNA or CODIS must also include the DNA Technical Leader. A copy of the authorization to conduct technical review will be provided to the SAS Manager. Supervisors will be responsible to assign reviews commensurate with the analyst’s experience.

- Concluding the Training Process: At the conclusion of the training program, the effectiveness of the training program shall be evaluated by the trainer, appropriate technical lead and/or trainee based on the trainee’s performance on written exams, competency tests, practical exercises, and/or mock trial results. Improvements to the program are made as needed and the review is documented through the annual review of controlled documents.

7.9 **Requests for Training**
Refer to WSP Regulation Manual 10.12.020 TRAINING DIVISION for instructions on completing a Training/Travel Request (WSP Form 3000-320-016). Training requests will be routed through the chain of command as instructed in the Regulation Manual. Training requests incurring costs specific to a functional area will include approval by the Management Liaison. All training requests will be approved or denied within thirty (30) calendar days from the submission of a properly completed request. If a request is denied, the person denying the request will provide a reason for the denial to the employee.

Employees engaged in their planned training program in their laboratory generally do not need to submit Training/Travel Requests (TTR) for the training modules in their program unless the above conditions apply. All training/travel events incurring costs will be documented on the CLD Training Spreadsheet.

7.10 COMPLETION OF TRAINING

Refer to WSP Regulation Manual 10.12.020 TRAINING DIVISION for instructions on submitting a Report of Training using the TTR form. In addition, a Report of Training must be completed for training received at functional area meetings. The completed form must be routed through the supervisor and then appropriately routed for entering the information into the employee training records by the local eTrain user or Administrator. Copies of all training and continuing education records for completed training will be maintained at the employee’s laboratory. An electronic version of each employee’s training records is maintained indefinitely in the Washington State Patrol eTrain program. It is recommended that employees maintain their own training records for career planning, performance appraisals, individual certification, and other uses.

7.11 REQUESTS FOR TRAVEL

Refer to WSP Travel Regulations for information on completing a Training/Travel Request (WSP Form 3000-320-016).

7.12 LABORATORY LIBRARY

Each laboratory will have access to a library containing current books, journals, and reference materials for each discipline. Each analyst is responsible for taking time to read periodicals, journals, articles, books, and laboratory memorandums in order to keep current with information and developments in their respective disciplines. A list of the contents in each library is maintained by the FLSB Librarian. The FLSB Librarian distributes by email the table of contents of various journals, magazines and publications. The FLSB Librarian is a resource for obtaining journal articles and
other needed reference material and should be contacted when necessary. These may also be found on the FLSB Portal.

### 7.13 Courtroom Testimony Training

CLD management is responsible for ensuring that testimony training is provided to employees who testify in court. Topics such as evidence handling, chain of custody, casework results and interpretation of examination or test results should be discussed during the training. Property and Evidence Custodians should also be given courtroom testimony training regarding evidence handling and chain of custody. This training can be given internally by a CLD employee or by an external source. Documentation will be maintained for each individual with their regular training records.

### 7.14 Job Performance

#### 7.14.1 Documenting Job Performance

Supervisors will document the work performance of each employee they supervise and maintain those records in a supervisory desk file. Supervisory desk files will contain positive and/or negative supporting documents, counseling, work directives, evaluations, or records relating to an employee’s job performance throughout the performance period. Supervisory desk files are required to be purged each year following an annual evaluation; training records will be maintained separately.

Employees will have access to and be made aware of the contents of the Supervisory Desk File. (See the Collective Bargaining Agreement.) Annual performance appraisals are required and will be completed for each employee.
8 DOCUMENT CONTROL POLICY AND PROCEDURES

8.1 POLICY

Documents that form the CLD Management System are controlled to ensure that only current, up-to-date documents are being used. All CLD management system documents will be made available to CLD staff via the FLSB Portal. The following procedure provides instructions concerning the creation, revision and distribution of these controlled documents. WSP agency documents are controlled and distributed by the agency.

8.2 DEFINITIONS

8.2.1 Document
A document is any writing or other item that conveys information in any medium including, but not limited to, paper copy, electronic file, audio or videotape, or photograph.

8.2.2 Document Control
Document control is the process for ensuring that controlled documents, including revisions, are reviewed, approved and released by authorized personnel (Issuing Authority), and distributed to personnel performing the prescribed activities.

8.2.3 Controlled Document
A controlled document is any document that forms a part of the CLD management system, and is subject to the requirements of this document control policy. Examples of controlled documents include policy, procedure, technical and training manuals, and required-use forms.

8.2.4 Document Review and Approval (DRA) Form
Document Review and Approval Form, used for all proposed modifications to controlled documents.

8.2.5 Form
A form is a printed, typed or electronic document with blank spaces for insertion of required or requested information. When required for use by policy or procedure, forms will be controlled documents. Adding lines, rows, columns or spaces is allowed as needed, provided the additions maintain the format and integrity of the form (such as retaining column/row headings when adding columns/rows).

8.2.6 Worksheet
A worksheet is a printed, typed or electronic document with blank spaces for insertion of required or requested information. Worksheets may be used to assist in collecting and recording information in casework, and are part of the case record.
When used for convenience but not required, worksheets need not be controlled. When required for use by policy or procedure, worksheets become controlled forms.

8.2.7 Record
Documents, including logs, forms, worksheets and electronic files that provide support of conformity to the quality management system. They may be held in the individual laboratories or at CLD Headquarters. These documents include, but are not limited to, method and equipment validation documents, equipment verification records, reagent and chemical QC logs, training records, proficiency test records, courtroom testimony monitoring records and audit records.

8.2.8 Uncontrolled Copy
A copy of a controlled document provided for informational purposes only. Examples include copies provided to external inspectors or copies required for legal discovery.

8.2.9 Issuing Authority
Personnel that are authorized to approve the posting of controlled documents on the FLSB Portal. The issuing authorities are as follows:

- The Bureau Director for Bureau-wide controlled documents;
- The Assistant Division Commander for the Quality Operations Manual;
- The Quality Assurance Manager for other Division-wide documents;
- The Laboratory Manager for laboratory-specific policies and procedures.

8.2.10 Master Document List
An electronic file maintained by the Quality Process Manager and available to all employees via FLSB Portal which contains the current revision status of any controlled document.

8.3 Procedure

8.3.1 Controlled Document Format
Each controlled document will have the following format requirements:

- A header on each page containing, at a minimum:
  - Washington State Patrol Crime Laboratory Division
- A footer on each page containing at a minimum:
  - Page _ of _
  - A statement indicating that “All Printed Copies are Uncontrolled”
  - The unique document identification
  - Revision number and revision date
- Controlled forms do not require the “All Printed Copies are Uncontrolled” statement, as they are intended to serve as a template for entering data or
information. No modifications to the form, except as noted in the above definition, are allowed without going through the document revision process.

Revision number indicates the total number of times the document has been revised since adoption of original document.

All controlled documents will have a history table indicating when the document was originally adopted and any revisions that have occurred since date of adoption. The table will include the following:
- Document Name
- Revision Number
- Date of Revision

This history table for each controlled document will be maintained on the FLSB Portal and will be updated with each revision. The master document file shall be maintained by the Quality Process Manager and will be available to all employees via the FLSB Portal, listing all the current approved documents.

8.3.2 Controlled Document Preparation

Documents should be prepared by personnel with adequate expertise in the subject. The detail of the document should be commensurate with the complexity of the activity and the background of the intended user of the document. The document must include enough detail and specificity to ensure that the activity conforms to quality specifications and/or expectations.

Recommended changes must represent the objectives of the CLD and not conflict with the WSP Regulation Manual, Revised Code of Washington, or Washington Administrative Code. Laboratory specific policies and procedures cannot supersede the CLD Quality Operations Manual. All affected members of the CLD should have the opportunity to provide input on the proposed changes prior to submission.

The preparer of the new or revised document is responsible for:
- Preparing the document in the proper format
- Acquiring copies of listed references
- Addressing or resolving comments from reviewers
- Submitting for review and approvals using the Document Review and Approval Form.

All proposed revisions must be submitted on form 4002, Document Review and Approval (DRA). The following information must be provided:
- The manual/controlled document name and the specific section of the manual/controlled document to be modified, or the proposed new document or section;
• A statement briefly describing the need for the procedure modification or incorporation of a new procedure.

Proposed changes should be clearly marked and submitted as an edited version tracking all changes made to the current document as follows:
• Deleted portions will have a strikeout
• Additions will be highlighted in yellow
• “Track changes” can be used as an alternative

Amendment of documents by hand pending the electronic re-issue of the revised controlled document is not allowed.

Note: Typographical or grammatical errors do not require a DRA. These types of errors will be brought to the attention of the Quality Process Manager for correction.

8.3.3 Controlled Document Review
Review is required for each new or revised controlled document prior to approval. For DRA’s unrelated to technical manuals/forms, the DRA will be reviewed through the supervisor and laboratory manager. For changes to technical manuals/forms, the Functional Area Supervisors, Technical Leads, DNA Technical Leader and/or Management Liaison will be responsible to ensure that the recommended changes represent the accepted body of scientific knowledge, both internal and external to the CLD. Technical review is for accuracy and clarity. The reviewer(s) must have adequate technical expertise in the discipline to evaluate the document.

A quality review is conducted by the SAS to ensure that the document conforms to accreditation and quality standards.

8.3.4 Controlled Document Approval
Each controlled document issued will be approved through the review process outlined above, with final approval authority by the SAS. All DNA and CODIS documents will be approved through the DNA Technical Leader. Review and approval of controlled documents are recorded on the DRA.

8.3.5 Controlled Document Issue
After the documents are approved, the document will be issued through the SAS. The approved controlled document will be posted on the FLSB Portal and CLD management will be notified by the SAS via e-mail. The notification will include the DRA, effective date, and affected staff. Lab management will be responsible for notifying affected staff and ensuring email acknowledgement of review and receipt is completed. In lieu of email acknowledgement, a Document Control Sheet (DCS) may also be completed. The next revision of the entire manual will include the approved changes from this procedure and will be posted on the FLSB Portal.
Once a document is adopted, it will be the responsibility of laboratory management to ensure it is implemented. Lab managers will be responsible for maintaining the DCS, or email acknowledgement, which will be subject to the audit process.

If a recently approved DRA needs to be altered within a very short time frame after being posted, the SAS can make the necessary changes and document on the approved DRA the sections which were modified. The SAS will be responsible for notifying staff of these changes or if a new DRA is required.

The document history will note the changes as described above and include the date of the entire revision. All personnel will have access to the official electronic documents. However, administrative access to the official electronic controlled documents will be restricted to prohibit unauthorized changes.

8.3.6 Archiving Controlled Documents

Obsolete documents will be archived on the “Archived Manual” section of the FLSB Portal. The date that a document is removed from active status and placed into “archived manuals” will be recorded on the specific document. The labeling of a document with this “archived date” will be used to denote that a document has been officially placed in archived status.

Employees shall use current versions of approved documents. Invalid or obsolete documents shall be promptly removed from all points of issue or use, or otherwise assured against unintended use; obsolete documents retained for either legal or knowledge preservation purposes shall be suitably marked.

8.3.7 Annual Review of Controlled Documents

Controlled documents will be annually reviewed and revised if needed to ensure they reflect current policies, practices, and technology. The revised documents are subject to the same review, approval, documentation and issuance requirements of the original document as stated above.

Technical leads and supervisors or the DNA Technical Leader will conduct this review for their respective technical and training manuals and forms. The Standards and Accountability Section will review administrative manuals. Laboratory managers will review their individual local lab policies. Documentation of this review will be provided by the reviewer to the QPM, who will record the review on the document history table.

8.3.8 Official Controlled Documents

The official controlled documents to be used by personnel are those posted on the FLSB Portal. All employees will have access to this site. Any copies of documents from this site represent unofficial copies and will be designated as such. The Quality Process Manager or designee will maintain the official controlled documents and archived versions of all controlled documents on the FLSB Portal.
8.3.9 Returned
Any DRAs submitted to the SAS that need to be returned, will be accompanied by a written explanation and/or suggestion for modification.

8.3.10 Tabled
Any DRAs submitted to the SAS that need to be tabled, will be accompanied by a written explanation along with the estimated date for reconsideration if applicable.

8.3.11 Submission of Requests for LIMS Manual Revisions
Recommended changes to the LIMS Manual are made using the same process noted above for document revisions, but are additionally routed through the Administrative Management Liaison.
9 QUALITY SYSTEM RECORDS: ACCESS, FILING, STORAGE, RETENTION AND DISPOSAL

Quality system records are any logs, worksheets, electronic files, spreadsheets or databases that provide documented support of conformity to the Quality Management System. These records include, but are not limited to:

- Method validation documents
- Equipment repair and verification records
- Reagent and chemical QC logs
- Training records
- Proficiency and competency test records
- Courtroom testimony monitoring records
- Chemical inventory records
- Reference collection records
- Audit records
- Laboratory Information Management System(s) (LIMS)

These records are maintained by the CLD staff. Filing, storage and retention of these records are as described below. Access to those records maintained at CLD Headquarters is given to the CLD Commander, the Standards and Accountability Section staff, lab managers, auditors, and supervisors for information pertaining to their respective assignments and responsibilities. Individuals may request copies of quality documentation pertaining to themselves. There may be overlap between records held at CLD Headquarters and the individual labs.

All records shall be legible and shall be stored and retained in such a way that they are:

- readily identified and retrievable
- protected from unauthorized access
- safeguarded against tampering and loss
- for LIMS, operated in an environment that complies with provider or laboratory specifications or, and for non-computerized systems, provides conditions which safeguard the accuracy of manual recording and transcription
- maintained in a manner that ensures the integrity of the data and information
- able to record system failures and allow appropriate immediate and corrective actions, as needed.

Records stored electronically shall be routinely backed up to prevent loss. If a LIMS is managed and maintained off-site or through an external provider, the laboratory
shall ensure that the provider or operator of the system complies with the above requirements.

9.1 **RECORDS FILED, STORED AND RETAINED AT CLD HEADQUARTERS BY THE STANDARDS AND ACCOUNTABILITY SECTION**

- Proficiency test answer sheets
- Method validation approvals
- Corrective actions
- Job performance improvement plans
- Policy and Procedure manual document review and approval forms
- Audit records and reports
- Laboratory safety inspection reports
- Official electronic controlled documents

9.2 **RECORDS FILED, STORED AND RETAINED AT CLD LABORATORIES**

- Appropriate personnel records
- Equipment performance verification, associated equipment method validation and maintenance records will be maintained at the laboratory in close proximity to the equipment (on-site),
- Case files and records, and any associated examination or administrative documentation according to retention schedules
- Records on deviations from procedure
- Chemical and reagent quality control logs and worksheets
- Temperature logs and other Quality Control documentation
- Standards and reference collection inventory records and verification logs
- Chemical Inventory databases
- Key control records
- Equipment Inventory
- Lab facility maintenance and security records and logs
- Safety records
- Training completion records
- Courtroom testimony monitoring records
- Visitor logs

9.3 **RECORDS MAINTAINED IN DIVISION-WIDE DATABASES/SPREADSHEETS**

- Laboratory Library Collection
- Firearms Reference Collection database
- Laboratory Information System (LIMS)
- Quality Process Improvement Tracker (QPIT)
- CLD Vehicle Maintenance Records (maintained in Remedy)
9.4 **ARCHIVE AND RETENTION OF QUALITY SYSTEM RECORDS**

Retention and disposal of quality records will follow the Washington State Patrol Archive Record Retention Schedule through at least one cycle of accreditation (four years). A current copy of the Archive Retention Schedule may be found on the FLSB Portal.

9.5 **CASE RECORDS: FILING, STORAGE, ACCESS, RETENTION AND DISPOSAL**

The laboratory will maintain all original case documentation (administrative and examination) in files bearing unique laboratory case numbers. All records shall be legible.

If an original record, paper or other media is captured as an electronic record, and the original record will be destroyed, the laboratory staff shall ensure that the electronic record is complete prior to destruction of the original record.

9.6 **CUSTOMER COMMUNICATIONS**

All communications with customers that form the basis for decisions in casework will be documented in the case record either in the examination documentation or in LIMS (case information field). The date, name of contact, name of laboratory staff, and the conversation (in substance) will be recorded.

9.7 **STORAGE AND ACCESS**

Case records will be stored in a manner that they are readily retrievable and protected from damage, deterioration or loss. Case records may be retained in the case folder as part of the case file, may be in a reagent or instrument log, or may be retained electronically (e.g., LIMS information and digital images). Upon completion, all case files will be stored only in designated areas. No completed case file will be kept in the possession of scientists unless replaced by a sign-out file locator. If case files are removed from the laboratory, for example when going to court, due diligence and caution will be exercised to preserve and protect the file and its contents. The laboratory will protect and make back-ups of case records stored electronically and will secure such records to prevent unauthorized access to or amendment of these case records.

9.8 **ARCHIVING**

All case files will be maintained under the control of the CLD until they are archived. Each laboratory will maintain a minimum of the previous two years plus current year of case files. Case files older than five years should be sent to the State Records Center for secure storage, unless there are reasons to retain the case files.
in the lab. Each laboratory maintains records of files that are stored at the State Records Center.

Prior to archival at State Records, the contents of case files will be secured against loss.

9.9 Expungement of Records

On receipt of a court order for expungement, the Division Secretary at CLD Headquarters should be contacted. Division staff will make any appropriate contacts with the WSP Risk Management Division who will provide guidance to the laboratory for conformance with the order.

For expungement of a convicted offender DNA sample, the CODIS Manager will follow the protocol in the CODIS Manual.

9.10 Laboratory Information Management System (LIMS)

The Laboratory Information Management System (LIMS) maintains the chain of custody and provides meaningful information and statistics on the type and number of requests received, on case turnaround time, and case work backlogs that will assist laboratory management in evaluating Division and laboratory objectives.

In addition, there are a series of reports designed to assist both users and management staff in the performance of their duties. Information and instructions may be found in either the LIMS Procedures Manual or in the on-line Help file of LIMS.
10 CASE MANAGEMENT

The CLD strives to complete the analysis of submitted evidence accurately and within the time constraints required by the submitting agency. The analytical process must not be open-ended. Supervisors will be responsible for monitoring the status of all cases in their section. The laboratory should inform the customer of any major delays in analyses. If there is not an expectation that evidence will be needed for examination/comparison within a reasonable time frame, the evidence should be sent back to the submitting agency until a mutually agreed upon time can be established for resubmission, if necessary. If for any reason the crime lab is not able to honor a request, the customer will be notified. Documentation of this communication must be entered in the case file.

10.1 CASE RECORDS AND DOCUMENTATION

10.1.1 Policy

Case Records (i.e., all administrative and examination documentation) will be identifiable, accessible to authorized personnel and properly stored to prevent damage or loss. Technical records, which exclude administrative documentation, will contain sufficient information to facilitate, if possible, identification of factors affecting the measurement result and its associated measurement uncertainty and enable the test to be repeated under conditions as close as possible to the original. Case records will include the date and the identity of personnel responsible for the sampling, performance of each test and reviewing and reporting of results.

10.2 PROCEDURE

10.2.1 Administrative Documentation

Administrative documentation must bear the laboratory case number and initials or signature in order to be placed back into a case file if it becomes separated. If signatures are present, initials are not required. It is the responsibility of the staff member adding the administrative documentation to the case file to ensure it is properly initialed or signed and has the correct lab number. Page numbers are not required for this documentation; however nothing in this section precludes the optional use of page numbering if desired. If the administrative documentation is a packet of material (such as an officer’s report) that is securely stapled together, the case number and initials or signature only need to be on the first page.

Examples of administrative documentation include:

- Request for Laboratory Examination (RFLE)
- Cover letters, officer’s reports, and other information relevant to the case
- Evidence transaction records including any submitting agency forms used in lieu of the RFLE for documenting the chain of custody
- LIMS printouts
• Court orders

10.2.2 Examination Documentation

Examination documentation must support the conclusions stated in the laboratory report. Observations, data and calculations shall be recorded at the time they are made and shall be identifiable to the specific task. Nothing in the examination documentation may be erased, made illegible or obliterated. In the case of records stored electronically, equivalent measures shall be taken to avoid loss or change of original data. Examination documentation will be permanent in nature and recorded using ink unless the use of ink is made impractical by environmental conditions such as cold or rain. If notes are recorded in pencil, they must be duplicated in ink as soon as possible. Pencil is appropriate for diagramming or making tracings. Pages of notes that will be destroyed due to chemical or biological contamination can occur only after the pages have been photographed or otherwise reproduced for preservation, and the reproduced record is reviewed and complete prior to destruction of the original record.

Changes, additions, or any other form of alteration must be initialed by the person making the alteration. When mistakes occur in examination documentation, each mistake shall be crossed out and the correct value entered alongside. Note that contemporaneous revisions are not considered amendments (see amended reports). If an observation, data, calculation or test result is rejected, the reason, the identity of the individual(s) taking the action and the date shall be recorded in the technical record. Selection of a stronger or clearer test result or observation from replicate test results or observations is not a rejection of the other test results or observations. If examination documentation is created electronically, the documentation will be considered completed and “stored” when the draft report is prepared and before the case file is technically reviewed. If changes to the electronic notes or documentation are required after being initially stored, the original will be “archived” in the electronic case file and the updated version renamed and saved.

Examination documentation must include:
- Unique laboratory numbers
- Handwritten/digital initials or signature of the examiner
- Dates of examination
- Page numbers

The unique identifier for each case for which data is generated shall be appropriately recorded on the printout when data from multiple cases is recorded on a single printout.

The page numbering system:
• Must readily show if a page of documentation is missing
• Will show the individual page number on each page and the total number of pages on the first page
• Will have any double-sided page treated as a separate page and numbered separately
• Will be numeric, except for additions that occur after the document was originally numbered. These additions will be expressed by a letter (a, b, c upper or lower case) after the page number and the additions will be noted along with the total number of pages on the first page.

The start and end dates of testing must be documented in the case record to allow for traceability of materials used in analysis. The start date of testing is documented as the date examination documentation begins and the end date is indicated by the draft complete milestone in LIMS. For CODIS, the end date is the “Analysis Complete” date which is documented on the last page of the case file.

Abbreviations are acceptable if they are readily comprehensible to a reviewer. Where abbreviations or symbols specific to the CLD are used, the meaning of the abbreviations or symbols shall be defined.

Examination documentation includes but is not limited to the following:
• Descriptive information pertaining to the case and the evidence:
  o case numbers
  o evidence descriptions (item numbers, packaging, seals, quantity, size, physical appearance, condition, and adhering materials)
  o currency denomination and count
• Photographs, digital images, and diagrams
• Discrepancies between evidence listed on the request form or packaging and the actual evidence
• If different from the report letterhead name and address, the location of testing, including when performed at a customer facility or at sites away from the laboratory’s permanent facilities, or in associated temporary or mobile facilities
• Examination procedures used and the parameters for those procedures
• Records of data; results of examinations; handwritten or machine-generated notes, forms, and observations; chromatograms, spectra, and other instrumental printouts; photographs, drawings, and other illustrations; identity and source of any standards or references used.
• Test results, conclusions, opinions and interpretations
• Notations showing the generation (and any additional details as necessary) of new evidence items such as trace collections, substrate controls, etc.
• When analytical equipment or instrumentation is used, the procedure and/or any specific operating parameters and the instrument used must be noted in the case record. This will be addressed in the specific functional area.
technical manual. In a laboratory with only one instrument for a specific test or procedure, the instrument’s identification is documented in the laboratory’s equipment list. In laboratories or units that have multiple instruments of the same make/model, one must record the unique identifier of the instrument used in the examination documentation or on the hard copy instrument data.

- Where appropriate, diagrams and/or photographs should be used in addition to narratives, to record observations. Photocopies or printouts of digital images may be suitable in some instances (i.e., thin layer chromatography, questioned documents, etc.)
- Observations, data and calculations must be recorded at the time they are made, and must be identified to the specific analysis or test
- Notes are intended to record observations made during examination. If it is necessary to rewrite notes to make them more legible, the original notes must be retained in the case file
- If a sampling plan is used, reference must be made to the sampling plan in the examination notes. Notes must include the date(s) sampling was performed, location of the sampling, clear indication of what evidence was sampled (item numbers), environmental conditions that may affect sampling
- Documentation to support conclusions shall be such that in the absence of the analyst, another competent analyst or supervisor could evaluate what was done and interpret the data

10.3 Digital Images in Casework Documentation

This section covers digital imaging used for the purpose of documenting casework: it does not address digital images that are considered to be evidence, which is covered under the Evidence section of this manual. Image capture devices should be capable of rendering an accurate representation of the item of interest. Different applications will dictate different levels of resolution for both the image capture and final output. All equipment should be maintained according to the manufacturer’s specifications.

Output devices should be capable of producing accurate representations of input images. The levels of resolution for printed or transmitted images will depend upon the application and need.

Original images will be stored and preserved in a native file format. Duplicates or copies will be used for working images when image processing is required.

Digital images must be saved utilizing a secure agency approved server (typically the authoring lab's designated storage location) that is automatically backed-up, or an agency approved CDR/DVDR/USB flash drive that must be retained in the casefile.
If the images are stored on an external hard drive or secure server, the parent folder will contain a unique identifier (typically the case number and request number) to readily enable locating. If something other than the case number and request number is chosen, this will be documented in the case file.

10.4 **LABORATORY REPORTS**

10.4.1 **Policy**

All casework results and conclusions will be documented in original written reports that are printed on departmental letterhead and signed by the scientist who analyzed, reviewed and authorized test results. The laboratory report is written for the laboratory’s customers as a response for a request(s) for service. Scientists will ensure that the results of each test performed for the request will be reported accurately, clearly, unambiguously and objectively and will include all the information requested by the customer, any information necessary for interpretation of the test results, and all information required to be included by the method used.

10.4.2 **Procedure**

10.4.2.1 **Laboratory Report Format**

The report should be as brief and clear as possible to facilitate understanding by the customer and adhere to the following format:

- Overview [optional]
- Results and Conclusions [May include evidence listing for some seized drug cases]
- Evidence [if applicable, and optional for some seized drug cases if listed under Results and Conclusions]
- Methods and Observations
- Remarks [Optional]

10.4.2.2 **Laboratory Report Content**

Each laboratory report shall include at least the following information, unless the laboratory has valid, documented reasons for not doing so (such as a legal requirement). Any additional information pertaining to the case and the tests performed not specified below will be maintained in the case record as it is not practical to include all the case related information in the laboratory report. When agreed with the customer through signature of the RFLE, the results may be reported in a simplified way described to the customer. Any information not listed in the laboratory report, including the date(s) of performance of specific laboratory activities, shall be readily available to the customer upon request.

- A title, Crime Laboratory Report
• The name and address of the laboratory
• If different from the report letterhead name and address, the location of performance of the laboratory activities, including when performed at a customer facility or at sites away from the laboratory’s permanent facilities, or in associated temporary or mobile facilities (e.g. WSP Crime Laboratory Seattle, University of Washington – Tacoma)
• The Lab number unique to this report, on each page or component of the report
• Clear identification of the end of the report (typically by including the page number and total number of pages)
• The request number(s) associated with the Lab number for the work covered in the report
• Submitting agency name and representative
• A reference to the analytical methods used
• Description of the sealed state of the evidence as received by the examiner
• A description, unique identifier, and when necessary for the interpretation of test results or when tests are adversely affected, the condition of the items analyzed
• Disposition of all evidence items created, recovered, retained or consumed by the analyst
• Reference to other items of evidence received by the analyst but not examined (if applicable)
• Results of analyses including the units of measurement where appropriate
• Any deviations, additions or exclusions from the procedure and information on specific test conditions when necessary
• Where necessary for the interpretation of the test results, a statement on the estimated measurement uncertainty; information on uncertainty is needed in lab reports when it is relevant to the validity or application of the test results, when a customer’s instruction so requires, or when the uncertainty affects compliance to a specification limit stated by a regulatory body, a statute, case law, or other legal requirement
• If applicable, description of any appendices to the report
• Results of tests performed by subcontractors will be clearly identified in the report
• If applicable, initial database entries (e.g., CODIS, ABIS, NIBIN)
• Reporting of associations resulting from a database search
• If applicable, the extent of database (e.g., CODIS, ABIS, NIBIN) searches, unless otherwise communicated to customers in the Forensic Services Guide
• When associations are made, the significance of the association shall be communicated clearly and qualified properly in the report (see discipline technical procedures for how to properly qualify the significance of associations)
• When comparative examinations result in the exclusion of an individual or object, the report shall clearly communicate the exclusion
• When no definitive conclusions can be reached, the test report shall clearly communicate the reason(s)
• A signature block including name and title of the person who analyzed, reviewed and authorized test results. The signature block for seized drug reports shall comply with all current applicable Washington State court rules, as currently found in CrRLJ 6.13(b). The analyst’s email address may be included as part of the contact information at the analyst’s discretion.
• Any analyst not signing the report, but performing a testing activity including, but not limited to evidence screening, shall be identified in the report (in the Remarks section for example). This does not include an additional analyst being involved in part of a batch/team approach such as loading an instrument, starting a batch run, or moving samples between analysis steps.

10.4.2.3 Additional Laboratory Report Content
Where the following is critical to the validity, application or interpretation of the test results, these additional items will also be reported:

• Date of receipt of the evidence item
• Where applicable, reference to the sampling plan and sampling method used (see below for additional sampling reporting)
• Where applicable, a disclaimer indicating which results may be affected by a customer-acknowledged deviation of the evidence items
• Where applicable, identification of data provided by a customer that was used and could affect the test results
• Information on specific test conditions, such as environmental conditions;
• Where relevant, a statement of conformity with requirements or specifications
• Where appropriate, opinions and interpretations
• Where appropriate, assumptions and limitations
• Any additional information that may be required by specific methods, authorities, customers or groups of customers.

10.4.2.4 Reporting of Opinions and Interpretations
The laboratory shall ensure that only personnel authorized for the reporting of opinions and interpretations report and authorize their release. The basis upon which the opinions and interpretations have been made must be documented in the examination documentation. The opinions and interpretations expressed in reports shall be clearly identified as such and shall be based on the results obtained from the tested items. For example:
• “Identification is the opinion of an examiner that there is sufficient quality and quantity of detail in agreement to conclude that two impressions originated from the same source.”
• “This report contains the opinions and interpretations of the analyst whose signature appears on the report.”

Any assumptions or limitations, and where necessary, their significance, that are critical to the validity and understanding of the opinion or interpretation must be clearly stated in the report.

Within the CLD accredited disciplines of forensic science, it is possible to see at least four categories of laboratory reports:

1. Reports which contain only test results:
   For the purposes of this discussion and interpretation, a “test result” is generally one generated without human intervention to evaluate test data. No conclusion, opinion or interpretation is offered as to what the test means or may mean.
   
   Example: A search of the hard drive failed to locate any occurrence of the phrase “unknown suspect.”

2. Reports which contain test results enhanced with the analyst’s or examiner’s conclusion, opinion or interpretation as to what the test result means or may mean:

   Example: The markings on the questioned bullet (Q1) were consistent with the markings on the bullet (K1) fired from the weapon (K2) submitted as Item 1. There is sufficient agreement to conclude that Q1 was fired from K2.

3. Reports which contain only a conclusion, opinion or interpretation:

   Example: The latent print removed from the beer can (item 1) was identified as having been made by the same individual who made the known inked left index finger impression bearing the name John Doe (item 2).

   Example: Item 2: White Powder – Identified as heroin

4. Reports which do not contain a test result, conclusion, opinion or interpretation:
   For a variety of reasons, a laboratory may generate a report to close out a submission that a customer no longer needs worked. A simple crime scene
The report may not contain any test results, conclusions, opinions or interpretations.

**Example:** At the request of submitting officer John Doe, all evidence is being returned with no analysis being conducted.

### 10.4.2.5 Laboratory Reporting of Measurement Uncertainty

Where applicable, the measurement uncertainty will be reported and presented in the same unit as that of the measurand or in a term relative to the measurand (e.g. percent) when:

- it is relevant to the validity or application of the test results;
- a customer’s instruction so requires, or
- the measurement uncertainty affects conformity to a specification limit

See also the section on Measurement Uncertainty for additional requirements.

### 10.4.2.6 Laboratory Reporting of Sampling

If sampling is used in testing where it is relevant to the validity or application of the results, the laboratory report shall contain the following where necessary for the interpretation of results:

- Date of sampling
- Unique identification of the item or material sampled (including the name of the manufacturer, the model or type of designation and serial numbers, as appropriate)
- Location of sampling, including any diagrams, sketches or photographs
- Reference to the sampling plan and sampling method
- Information about the sampling plan, including confidence levels and corresponding inference(s) regarding the population
- Details of any environmental conditions during sampling that affect the interpretation of the results
- Information required to evaluate measurement uncertainty for subsequent testing

If applicable, where the laboratory has not been responsible for the sampling stage (e.g. the sample has been provided by the customer, the laboratory report shall state that the results apply to the sample as received.

See also the section on Sampling and Sample Selection.

### 10.4.2.7 Laboratory Reporting of Work Done by Others

Crime Laboratory personnel who issue findings or conclusions, including writing test reports and providing testimony, based on examination records and/or reports
generated by another FLSB forensic scientist(s), may do so only after the referenced report has gone through technical and administrative review. (Police reports or medical examiner/autopsy reports are not subject to Division administrative reviews but these reports should also be reviewed if used as reference documents.) All such findings or conclusions must be based on information contained within the referenced report(s), associated notes within the referenced case file, or further examination(s) by the author. The laboratory case file will contain a copy of the referenced report(s). Relevant pages or material that was referenced in the case file will be documented in the case file with the analyst’s initials.

10.4.2.8 Laboratory Reporting – Cross Referencing Other Cases

Documented permission must be present in the case record to reflect submitting agency approval of crime laboratory evidence use in related cases that have a different submitting agency or the same agency but different agency case number. Permission must be provided by all submitting agencies involved. The permission requirement does not apply to reporting CODIS hits or use of exemplars.

When reporting a comparison of multiple items of evidence or analytical data from two or more different agency case numbers, a separate report will be prepared for each agency case number. This reporting is optional when the comparisons were with exemplars, whether regarded as evidence or non-evidence exemplars. A statement should be included in each report that includes the other Crime Laboratory number, request number, agency name, and agency case number of the related requests. A Justice Trax entry will be made to relate the case to relevant cases. One case file will hold the original case notes. The related case file(s) will include either a copy of the related original case notes or reference to the Crime Lab number and request number that contain the original case notes. Any copied pages must be readily recognizable as a copy.

When reporting a comparison of an item of evidence or data to previously reported testing from another request, the new report should include reference to the relevant previous report by Crime Laboratory number and request number. This reporting is optional when the comparisons were with exemplars, whether regarded as evidence or non-evidence exemplars. For review purposes, the current case file for this comparison must include the pertinent examination documentation from the previous case file. Any copied pages must be recognizable as a copy.

10.4.2.9 Laboratory Reports – Special Circumstances

If analytical work has already been performed and the customer informs the laboratory that a report is not needed, a technically and administratively reviewed report will still be issued.

The CLD does not require a written laboratory report for the database entry of fired cartridge cases into IBIS/NIBIN. Refer to the Firearms and Toolmarks Technical
Procedures Manual for additional details on crime lab reporting of IBIS/NIBIN entries.

Should a submitting agency request a statement of conformity to a specification or standard for the test result (such as from the Washington Administrative Code or Revised Code of Washington), the laboratory shall report on the statement of conformity, such that the statement clearly identifies:

- to which test results the statement of conformity applies
- which specifications, standards or parts thereof are met or not met
- the decision rule applied (unless it is inherent in the requested specification or standard)

Unless inherent in the requested specification or standard, the decision rule selected shall be communicated to, and accepted by the submitting agency. Where necessary, the laboratory shall take into account the level of risk (such as false accept and false reject and statistical assumptions) associated with the decision rule employed, and apply the decision rule. See also section on Measurement Uncertainty.

### 10.4.3 Authorizing Laboratory Reports and Test Results

All completed laboratory reports will be signed on each page by the forensic scientist who analyzed, reviewed and authorized test results. By signing and dating the report, scientists:

- authorize the results, conclusions, opinions and/or interpretations;
- approve release of the report, and certify that:
  - they were authorized and competent in the testing performed in the case;
  - the report is a true and complete account of the testing performed;
  - the results, conclusions, opinions and/or interpretations are supported by the testing;
  - they reviewed the laboratory report;
  - the report has been technically reviewed; and
  - they are qualified, through training and experience, to perform the testing and make the conclusions as detailed in the report.

### 10.4.4 Draft Reports

In preparing a laboratory report, the initial attempt may require modification or multiple draft reports before the analyst and technical reviewer are satisfied that it accurately conveys the work done and the analyst’s conclusions. The document is considered a work in progress until the technical review is completed. A draft report should be clearly marked as a “Draft”; all pages of the draft report will bear
the author’s initials and date. Only the Final Draft Report with both the author’s and the technical reviewer’s initials or signature and date will be retained in the case file.

10.4.5 Releasing the Report

After the technical and administrative review of a report has been completed and documented, an original signed and dated report will be released to the submitting agency. Copies of the report, reproduced in full, may be provided to a prosecuting attorney with jurisdiction and to other parties or by court order. If the report is sent initially by electronic means, the electronic transmission must be followed by a hardcopy original signed report. Upon distribution of the original report, the Distributed milestone in LIMS will be marked. A copy of the original signed report must be kept in the case file.

10.5 REVIEW OF REQUESTS

10.5.1 Policy

The WSP CLD will ensure that the customer’s requirements, including methods to be used, are adequately defined, documented and understood; that the laboratory has the capability and resources to meet the requirements of the request and that the appropriate methods or procedures are selected and capable of meeting the customers’ request requirements. The review of the request will also cover any work that is subcontracted by the laboratory. The WSP CLD will have initial discretion over the selection of methods for analysis, the totality of the analysis and the items tested.

10.5.2 Procedure

A Request For Laboratory Examination (RFLE) form must generally accompany all evidence submissions. A request form may be submitted prior to the submission of the actual evidence, but all evidence received by the laboratory must have an associated request form.

Prior to the start of testing of evidence, CLD personnel will review the evidence and case information against the services requested on the RFLE to confirm that the CLD has the appropriate test methods, capability, and resources to perform the services requested.

The review will be documented by initialing and dating of the request by the supervisor or scientist, by creating documented case communication with the investigator, completion of the request, or by other demonstrable evidence that the review took place within the first 60 days. All accepted requests will be assigned by supervisors or designee as soon as practical. The date that cases are assigned will be documented in LIMS.
Although timeliness is important, quality will not be sacrificed in order to meet a deadline. If meeting a deadline is likely to compromise quality, staff may consider options including transfer of the evidence to another laboratory. If options are not feasible, staff will consult with the submitting agency and/or the prosecuting attorney and advise them the work cannot be satisfactorily completed within the imposed timeframe. If a resolution cannot be reached, this will be communicated up the chain of command as appropriate. The agency contact and resolution communications will be documented in the case record.

Lab managers will work with the supervisors to ensure that good customer service is provided on all unassigned cases. If appropriate, the evidence should be returned and the request cancelled.

After the request has been reviewed, a scientist will inform the customer and notify their supervisor if the CLD is unable to fulfill the request for services and the reason. Any differences between the requested services and the services that the CLD can provide will be resolved before any work commences. In addition, any deviations requested by the customer, if implemented, shall not impact the integrity of the laboratory or the validity of the test results. Documentation of this contact, at a minimum, will be placed in the case info file tab in LIMS. If a requested service is cancelled or not performed, this will additionally be noted on the RFLE. Once the CLD has received and accepted the evidence (through submission on the RFLE), it is the CLD responsibility to determine the best analytical approach for the evidence. It is understood that the most effective use of resources may not allow all evidence items to be examined. For example, if multiple items are submitted in a single-suspect seized drug case, only a single item may be analyzed and the agency would not receive pre-notification beyond what is described in the Forensic Services Guide.

Any subsequent amendments or deviations from the initial agreed upon services will be documented at minimum in the case info file tab in LIMS. The customer and affected personnel will be notified.

### 10.6 REVIEW OF CASEWORK

#### 10.6.1 Policy

Each laboratory will ensure that conclusions are reasonable and supported by the examination documentation and that established policies and procedures are being followed. All laboratory reports and associated case documentation will be subject to technical and administrative reviews.

#### 10.6.2 Analyst Review

Analysts will conduct a thorough review of their own work prior to authorizing technical review. This review is done after all analyses for that request are
complete and the draft report has been printed where applicable. The analyst documents this review by selecting Draft Complete in LIMS and by documenting a completion date in the case file. (For the CODIS Laboratory, this documentation is the “Analysis Complete” date found on the last page of the case file). The analyst review is a complete review of the case file consisting of all the elements of the technical and administrative reviews. Producing quality casework is a normal job function of all scientists qualified to perform casework, and the quality of casework will be documented and evaluated by supervisors. Every effort will be made to prepare a draft report that holds up to technical and administrative scrutiny. The author of the lab report has the primary responsibility to make sure that their draft report is as free from errors as possible and that case documentation is complete and of the highest quality.

The reporting analyst is the person responsible to ensure that the distributed report is an accurate representation of the case results. The analyst’s reported conclusions must:

- be able to stand alone without the necessity of explanation by supporting verbal testimony
- be accurate, thorough, and clearly stated
- be supported by the data
- clearly distinguish fact and inference
- appropriately address the customer’s questions

### 10.6.3 Technical Review

- Technical review will be conducted on all cases before release of written and verbal/email reports. This is to ensure that the results, opinions, interpretations and conclusions stated in the draft report are properly qualified and supported by the case record. The technical review is also performed to ensure examination documentation is complete and accurate and that the final report will be free of omissions and errors. Technical review is a normal job function of all scientists qualified to perform that function, and will therefore be subject to documentation and evaluation by supervisors. While the final responsibility for the scientific findings in the report rests with the analyst, the technical reviewer is equally responsible for the quality of the report and both will be held accountable.

- Assignment of cases for technical review is the responsibility of section supervisors. Technical review is to be conducted by authorized individuals who have been competency tested in the testing being reviewed and who are currently performing casework or have completed proficiency tests in that category of testing within the last four years. For technical review of DNA cases, the technical reviewer must be current with their proficiency testing. Technical reviews shall not be conducted by the author or co-author(s) of the examination documentation or draft report under review.
• The technical review process should be undertaken as soon as practical after the case is completed. Complex or difficult cases may require more time in order to do a thorough review. Supervisors are responsible for ensuring that cases are reviewed in a timely manner.

• The technical reviewer will ensure:
  - Examinations conducted are appropriate to satisfy the request made by the customer
  - Conformance with test methods and applicable policies and procedures
  - If an analysis was not conducted, the reason is supported by established laboratory policy
  - Communications and phone notes are present if applicable
  - All procedures, data, results, conclusions, opinions and interpretations are documented
  - Results, conclusions, opinions and interpretations are accurate, properly qualified and supported by the examination documentation
  - Conclusions are reasonable and stated unambiguously, neither overstating the significance of the findings nor omitting any reasonable conclusion
  - Opinions and interpretations are clearly identified as such, are accurate and properly qualified
  - All relevant case information is included
  - Descriptions of evidence and evidence packaging are complete
  - All calculations and data transfers are verified for accuracy
  - Appropriate procedures were used and test parameters (for example, instrument operating parameters) were appropriate for the examination.
  - Any deviations from established procedures are recorded in the case file, technically justified, authorized, and accepted by the customer.
  - Actions taken when discrepancies are found are described
  - Appropriate standards and controls are used when necessary and documented
  - Other items of evidence received by the analyst but not examined are referenced (if applicable)
  - Generation and disposition of new evidence items such as trace collections, substrate controls, etc., is documented
  - All strikeouts or insertions are noted with the examiner’s initials. Overwrites must be struck-through, rewritten, and initialed. No obliterations should be present.
  - All pages of examination documentation are labeled with the case number, dates, examiner’s handwritten initials, and page
number. The total number of pages of notes is documented on the first page.
- The draft report is clear, concise, and initialed and dated
- The answer sheet for proficiency tests has been fully completed and is free of errors
- Excessive errors or insufficient data to support the conclusion are brought to the attention of the supervisor
- Discipline-specific requirements for technical review are met.

An approved discipline specific technical review checklist will be used to facilitate the review process and be retained in the case record as administrative documentation. If during the technical review process, an observation, data, calculation or test result is rejected, the reason, the identity of the individual(s) taking the action, changes made, and the date shall be tracked by recording in the technical record. Tracking can be accomplished in a variety of ways, including but not limited to noting the changes on the technical review checklist or examination documentation and through document track change functions. The analyst must address all the observations and recommended corrections of the technical reviewer.

### 10.6.3.1 Technical Review Issues

If during the technical review process, there are significant concerns regarding technical or quality issues, such as those listed below, the case file must be turned over to the supervisor.

- The examination documentation does not support the conclusions stated in the report
- The examination documentation is not clear in content, intent, or purpose
- The examination documentation contains procedural errors
- The examination documentation or report exhibits numerous errors not appropriate for the complexity of the case
- The examination documentation contains inappropriate strikeouts, obliterations or overwrite or cut-and-paste errors
- Issues or discrepancies are not successfully resolved

The supervisor will evaluate the concerns and, if appropriate, notify the Laboratory Manager and the Standards and Accountability Manager. If the case involves DNA analysis, the DNA Technical Leader will also be notified (see also the section on Nonconforming Work and Corrective Actions). Substantive nonconformities or recurring nonconformities discovered during technical reviews are to be brought to the attention of the SAS Manager and Quality Assurance Manager through the chain of command as soon as possible. The Corrective Action process will be followed.
Errors discovered after the technical review process may be addressed by Corrective Actions and will involve both the originating scientist/author and the technical reviewer.

10.6.3.2 Documenting Technical Review

Technical Reviews will be documented with the reviewer’s initials and date on each page of the final draft report, and in LIMS. (For the CODIS Laboratory, the reviewer’s initials and date are on the first page of the case file). The presence of the reviewer’s initials indicates that the bench notes, data, spectra, photographs, and other documentation found in the case file clearly support the conclusions stated in the report.

Any alterations made through technical review of the final draft report bearing the analyst and technical reviewer initials/signature shall be crossed out, not erased, made illegible or deleted, and the correct value entered alongside. All such alterations, including adding information, shall be signed or initialed and dated by the person making the alteration, the analyst and by the technical reviewer. The final report must reflect these alterations. If the analyst disagrees with the changes indicated on the altered draft, the report cannot be released and the analyst will need to contact the technical reviewer and resolve the disagreement or follow the mediation procedures described below in section on Resolution of Technical Differences of Opinion.

10.6.4 Inter-lab Technical Review

In some labs in the CLD there are insufficient analysts to conduct technical reviews of all cases. It is therefore necessary to submit case files to other labs for review. In addition, it has been determined that a quality improvement can be obtained by submitting routine casework for review to scientists outside one’s own laboratory. Therefore, the Quality Process Manager, working in conjunction with the functional area supervisors, will develop a schedule for inter-lab technical review.

All cases submitted for inter-lab technical review will be handled using the following guidelines:

- When submitting case files to other laboratories for inter-lab technical review, precautions must be taken to avoid the loss of case file documentation, particularly that which is difficult if not impossible to reproduce. Original case file documentation which cannot be reproduced (i.e. RFLE, case notes) will be retained by the submitting laboratory unless hand delivered and returned; otherwise only copies of this documentation will be sent out for review. Original case file documentation which can be easily reproduced (i.e. instrument data) may be sent out for review and does not need to be hand delivered or photocopied.
- For scheduled inter-lab reviews, rush cases will not be submitted unless both parties agree.
• Supervisors will be responsible for selecting cases for inter-lab technical review and will monitor the program.
• Reviewers will conduct technical reviews in a timely manner. Supervisors will be apprised of any delays.
• Documentation of the technical review will be by dated signature or initials on all pages of the final draft report or photocopy. If a photocopy, the photocopy will be retained in the case file.

### 10.6.5 Administrative Review

An administrative review will be conducted on the case file and final report prior to the release of laboratory reports, amended reports and proficiency test answer sheets. The administrative review is designed to ensure that:

- The report or answer sheets being released correctly and completely reflect the final draft report, including any minor corrections indicated on the final draft report by the reviewers
- The report or answer sheets being released do not contain misspelled words or grammatical errors
- The evidence item numbers and case numbers are correct
- Examination documentation is identified with the case number, initialed, dated, and page numbered
- The total number of examination documentation pages is documented on the first page
- Administrative documentation is identified with the case number and initials/signature
- A technical review has been completed and is documented on the final draft report and where applicable, on the technical review checklist
- Proficiency test files contain a copy of the answer sheet with case numbers, initials and documentation of technical review; administrative review may also be documented on the answer sheets

During administrative review, any suggested alterations to be made on the final draft report bearing the analyst and technical reviewer initials/signature shall be brought to the attention of the analyst. The alterations shall be crossed out and the correct value entered alongside. All such alterations, including adding information, shall be signed or initialed and dated by the person making the alteration (analyst), and if technical in nature, by the technical reviewer.

Administrative reviews will be conducted by technical staff, supervisors or lab managers. The administrative reviewer does not have to be technically proficient in the functional area, but may not be the author of the report. The administrative review is documented in LIMS and may additionally be documented in the case file.
10.6.6 Verification of Physical Comparisons

A verification of physical comparisons is an examination of the evidence to verify another analyst’s conclusions. Those cases where such a verification is required will be identified in the technical manuals for each forensic discipline.

Verifications are conducted by individuals currently authorized to perform casework in the category of testing. Verifications will be a separate process from technical review, but may be conducted by the technical reviewer.

Verifications, including off-site verifications, will be documented in the case file and include the identity of the analyst performing the verification, when it was performed, and the results of the verification. The verifier will provide sufficient documentation to support his/her own independent observations and conclusions. Minimally, if the verifier draws the same conclusion as the primary analyst, documentation must include the verifying analyst’s initials and date of examination. Where applicable, the scanned copy with the verifier’s initials and date will be retained as the original. For situations where the verification does not agree with the original test result, the resolution of any discrepancy shall be recorded. Refer to the section on resolution of technical differences of opinion and nonconforming work, as applicable, for actions to resolve discrepancies.

If an observation, data, calculation or test result is rejected, the reason, the identity of the individual(s) taking the action, changes made, and the date shall be tracked by recording in the technical record. Tracking can be accomplished in a variety of ways, including but not limited to noting the changes on the technical review checklist or examination documentation and through document track change functions.

10.7 Technical Review in Special Situations

10.7.1 Amended Laboratory Reports

Contents of issued laboratory reports may occasionally require modification by the generation of Amended Reports. The report must be titled “Amended Report,” and a brief explanation describing the need for the amendment must be the first sentence of the report. If the amended report revises a result or conclusion, the revised result or conclusion and the reason for the revision must be clearly stated in the amended report. Where additional information is critical to the validity, application, interpretation or understanding of the revised result or conclusion, refer to section 10.4.2 for additional reporting requirements.

All amended reports shall be uniquely identified and shall contain a reference to the original that it replaces. Amended reports will, at a minimum, include the submitting agency and laboratory case numbers, request number of the original report, and the corrected report language.
All amended reports must be documented in the Case Information field in LIMS. A copy of the original signed report and the amended report shall be retained with the case record, including the date of alteration, an indication of the altered aspects and the personnel responsible for the alterations.

If the reason for the amended report is clerical in nature, the supervisor’s review and approval is required. The supervisor will document their review and approval by initialing and dating the draft of the amended report. An example of a clerical error would be a typo (e.g. incorrect case or item number, or word misspelling) that generally does not alter or jeopardize the correct understanding of the reported content, conclusions and results of the forensic testing.

If the reason for the amended report is due to a technical error or oversight, the amended report must be technically reviewed prior to release. The technical reviewer will document their review in the Case Information field in LIMS. The technical error or oversight must be documented and treated as a nonconformance (refer to the section on Nonconforming Work and Corrective Actions). Both the original and amended reports will be attached to the entry in the Quality Process Improvement Tracker (QPIT). Copies of all amended reports involving technical issues will be available to the supervisor. An example of a technical error would be an incorrect conclusion, method, or observation or any error that alters or jeopardizes the correct understanding and interpretation of the reported content, conclusions and results of the forensic testing.

10.7.2 Verbal/Email Reports

When necessary, a verbal/email report may be issued provided that the section supervisor or designee has approved that course of action and that the technical subject matter contained in the verbal/email report has been technically reviewed.

The issuing scientist must first draft a written version of the verbal/email report.

The verbal/email report will be submitted for technical review. The data supporting the verbal/email report will be initialed and dated by the technical reviewer. The technical reviewer will document the technical review by initialing and dating the written version of the verbal/email report. The supervisor and technical reviewer may be the same person provided that the supervisor is qualified in the relevant forensic discipline and is current with the required proficiency tests.

If fully approved through the processes described above, the scientist may release the verbal/email report. The verbal/email report will be documented by making a notation in the Case Information field in LIMS. The technical reviewer will also make a notation documenting their review in the same section in LIMS. The scientist will document on the written version of the verbal/email report the date, time, and name of the person to whom the verbal/email report was released.
written version of the verbal/email report must be maintained as a permanent part of the case file.

In some instances, not all of the casework will have been completed; however, the technical review must be appropriate for the level of completion of the analytical work and documented as such.

Results containing non-comparative information such as semen identification, indications of blood, presence of trace evidence, etc., may be released without technical review.

A verbal/email report must be followed by a formal written report once the casework has been completed.

The final technical reviewer must ensure that the formal written report is consistent with the verbal/email report issued previously. The technical reviewer will notify the supervisor of any inconsistencies or problems.

10.8 Resolution of Technical Differences of Opinion

Disagreements may sometimes arise between scientists during the technical review process. Every effort will be made to resolve these issues at the peer level. Technical reviewers may request changes in draft reports, further work to clarify issues, or further work to complete cases. If there are unresolved differences during the review, the following process will be used:

- The reviewer and the scientist will bring the issue to the attention of their supervisor who will act as a mediator. More than one supervisor, or one selected supervisor, may serve in the mediation.
- If not resolved, the Technical Lead will review the issues and make a recommendation to the supervisor, scientist, technical reviewer, lab manager and the SAS Manager.
- If no agreement is reached, the SAS Manager will then form an arbitration review committee and notify the CLD Commander.
- The arbitration review committee members shall be scientists with independent casework experience in the subject matter, and at least one laboratory manager not associated with the issue.
- Recommendations by the committee must be documented and may include re-analysis, additional work, revised draft report wording, or other suitable action. The decision of the review committee concerning the resolution of the case shall be binding.
- NOTE: For cases involving DNA analysis, if mediation is not resolved at the supervisor level then a second mediation is to be done by the DNA Technical Leader or by a qualified DNA analyst approved by the DNA Technical Leader. If differences exist after this mediation, the DNA Technical Leader, in
consultation with the SAS Manager, is empowered either to arbitrate the issue(s) or decide that a review committee should arbitrate. The supervisor(s) and lab manager will be notified when mediation or arbitration is necessary, and the result of the process.

- The resolution will be recorded in the case file and concluded prior to the release and distribution of the laboratory report.

10.9 **FOCUSED CASEWORK REVIEW**

When quality processes uncover significant errors in casework, or there is a complaint alleging misconduct or incompetence, the CLD Commander or FLSB Director may initiate a focused casework review. If a root cause analysis has been completed, the CLD Commander or FLSB Director will review the analysis and its recommendations and any other input from the SAS Manager as part of their deliberation as to the necessity of a focused casework review.

10.9.1 **Review of Affected Cases**

The focused casework review will be conducted by an appropriate scientist or panel of scientists chosen by the CLD Commander. The reviewing scientist(s) will prepare an IOC report summarizing the findings and forward the report to the SAS Manager who will review and discuss with the CLD Commander or FLSB Director.

10.9.2 **Notifications**

The SAS Manager or designee will notify ANAB within 30 days of the initiation of a focused casework review.

The FLSB Director will notify the Forensic Investigations Council within 30 days of the focused casework review.

Disclosures to prosecuting attorneys and the judiciary will be in accordance with WSP Regulation Manual section 6.01.065.

10.9.3 **Removal from and reinstatement to Casework**

The scientist who is under a focused casework review will be removed from casework by the CLD Commander until the matter is resolved as required by the section on Nonconforming Work and Corrective Actions. In addition to the fact finding, technical review, re-examination of evidence, or other action taken by laboratory management, amended laboratory reports may be issued in the affected cases to the submitting agency with copies to the prosecuting attorney’s office. Reinstatement to casework will also be by the CLD Commander.
10.10 Courtroom Testimony

Testimony should be limited to the results of the staff member’s direct work on the case in question, direct knowledge of the case events or an area of their expertise. Most often requests for appearance will be through a subpoena. Every effort should be made to comply with requests for appearance regardless of whether a subpoena is received as this is the legal culmination of our laboratory analysis.

Subpoenas received that pose a scheduling conflict must be resolved. Resolution is generally done via conversations between the staff member and the person issuing the subpoena.

10.11 Court Testimony Review

10.11.1 Policy

The testimony of each staff member, and former employees that provide testimony while under contract with the CLD, must be reviewed annually and technically reviewed at least once during an accreditation cycle. The laboratory must provide clear documentation identifying individuals who did not testify over the course of the year. For scientists authorized in multiple disciplines, technical review of testimony shall be performed for all disciplines in which an analyst is authorized to conduct casework, and for which they testify, during an accreditation cycle.

Mock trials/moot courts are not a substitute for actual testimony for this review requirement.

10.11.2 Procedure

10.11.2.1 Court Testimony Review Methods

Testimony review methods include:

- Direct observation of the testimony by a qualified individual;
- Review of a video recording of the testimony;
- Use of a video conferencing system to observe the testimony “live”;
- Review of an audio recording;
- Review of testimony transcripts;
- Solicitation by a laboratory manager or supervisor to one or more officers of the court for evaluation of the testimony.

10.11.2.2 Requirements

Prior to going to court to testify it is the duty of the staff member to inform their supervisor. After testifying, each person is responsible for entering the time spent related to testimony into the Activities Section of LIMS, including travel and preparation time.
10.11.2.3 Technical Reviewer Requirements

Technical review of testimonies is to be conducted by authorized individuals who have been competency tested in the testing being testified to, and who are currently performing casework, or have completed proficiency tests in that category of testing, within the last two years. The technical reviewer must be a Forensic Scientist 3 or higher.

10.11.2.4 Supervisor Requirements

If the testimony was directly observed, the testifying staff member will be given feedback through their supervisor and/or the reviewer on the positive aspects of the testimony as well as the areas that may need improvement. If a court testimony was not directly observed, the supervisor may consult with an officer of the court who was present for feedback on the testimony, except for technical reviews. Information received by the reviewer in the form of transcripts, audio or video will be shared with the staff member.

Time spent by the supervisor and/or testimony reviewer in monitoring the testimony will be entered into the Activities Section of LIMS for the lab number addressed by that testimony.

Testimony review will be documented on the court testimony evaluation form and placed in the individual’s supervisory desk file. The supervisor and/or designee will discuss the assessment with the staff member; the staff member and supervisor/designee will sign and date the form. Any problems identified from the review of testimony will be addressed by the supervisor and documented in the supervisory desk file. (See section on Nonconforming Work and Corrective Actions)

NOTE: Copies of the court testimony evaluation form shall be retained for at least one accreditation cycle.

10.11.2.5 Laboratory Manager Requirements

Provided that testimony occurred, it is the responsibility of the Laboratory Managers to ensure that testimony of all staff members be reviewed and documented annually and technically reviewed at least once in an accreditation cycle. Laboratory Managers will also maintain testimony review records for former CLD employees who provide testimony while under contract with CLD.

10.11.2.6 Evaluation Criteria for Acceptable Performance

Evaluation criteria will include:

- Effective communication skills, clarity, appropriate eye contact
- Appropriate, cooperative and professional demeanor
- Objectivity, impartiality
- Professional appearance
- Being well organized, prepared, familiar with the case file and on time
• Sound technical knowledge
• Technical accuracy and clarity
• Results, opinions and interpretations are accurate, properly qualified and supported by the technical record
• Other relevant observations

10.12 DISCLOSURE AND RELEASE OF INFORMATION

10.12.1 Policy

The CLD is required by law to disclose documentation and information when it is requested by the media, defense counsel, or other parties designated by the Public Records Act. These inquiries typically involve requests for forensic test results, test methods, and/or validation data.

All employees are responsible for maintaining confidentiality in all cases. It is the policy of the CLD to follow the WSP Public Disclosure procedures and to work with customers to honor requests.

10.12.2 Procedure

Discovery requests from the prosecutor will be referred to the appropriate scientist. The section supervisor will be informed of all discovery or public disclosure requests for their section. If appropriate, the supervisor may notify the Laboratory Manager (e.g. for excessive requests, requests for staff DNA profiles). Discovery requests are usually received from the prosecutor's office and can be handled by the assigned scientist by providing the requested documents. In most instances, a copy of the report and all requested examination documentation will be transmitted as soon as possible through the prosecuting attorney’s office unless specifically ordered otherwise by the court or authorized by the prosecuting attorney.

If requested, these items can be provided via discovery request(s):

Note: The items listed below are not all-inclusive.

• Copy(ies) of the case file including electronic data
• Quality assurance records directly related to the requested case (quality variance entries, contamination log entries, corrective actions)
• Evidence Chain of Custody
• Proficiency test evaluation forms for up to two years
• Reporting scientist and technical reviewer CV’s
• Copy of laboratory accreditation certificate and scope of testing
• Summary audit reports
• Current CLD manuals are available on the WSP external website. Archived versions of CLD manuals are on the FLSB Portal
• Copies of manufacturer product inserts
Copies of internal validation (summaries only)

Requests for information received from anyone other than the Prosecutor’s Office will be coordinated and tracked through the Public Disclosure Tracking Coordinator (PDTC).

For a subpoena duces tecum, the scientist/individual served will produce the required case records and forward to the PDTC. The PDTC will track the request, provide the requested records in response to the subpoena duces tecum, and notify the served individual when the records have been provided.

Requests for records kept outside the CLD (HRD, OPS, etc.) will be forwarded to the PDTC for response. Court orders for non-casework laboratory documentation will be forwarded to the PDTC.

If requested, the following items can be made available for on-site (in lab) inspection. Pertinent pages viewed at the lab may be made available to the requestor by digital scan or photocopy. If the request is unduly burdensome, the analyst’s supervisor and the prosecutor will be contacted.

- Original case file can be viewed with the exception of any "no match" (see below) candidate match paperwork. Documentation associated with an elimination DNA profile may only be viewed if a court order providing protection of the elimination DNA profile and/or name has been received.
- Quality assurance records: quality variance logs, contamination logs, corrective actions, and detailed audit documentation and responses beyond those that are directly related to the specified case. The specific area of interest shall be requested in writing. Only entries in the log(s) dating from three months before the date of testing of the samples to three months after may be viewed.
- WSP Validation notes, electronic data, and data analysis tables
- Analyst training records
- Proficiency test files, electronic data and answer sheets
- Instrument and equipment maintenance and calibration records
- Quality Control testing data for reagents and chemicals

For case files involving "No Match" candidate paperwork or staff/elimination DNA profiles:

- If the case file contains "no match" candidate match paperwork, the "no match" candidate profile shall be redacted by the PDTC.
- If the case file contains a match to an elimination DNA profile belonging to a past or present employee, the case file may be sent with the past or present
employee’s DNA profile redacted by the PDTC. If the requestor requires the redacted information then a court order providing protection of the elimination DNA profile shall be sought prior to release of the documentation.

- If the case file contains a match to an elimination DNA profile, not belonging to a past or present employee, the case file may be sent with the elimination DNA profile, but the name of the person redacted by the PDTC. If the requestor requires the redacted information then a court order providing protection of the elimination DNA profile and the name of the contributor shall be sought prior to release of the documentation.
- If a case file request requires redaction of "no match" candidate paperwork or staff/elimination names and/or profiles, the following steps will be followed:
  - The scientist shall notify their supervisor, lab manager, PDTC, and CODIS Manager of the discovery request. The DNA Technical Leader and DNA Operations Manager can also be notified if appropriate.
  - The lab manager or CODIS Manager (or designee) shall consult with the Assistant Attorney General as soon as practicable.
  - Scientist shall notify the prosecutor and defense attorney that a copy of the redacted information will be released by the PDTC upon receipt of a court order providing protection of the elimination DNA profile and/or name.
  - If the elimination DNA profile is from a current WSP employee, the lab manager (or designee) shall provide notice to the affected employee prior to the release of their DNA profile.
  - Documentation associated with the release of a match to an elimination DNA profile (e.g. protective court order, notice to employee) shall be retained in the case file.

The case record will contain a record of the date the request was received, the date(s) the information was provided, and the records and information provided. If the request is excessive, the scientist involved should immediately notify their supervisor and mitigation will be sought. The assigned prosecuting attorney should be consulted first; the WSP Public Records Officer and the WSP representative in the Attorney General’s Office (AGO) may also be valuable resources. Any contacts with the AGO will be routed through the chain of command to the CLD Commander or designee.

The prosecuting attorney and/or defense counsel may request a pre-trial conference with a scientist to discuss a particular case. Scientists should participate in trial preparation with attorneys, whether in face-to-face meetings or by teleconference.
10.12.3 Media Contacts

The CLD recognizes the need for a positive and open relationship with the media by maintaining a Public Information Officer (PIO). The CLD PIO is the QPM unless otherwise designated by the CLD Commander.

Media requests for information will be directed to the Division PIO who will notify the CLD Commander as appropriate. The Laboratory Managers or their designees will also be notified of such requests.
11 EVIDENCE MANAGEMENT

11.1 DIVISION POLICY

The procedures used for the transportation, receipt, handling, protection, storage, retention, disposal or return, and examination of evidence must be designed and carried out to protect the integrity of evidence, and to protect the interests of the laboratory and the customer. Precautions must be taken to avoid evidence deterioration, loss, cross contamination or deleterious change during handling, transporting, storing/waiting, and preparation for testing. Care and handling instructions to preserve the evidence provided shall be followed.

Laboratory staff will use a best practice approach when selecting examination order and techniques so that evidence is not compromised or unnecessarily consumed. All employees will share in the responsibility of ensuring that evidence is not lost, contaminated, or cross-transferred. Universal precautions to prevent contamination will be used when necessary.

11.2 DEFINITIONS

11.2.1 Chain of Custody

Documentation demonstrating the receipt of, internal transfers, and the return of evidence to the submitting agency by the CLD laboratories.

11.2.2 Convenience Packaging

Convenience packaging is defined as a container used primarily to aid in the transport of the evidentiary items contained within. Convenience packaging will not have an evidence seal.

11.2.3 Evidence Packaging

Evidence packaging is defined as packaging that contains an evidence seal.

11.2.4 Evidence

A physical object, material or test item believed to have some investigative or forensic significance and defined as such by law enforcement personnel or forensic analysts.

- Items created from evidence that cannot be reproduced are generally considered evidence. When evidence, such as latent prints and impressions, can only be recorded or collected by digital imaging or photography and the impression itself is not recoverable, the digital image, photograph or negative of the image shall be treated as evidence.
- Digital images captured at crime scenes are considered evidence.
- Test fires submitted or created by laboratory personnel for comparison and/or IBIS entry are treated as evidence. Test fires submitted by user
agencies for NIBIN entry only, or brought in as “walk ins,” are considered exemplars and are not treated as evidence. Refer to the Firearms and Toolmarks Technical Procedures Manual.

- DNA samples (e.g. extracts produced in the lab), bloodstain cards (e.g. stains for DNA analysis made from reference samples) and microscope slides for DNA analysis are not considered to be evidence.
- Convicted offender DNA samples submitted to the CODIS Laboratory are not considered evidence.
- Items that are created from evidence by an analyst, and can be reproduced, are not considered evidence (i.e. photographic prints made from negatives, extracts where additional material is available to reproduce the extract, etc.).

Individual technical manuals will address specific issues for DNA analysis work products and items created from evidence that can be reproduced.

11.2.5 Evidence Seal
An evidence seal is a device or material that is used to close or fasten an opening or connection in order to protect evidence from loss, cross contamination or deleterious change.

11.2.6 Exemplar Materials
Exemplar materials are representations illustrating the class and individual characteristics of a known object or source within a specific context. Exemplars are created in order to allow a comparison between the known object or source and a questioned item of evidence. Exemplars may or may not be considered to be evidence.

11.2.7 Final Disposition
Final Disposition is the last step in the chain of custody. This is most commonly the return of an item of evidence to the submitting agency. However, there must be documentation in the case record if the item is to be retained in the laboratory or if it is to be destroyed.

11.2.8 Laboratory Information Management System (LIMS)
LIMS documents the official chain of custody for evidence submitted to the laboratories. JusticeTrax LIMS-Plus is the CLD’s Laboratory Information Management System which is used for tracking cases and evidence, and for generating analytical and statistical reports. LIMS is a secure electronic database and requires a user identification and password. A full description of LIMS is contained in the LIMS Operations Manual.

11.2.9 Limited Sample
A limited sample is one that is likely to be completely consumed during analysis.
11.2.10 Request for Laboratory Examination (RFLE)
The form (Form 3000-210-005) used by the submitting agency to formally request forensic services from the CLD.

11.2.11 Submission
Items received as evidence for a laboratory analysis on one or more RFLEs. (See LIMS Operations Manual.)

11.2.12 Transferred Case
Evidence in a case sent to another laboratory for analysis.

11.3 Chain of Custody

The CLD shall document a chain of custody on all evidence received from time of initial submission of evidence to time of evidence return to agencies, and on items that are collected or created and preserved for future testing that are considered to be evidence. The chain of custody record shall document all internal transfers indicating each person taking possession of an item of evidence or the location of the evidence.

The chain of custody must be maintained at all times and shall securely and accurately identify:

- the individual(s) or location(s) receiving or transferring the item(s);
- all evidence items transferred, received and handled by the laboratory, including those items not tested; and
- the chronological order of transfers, minimally including the date.

Each laboratory will use a secure electronic chain of custody record through the use of the LIMS. This electronic record is the official chain of custody. However, the initial chain of custody is also documented on the RFLE. Additional chain of custody documentation is allowed; however, all evidence transfers must be entered into LIMS. EXCEPTION: Crime Scene Response – CSRT documents evidence collection on the CSR Evidence Inventory Log (Form CSR-EIL-11003) for release to the investigating agency at the crime scene. This evidence transfer will be documented on the form by both the Crime Scene Responder and a representative from the investigating agency.

11.4 Evidence Marking and Sealing

(See also the Forensic Services Guide.)
All evidence is authenticated by marking it with a unique case number and a unique item number. These identifiers must be on the evidence packaging or on the evidence item itself if the item is unpackaged. Laboratory staff will use care when marking items to minimize devaluing or defacing them.

Evidence can be protected from loss, cross contamination or deleterious change by sealing the evidence in a container using tamper-indicating tape or similar device. If the evidence needs to be sealed in a container in order to protect it, all of the openings in the container must be sealed.

Staples are not considered tamper proof and do not constitute a proper seal. Evidence seals, including heat seals and pressure sensitive seals, must bear the initials of the person sealing the evidence. The initials must extend across the evidence seal onto the packaging.

It is not always practical or necessary to seal evidence in a container in order to protect it from loss, cross contamination or deleterious change. Large items such as furniture, doors and windows, and automotive components cannot be containerized and sealed in a practical manner. In these situations, the area of the item that has forensic importance should be covered so that the area is protected. The covering should be clearly marked indicating that this is the area of interest.

As a general rule, evidence that is received in a sealed condition should remain sealed except during analysis. If a sealed item needs to be opened prior to analysis (safety check, retrieve an RFLE, etc.), it should be resealed before storage in the evidence vault.

11.5 EVIDENCE DESTRUCTION

Evidence is the property of the submitting agency and laboratory staff will not destroy items of evidence. Laboratory staff may destroy non-evidence proficiency test samples that are no longer needed as discussed in the section on Proficiency Test Samples. It may be necessary for evidence to be entirely consumed in an analysis (see Limited Samples below).

11.6 EVIDENCE RESPONSIBILITIES

Property & Evidence Custodians (PEC) have primary responsibility for the receipt, storage, and release of evidence. They are also primarily responsible for transfers of evidence between laboratories and in and out of the evidence vault. These responsibilities include ensuring that the chain of custody is maintained on all evidence in their control and that all evidence is properly sealed.
Analytical staff is responsible for the security of evidence in their possession. This includes ensuring that the chain of custody is maintained and that evidence is properly sealed and documented for return to the submitting agency.

Evidence audits will be conducted in accordance with the WSP Regulation Manual and the Audit section of this manual.

Laboratory personnel will not routinely transport evidence to court. Transporting evidence between labs can be authorized by the lab supervisor or manager.

11.7 **Evidence Receipt**

The RFLE is the source document for user agencies to submit evidence and request laboratory examinations. The RFLE must accompany all evidence submissions. This form contains the information initially entered into the Division’s LIMS such as the name of the submitting agency, agency case number, suspect and victim names, requesting official, detailed list of evidence submitted, examinations requested, and chain of custody. When the casework is completed, the RFLE becomes a permanent part of the case file.

Evidence submission should be done by hand delivery or by means of a secure transport system. Evidence is routinely received and/or released by the laboratory via secure transport carriers. A secure transport carrier is a company such as United Parcel Service, Federal Express or the United States Postal Service.

Received evidence must conform to the preservation and integrity principles outlined in this manual. Recommended packaging protocols can be found in the Forensic Services Guide.

The CLD does not accept syringes, hypodermic needles, razor or scalp blade, or any types of skin-puncturing or sharps evidence without prior documented management approval involving the appropriate forensic scientist supervisor. These objects may pose significant risk of blood borne pathogen exposure, cuts, or needle sticks to laboratory personnel. This does not apply to knives or other sharps reportedly used in violent crimes. When received, items previously approved for submission will be inspected. If the item or packaging is deemed unsafe, the item will be returned to the submitting agency without analysis. The crime laboratory will not accept any case that includes a needle alone or a syringe with the needle detached.

When cases are received with specific suspected contaminants suggested by the agency, such as certain poisons, supervisors or laboratory managers will have the authority and flexibility to accept such cases if Crime Lab personnel have authorization for safely handling the evidence and for conducting such analyses with accepted methods. Agencies should have prior approval from the local Crime
Laboratory prior to submitting such items. It may be necessary to refer the agency to another laboratory more fully capable of handling these analyses.

Submitted firearms will be checked to see if they are unloaded and are safe to handle. The safety check will be performed by a firearms examiner or a person trained to perform this procedure. If a firearms examiner or other properly trained person is not available, the firearm must be placed in a designated area of the evidence vault until it can be checked. Once the firearm is checked and determined to be safe to handle, a notation to that effect will be made on the RFLE and on the packaging.

Agencies are encouraged to submit test fires only for IBIS entry as exemplars, not evidence. Exemplar test fires may be physically received into the laboratory or brought in as “walk ins”. Test fires submitted without an RFLE will not be treated as evidence and will not be documented in LIMS. Test fires submitted by agencies with an RFLE will be treated as evidence. Refer to the Firearms and Toolmarks Technical Procedures Manual.

If shipped evidence is received with evidence packaging lacking proper initials across the evidence seal, laboratory staff will initial the seal, add their own seal with their initials, or apply additional tape along with their initials in order to create a proper evidence seal. This will be documented in LIMS.

If shipped evidence is received unsealed and the evidence is such that it requires sealing in order to protect it from loss or deleterious change, the following options are available:

- The evidence may be returned to the submitting agency at the discretion of the Laboratory Manager.
- Laboratory staff may inventory and seal the evidence. The submitting agency will be notified and the condition of the packaging will be documented in LIMS. Documentation of notification will be in the LIMS case synopsis/notes field under the “Case Info” tab including the initials of the person recording the information and the date.

Packages and envelopes with evidence received by secure transport, and all convenience packages containing multiple evidence items, will be opened in order to verify the contents are consistent with the RFLE and so that the evidence can be properly documented, labeled, correctly stored, and entered into LIMS.

If the evidence received does not conform to the description provided on the RFLE, this will be documented on the RFLE and in LIMS. The submitting agency will be notified and the discrepancy resolved prior to analytical work commencing on an evidence item when the discrepancy involves evidence listed on the RFLE not being present or any difference that calls into question the chain of custody. Supervisors will determine if contact with the agency is appropriate.
When there is doubt as to the suitability of an item for testing, or the test required is not specified in sufficient detail, the laboratory shall consult the submitting agency for further instructions before proceeding and shall document the discussion. (See the Chapter on Review of Requests)

When receiving evidence, laboratory staff will:

- Leave original seals intact when possible.
- Locate the RFLE.
- If hand delivered, ensure that the “Submitted by:” block at the bottom of the RFLE contains a signature.
- Some agencies routinely submit evidence using their own chain of custody form. The agency form may be substituted for the chain of custody blocks on the RFLE provided the form is properly filled out and contains appropriate signatures.
- If secure transport (UPS, Certified Mail etc.) is used, document the courier and the tracking number in the appropriate block in LIMS. If there is no signature in the “Submitted by:” block, LIMS shall have, as an option, an “Agency” in lieu of a named agency representative as the submitter.
- When evidence is received via secure transport carrier, the person that opens and inventories the package must sign the RFLE in the “Received by:” block at the bottom of the form and will be included in the LIMS chain of custody using a secure transaction as the person receiving the package from the carrier. The date and time entered in LIMS will be when they take possession of the package.
- A staff member that receives a package delivery, but does not open or inventory the package, need not sign the RFLE, but will be included in the LIMS chain of custody using a non-secure transaction. Documentation of the package receipt, and storage if applicable, will be made in the notes field of the evidence tab in LIMS and if appropriate, written documentation may be included in the case file. The secure chain of custody will begin when the person that opens and inventories the package takes possession of it.
- A laboratory number will be assigned to the case and will be entered into LIMS. All items received will be documented in LIMS. All evidence items will be marked with the laboratory number by affixing a bar code label. (See LIMS Manual).
- Secure the evidence in the laboratory evidence vault or alternate evidence storage facility.
- Document any discrepancies on the RFLE and in LIMS.
- Enter the evidence receipt into LIMS by:
  - Entering the agency name, agency case number, offense, Uniform Crime Reporting (UCR) code, and the offense date (if available)
  - Entering the suspect(s) and/or victim(s) name(s)
o Entering each evidence submission and its chain of custody
o Creating a lab request and linking it to the specific evidence item by “relating” the request
• A separate “request” bar code will be printed and affixed to the RFLE.

11.8 **STANDARD EVIDENCE ABBREVIATIONS**

Below is the approved list of abbreviations for describing the evidence received by the laboratory:

- **SE**  Sealed envelope
- **SPB**  Sealed plastic bag
- **SPPB**  Sealed paper bag
- **SPKG**  Sealed package
- **SBOX**  Sealed box
- **SSTYRO**  Sealed Styrofoam box
- **SCAN**  Sealed can
- **SSAK**  Sealed sexual assault kit

It is not mandatory to use abbreviations; if not using abbreviations, write out fully the description of the evidence received. If using abbreviations, use the standard list.

If the item received is not described in the abbreviation list, write out the description fully.

If the evidence packaging is glue-sealed, gum-sealed, self-sealed or heat-sealed, the type of seal may be described in LIMS but it is not required. It is only required to indicate whether or not the item is in an appropriate sealed condition (i.e. sealed) as defined in the Forensic Services Guide. The seal description may also be combined with the approved abbreviations. For example, one SE (glue-sealed) or one SPB (heat-sealed).

To better describe a received item, additional information may either be combined with the approved abbreviations or written out fully. For example, one SBOX (rape kit) or one sealed hard plastic container.
11.9 **Evidence Items Produced During Casework**

Any substance permanently removed, extracted, or otherwise physically separated from a submitted item of evidence for any reason can itself be considered evidence. Individual technical manuals will address specific issues in this regard.

Any such substance shall be collected in an appropriate container and handled in a manner that protects it from contamination, loss or deleterious change. If this substance represents a new item, then it must be accounted for by proper marking, sealing, and documenting in the examination documentation, laboratory report, RFLE and LIMS. All new items will be returned to the submitting agency with all of the original items upon completion of examination.

11.9.1 **Crime Scene Evidence**

Evidence collected by CLD crime scene responders will be fully documented and identified, and packaged as to protect it from loss, cross transfer, contamination and/or deleterious change. All evidence will be turned over to the agency with jurisdiction over the case. An exception can be photographs taken at the scene, which will be submitted to the agency as soon as possible.

11.9.2 **Evidence Transfers**

The location of all items of evidence while in the custody of the CLD will be documented in LIMS. **EXCEPTION:** Crime Scene Response – Crime Scene evidence collected at the scene is released directly to the investigating agency before subsequent submission to the Crime Lab for analysis.

Documentation of evidence in and out of the evidence vault and of the transfer of evidence items from one scientist to another must be recorded in LIMS. Secure transactions are mandatory whenever a laboratory staff member is involved in the evidence transaction, except as noted above for staff receiving but not inventorying evidence. Any exceptions to this policy must be approved by local laboratory management and documented in the notes field of the LIMS evidence tab. A transaction is considered “Secure” when a lab staff bar code is scanned and the lab staff member enters his/her PIN. “Via” is not a required field for secured transactions. Evidence transfers may also be recorded in the examination documentation.

Laboratories receiving transfers from other laboratories will follow the standard evidence intake procedures.

11.10 **Total Transfer**

A total transfer between laboratories occurs when one laboratory has transferred all evidence items on an RFLE to another for examination. The original RFLE
documents the chain of custody from the submitting agency and accompanies the evidence being transferred.

The transferring laboratory will:
- Assign a case number to the request;
- Retain a photocopy of the RFLE;
- Ship the original RFLE and the evidence to the receiving laboratory;
- Document the chain of custody transaction as detailed in the LIMS Operations Manual (see LIMS Manual 20.05 and 21.0);
- Provide instructions on the RFLE if the receiving lab should not return the evidence to the submitting agency.

The receiving laboratory will:
- Document the chain of custody as the receiving laboratory as detailed in the LIMS Operations Manual (LIMS Manual 20.04 and 21.0)
- Return the evidence to the submitting agency unless otherwise instructed by the transferring laboratory.

11.11 Partial Transfer

A partial transfer occurs when only a portion of the evidence on an RFLE is transferred between laboratories.

The transferring laboratory will:
- Generate a new RFLE that retains the original laboratory and agency case numbers, includes all other relevant information, and lists those items being transferred; retain a copy of the new RFLE;
- Ship the new RFLE and the evidence to the receiving laboratory, and include a photocopy of the original agency RFLE (which will document the chain of custody from the original submitting agency);
- Document the chain of custody transaction as detailed in the LIMS Manual (see LIMS Manual 20.05 and 21.0);
- Provide instructions on the RFLE if the receiving lab should not return the evidence to the submitting agency.

The receiving laboratory will:
- Document the chain of custody as the receiving laboratory as detailed in the LIMS Manual (LIMS Manual 20.04 and 21.0).
- Retain the original laboratory case number will be retained.
- Return the evidence to the submitting agency unless otherwise instructed by the transferring laboratory.

For transfers of evidence where the chain of custody is documented on an agency form, rather than an RFLE:
• The transferring lab will provide the receiving lab with:
  o Either the original agency chain of custody form for total transfers, or for partial transfers, a copy of the agency form;
  o The original RFLE for total transfers, or for partial transfers, a filled out new RFLE.
• The evidence will be returned to the originating lab ONLY in cases where the agency form was used in lieu of the RFLE chain of custody blocks. The originating lab will return the evidence to the agency using the agency form.
• Only the lab(s) receiving the evidence from the agency and releasing the evidence back to the agency signs the agency form. The RFLE will be used in all other situations.
• The agency will be provided a copy of the RFLE to show chain of custody from one lab to another.
• A copy of the agency form will be retained in the case file in instances where it was used in lieu of the RFLE.

11.12 LIMITED SAMPLE

If during the course of analysis a sample which cannot be reproduced is determined to be limited, the scientist must notify either the case detective/investigator or the prosecuting attorney, whichever is most appropriate, of the need to consume the entire sample. A representative of the law enforcement agency or the prosecuting attorney must provide written approval before the analysis may proceed. The written approval shall become a permanent part of the case file, and may be in the form of a note, memo, letter or e-mail.

11.13 EXEMPLAR MATERIALS

Exemplars may or may not be considered evidence. Please see the definition of evidence at the beginning of this chapter. Exemplars may be sent to the submitting agency along with the case evidence upon completion of examination.

These materials may include but are not limited to:
• Test prints from footwear or tires
• Test marks from tools
• Fingerprint exemplars
• Handwriting exemplars
• IBIS test fires

If considered evidence, these items will be given an evidence item number and entered into LIMS. These items will be sent to the submitting agency along with the case evidence upon completion of examination.
If the exemplars are not considered as evidence, these materials may be retained in the case file or in the laboratory (see the relevant Technical Manual). Copies of exemplar materials that are used as a mechanism to document the examination performed should be kept by the laboratory.

11.14 **Proficiency Test Samples**

Proficiency test samples will be handled in the same manner as case evidence until the Quality Process Manager determines that all proficiency test requirements have been satisfied or the samples are no longer needed for that purpose. Proficiency test samples received from an external proficiency test provider will be deemed sealed. See the section on Proficiency Testing.

11.15 **Evidence Storage**

Evidence will be stored in the laboratory evidence vault or in an alternate evidence storage location within the laboratory. Alternate storage locations include evidence refrigerators/ freezers, individual evidence lockers, or unit evidence storage areas. These facilities must be locked and secured during off-duty hours. When evidence items need to be stored or conditioned under specified environmental conditions, these conditions shall be monitored, recorded, and described in the respective functional area procedures.

Evidence items are to be resealed as soon as practicable after the requested testing is completed. Evidence in the process of examination may be left unattended for short periods of time but must be in a secure laboratory area and protected from contamination or loss. Examples of short periods of time may include, but are not limited to, rest breaks, meal periods, phone calls and short conferences. If evidence examinations cannot be completed in a day, the evidence may be returned to the vault in an un-sealed condition, on a cart, for example, or one of the alternate storage locations listed above, provided the evidence is protected from loss or contamination and is clearly identifiable as evidence. Alternately, evidence may be left on the bench or examination table un-sealed, again as long as it is protected and identified as evidence. Seized drug evidence may not be left in the open but must be placed in locked storage. Under certain circumstances it may be necessary to leave evidence unsealed, pending further work, such as latent fingerprint cards held for future comparisons. In these situations, it will be permissible to leave the evidence unsealed as long as it is being stored in a protected and secure location, with a limit of one year. Beyond a year the evidence must be sealed.

Evidence that falls under the following categories may be kept in open laboratory examination areas if marked as evidence and protected from deleterious change:

- Evidence too large for the vault or alternate storage area;
• Evidence that requires special handling because of chemical or biological hazards, possible cross-contamination with other evidence, or to maintain evidentiary value (such as drying an item to be examined for DNA).

Specific types of evidence will be stored in accordance with requirements in the Forensic Services Guide.

Convicted offender DNA samples submitted to the CODIS Laboratory are not considered evidence and are retained in the laboratory indefinitely.

11.16 Evidence Return

All submitted evidence including items generated during casework shall be returned to the submitting agency upon the completion of examination.

Evidence will be released only to the submitting agency or approved subcontracting laboratories. An exception is possible when there is a valid court order requiring the release of the evidence to a party other than the submitting agency, or the submitting agency directs the laboratory in writing. A copy of the court order or written request will be maintained in the case file.

Returned evidence must be properly documented on the RFLE (or on the agency chain of custody form if used) and in LIMS. If not all items on the RFLE are being returned in a given transaction, the “Item(s) being released/returned” section of the RFLE will be completed to list the items that are being returned. Use of this section is optional if all items are being returned or released.

Evidence items returned or transferred by secure transport carrier require confirmation receipts which will be retained in the case file. If delivery confirmation is not confirmed within ten days of shipping, the lab manager will be notified. The lab manager will ensure that affected parties are informed as appropriate.

11.17 Cancelled Requests

Employees may cancel a request and return the evidence to the submitting agency at any time when the circumstances of the case dictate it. These circumstances may include, but are not limited to, lack of probative value of the evidence; insufficient information accompanying the evidence to perform the request; lack of response by the agency, prosecutor, or other submitter.

Note: If any analysis has been performed, a lab report is required regardless of the agency or prosecutor canceling the request.
Cancellation of requests, other than for administrative corrections of data entry, will be communicated to the customer. The communication will be noted on the RFLE and documented in the “Case Info” tab synopsis/notes field in LIMS.

11.18 HAZARDOUS MATERIAL

Unidentified material is not considered hazardous until it has been determined to be so by analysis. In order to comply with Federal Department of Transportation regulations regarding the shipping of hazardous materials, it is recommended that the laboratory follows the requirements of the small quantity exception rule (See Code of Federal Regulations 49 CFR 173.4) if possible. If not possible, agencies which have submitted evidence to the CLD that has been determined to be hazardous shall be asked to retrieve the evidence from the laboratory in person.

11.19 EVIDENCE AUDITS

Evidence audits shall occur as described in the Property Inventory/Audit section of the WSP Regulation Manual (21.00.020) and WSP Property and Evidence Custodian Manual.

All evidence audit reports and the inventory sheets from which the audit is conducted will be retained and centrally located in each laboratory through at least one cycle of accreditation or per the retention schedule, whichever is longer. If the audit requires the inventory of evidence in the possession of a scientist or other personnel who are not present, this will be conducted with a witness who will initial and date the portion of the inventory sheet that they witnessed. All inventory sheets will be signed and dated by whoever participated in the audit. These audit reports will be subject to inspection during the annual Standards and Accountability Sections audit.
12 NONCONFORMING WORK AND CORRECTIVE ACTIONS

12.1 Policy

In the event that any laboratory member becomes aware of a nonconformity with any aspect of testing, work, or the results of this work (e.g., in analysis, proficiency tests, reports, testimony, or care and preservation of evidence), correction shall be taken immediately including any decision about the acceptability of the nonconforming work. When a nonconformity has been identified, an evaluation of the significance of the nonconforming work will be made. Where the evaluation indicates that corrective action is needed, the corrective action procedures shall be promptly followed.

All employees will be trained on the corrective action process at an appropriate time for their duties and responsibilities. In order to enhance the quality and effectiveness of root cause analysis and corrective actions, training will include root cause analysis principles and processes and its acceptance within the laboratory environment as part of a just culture.

12.2 Procedure

The Corrective Action procedure is a step-wise process as outlined below. The process is entered and tracked by entering into the Quality Process Improvement Tracker (QPIT).

12.2.1 Identification

While not an exhaustive list, identification of nonconformities may occur through any of the following:
- internal or external inquiries or complaints
- quality control
- instrument calibration
- staff observations
- supervisor observations
- technical and administrative review of reports and case files
- indications of inadequate technical review
- calibration certification checking
- management reviews
- internal or external audits

12.2.2 Notification

When a potential nonconformity has been identified, the Standards and Accountability Section will be notified. Staff working in the DNA functional area or CODIS will use the appropriate Quality Variance (QV) QPIT entry, which will be submitted to the DNA Technical Leader with a copy to the Lab Manager. The CODIS
Administrator will be notified of nonconformities that impact the DNA records entered into CODIS. All others, including those identified in a Court Testimony Performance Evaluation, will use the Notification of Nonconformance QPIT entry, which will be submitted to the Quality Assurance Manager with a copy to the Lab Manager. A nonconformity as described in an internal or external audit report is entered into the QPIT as a Corrective Action Request. A letter of inquiry from the ANAB Proficiency Review Committee is entered into the QPIT as a Proficiency Test Inquiry.

The notification may be documented and submitted using the QPIT by any staff member and is routed through their supervisor. A supervisor, Lab Manager, DNA Technical Leader, or SAS Manager can decide to propose a corrective action plan with the notification.

When a DNA analyst identifies DNA contamination by an employee outside of the DNA section the following notification process shall be followed:

- The DNA analyst shall inform their supervisor and Lab Manager of the observed contamination. The DNA analyst will complete QV QPIT entry. If the contamination is outside of the CLD, the analyst and supervisor will discuss the best approach for notification.
- The DNA analyst, and/or their supervisor, will consult with appropriate individuals (including, but not limited to, the contaminating employee and their supervisor) to determine the root cause of the contamination and possible preventive measures. The QV QPIT will be updated and completed by the DNA analyst, contaminating employee, or an appropriate supervisor (as determined by the Lab Manager) in a timely manner.
- The contaminating employee, their supervisor, the DNA analyst and the supervisor of the DNA analyst will acknowledge the QV using the QPIT, which will be updated as applicable.
- The DNA Technical Leader and the Laboratory Manager are notified through the QPIT as applicable.
- The DNA TL will review the QV entry, and if complete and accurate, will approve the QV in QPIT. The QV can be rejected if additional information is still needed for an adequate review. The DNA TL can mark the QV as nonsubstantive with or without further investigation, or substantive with an investigation needed.
- Substantive nonconformance requires disclosure to ANAB within thirty days of determining that nonconformance has occurred. If the nonconformance has been determined to be substantive, the Quality Assurance Manager will provide written notification to all involved parties and assign an individual to prepare the Root Cause Analysis and Corrective Action Plan. The thirty day clock will begin from the date of that notification. In consultation with all involved parties, the disclosure report will be prepared and submitted by the Quality Assurance Manager.
• The Quality Assurance Manager or, if DNA related, the DNA Technical Leader, may state in the QPIT, that follow-up is unnecessary if it is thought that the follow-up does not serve a useful purpose.

12.2.3 Root Cause Analysis

The corrective action shall start with an investigation to determine the root cause(s) of the problem. This is a process of fact finding used to evaluate all aspects of the incident, including the policy or procedure involved, to identify the basis of the nonconformity. This process is a tool designed to help identify what, how, and why an event occurred, or the underlying factors leading up to a casework error or nonconformity. There may be more than one cause for a nonconformity. Whenever a discrepancy or nonconformity occurs in casework or CODIS lab work, the cause(s) should be determined if possible. The Quality Assurance Manager may assign members of the CLD other than the supervisor to conduct the root cause analysis investigation. Those performing root cause analyses should have already completed root cause analysis training.

The investigation may include an evaluation of procedures, staff skills and training, consumable supplies, equipment and instruments, calibration status, customer requests and requirements, samples, reagents, controls, and other items as deemed necessary during the investigation. The investigator shall consult with all necessary personnel, including with the staff member involved, to determine the basis of the nonconformity as completely as possible.

Nonconforming work may be a systemic error rather than an employee error, or a combination of both. The root cause analysis may provide a platform for process improvement, and may help guide value-additive changes in policy and procedure.

Refer to Appendix 1 for root cause analysis guidelines and procedures.

12.2.4 Evaluation of the Significance of Nonconforming Work

The level of significance of the nonconformity will determine the appropriate corrective action. Nonconformities occur in a continuum of significance and severity. Because of this, they must be evaluated for their significance and a decision made regarding the appropriate corrective action for the nonconforming work. This evaluation is the responsibility of the Standards and Accountability Section: the DNA Technical Leader for DNA related incidents of nonconformance, and the Quality Assurance Manager for all others.

The evaluation of the significance level of the nonconforming work must consider risk levels (see table below):

• The severity of the nonconformance, including the impact and acceptability of the nonconforming work and on previous test results;
• The possibilities and implications of the nonconforming work recurring
• If there is/was laboratory conformance with its own policies and procedures
• The suitability of those laboratory policies and procedures

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For instances where the nonconforming work is determined to be nonsubstantive, the correction or immediate action taken, as reported in the QPIT entry, may be sufficient, and no (or limited) further action would be necessary. However, the supervisor, Lab Manager, Quality Assurance Manager or DNA Technical Leader (for DNA related incidents of nonsubstantive nonconforming work) may decide to implement a corrective action plan.

In instances where the Standards and Accountability Section determines the nonconforming work is substantive, the corrective action procedure, described below, will be implemented. If the nonconformity is determined to be related to a functional area wide, laboratory wide, or system wide deficiency, the SAS Manager and CLD Commander will ensure casework is discontinued as appropriate until the nonconformity is addressed and resolved. Where necessary, the customer(s) will be notified and work recalled.

If a Corrective Action is required, the Quality Assurance Manager will assign a person to prepare the Corrective Action Plan. The DNA Technical Leader can designate the person to prepare the Corrective Action Plan if the nonconformance is in DNA.

12.2.5 Corrective Action Plan and Implementation

The individual preparing the Corrective Action Plan will, in consultation with the functional area’s Technical Leads, supervisor, Management Liaison and/or DNA Technical Leader as necessary, identify potential corrective and preventive actions. They must select and implement the action(s) most likely to eliminate the problem and to prevent recurrence. Corrective actions shall be to a degree appropriate to the magnitude and the risk of the problem and be completed within a reasonable timeframe agreed to or determined by SAS. For the DNA functional area and CODIS, the DNA Technical Leader must approve all corrective action plans before they are implemented. The Quality Assurance Manager must approve all others. Once the corrective action plan has been approved, as designated in QPIT, the CAP will be implemented.
Difficulties with an employee’s individual work performance will normally be addressed by the employee’s supervisor with assistance and input from other appropriate individuals if necessary. The actions taken to correct the problem should be focused on the professional development of the employee, which normally includes remedial training and other assistance designed to help the employee overcome the problem.

12.2.6 Timeline

A timeline for completion will be included in the CAP. If the planned actions call for interim reviews prior to completion, these shall be scheduled and included in the timeline. The approved CAP should indicate who will prepare the final corrective action report (most often the author of the plan). The Quality Assurance Manager, Laboratory Managers and Supervisors will ensure that corrective actions are implemented and completed within a reasonable timeline agreed to and approved by SAS.

12.2.7 Corrective Action Plan Completion

Corrective Action Reports (CARs) are prepared after completion of the CAP and document the results of the CAP. The report will document any required changes from the corrective action and should include any further needed actions or recommendations. In lieu of an IOC, the information may be entered into the QPIT. All CARs require approval by the Quality Assurance Manager, and if concerning DNA, require the DNA Technical Leader’s approval. Courtesy notifications will be given to the CLD Commander and all involved parties. Internal and external DNA QAS audits are reported to NDIS by the CODIS Manager on the Lab Audit Certification form. Electronic copies of all external DNA QAS audits and associated remediation are sent to NDIS.

12.2.8 Follow-Up

In order to ensure continued conformance and that the corrective actions have been effective, monitoring of the employee’s performance, laboratory performance, or system performance will be conducted by the employee’s supervisor, or the individual responsible for carrying out the corrective action plan.

Internal auditors will review corrective actions taken during the Annual Internal Quality Audit, and will look for reoccurrences of problems or documentation demonstrating that there has been no reoccurrence.

Each functional area, at least once per year, will review reported nonconformances, including internal and external assessment nonconformances, for their respective functional area, from all labs. The review may be conducted among all staff or at the management liaison/supervisor/technical lead(er) level, and shall be documented. This supports continuous improvement and consistency among labs.
by making scientists aware of nonconformances from other labs and what was learned in the process.

12.2.9 Reporting Requirements and Responsibilities
ANAB requires a summary of nonconforming work events, the actions taken, and summary of any other substantive corrective actions completed, or in process, since the last on-site visit included in each laboratory’s annual provision of conformance documents. Each lab manager, with the assistance of the Standards and Accountability Section as needed, will prepare these summaries as part of the laboratory’s required reporting to ANAB.

Records of Corrective Actions will be retained for at least one accreditation cycle and in accordance with the agency retention schedule.

12.2.10 Notification of Customers
When substantive nonconformities occur in casework, it may be necessary to notify the customer of the facts surrounding the event. If necessary, an amended laboratory report will be prepared as soon as possible and provided to the submitting agency. The case file will contain documentation of the technical measures taken to resolve the nonconformity.

12.3 Responsibility for Authorizing Resumption of Work
In cases where an analyst has been removed from casework, or when required by the corrective action plan, a follow-up competency test will be issued by the Quality Process Manager following successful completion of the corrective action plan. An analyst may only return to casework on the authorization of the CLD Commander. If a process or procedure has been removed from use, it also will not be used until authorized by the CLD Commander. Authorization to return to casework from the DNA Technical Leader is additionally required if the actions involve DNA or CODIS.

12.4 Preventive Actions
Preventive actions are designed to eliminate the cause of a potential nonconformity and prevent occurrence. Identification of needed improvements, either technical or concerning the management system, evaluation and implementation of a preventive action may include one or more of the following:

- Research regarding policies and procedures in other crime laboratories or jurisdictions
- Consultation with customers to ascertain the extent of their needs
- Consultation with CLD employees to obtain developmental suggestions
- Validation of technical methods following the Method Validation section of the CLD Quality Operations Manual
• Monitoring of effectiveness with CLD employees and customers of laboratory services

Personnel are encouraged to identify preventive actions as opportunities to improve quality and correct potential sources of nonconformity before they become problems.

Preventive action proposals shall be brought to the attention of the Supervisor, the appropriate technical lead(s) (in addition, if DNA or CODIS, the DNA Technical Leader) for the discipline, Laboratory Manager, appropriate Management Liaison, and/or Quality Assurance Manager through written correspondence such as e-mail or IOC. The Supervisor, Laboratory Manager, and/or appropriate staff shall evaluate the suggestion and work with the submitting individual to develop an action plan. As the preventive action is implemented, it shall be monitored for effectiveness as outlined in the action plan.
13 TRACEABILITY AND QUALITY CONTROL

Many factors contribute to the accuracy and reliability of the tests performed by the CLD, including:

- The training and qualifications of personnel
- Technical/analytical methods
- Reference standards, reference materials, reagents and supplies
- The selection, calibration and maintenance of equipment

The CLD will take account of these and other factors and will ensure that the personnel are properly qualified and trained; that procedures are validated; that reference standards and materials are traceable and verified for performance; and that equipment is calibrated and/or verified. All procedures, reagents, reference standards and materials, supplies and equipment/instrumentation that affect the quality of the tests and/or calibrations will be controlled.

13.1 TRACEABILITY AND QUALITY CONTROL OF REAGENTS

13.1.1 Policy

All commercially and laboratory prepared reagents, as well as chemicals used to prepare reagents, used for casework analysis within the CLD will be of sufficient quality to assure the integrity of the analytical results. All reagents must be checked to ensure their reliability and that the quality will equal or exceed that necessary for the type of testing or use designated in the functional area technical manual. How this will be performed and the frequency of reagent checks will be determined by each functional area and will be found in the respective technical manuals. The reliability testing shall occur before use or, if appropriate, concurrent with testing provided reagents were verified as meeting specified requirements.

Reagents prepared in the laboratory shall be labeled with, at a minimum, the identity of the reagent, the date of preparation or lot number, and, as applicable, (for example, other than room temperature), storage requirements. Records shall be maintained identifying who made the reagent, the components used in preparation, and that the reagent was tested and worked as expected.

13.2 EQUIPMENT

13.2.1 Policy

Equipment and its software used for testing, calibration and sampling shall be capable of achieving the accuracy required and shall comply with specifications relevant to the tests and/or calibrations concerned. Before being placed into service for use, equipment (including that used for sampling) shall be successfully calibrated or performance verified.
to establish that it meets the laboratory’s specification requirements and complies with the relevant standard specifications.

13.2.2 Procedure

Equipment will have regular maintenance, calibration (if required) and performance verifications to ensure continued performance.

Laboratory Managers have final approval authority before new equipment is placed into service for casework and must ensure that the validation and/or performance verifications take place.

13.2.3 Personnel Equipment Use

Equipment shall be operated by authorized personnel as determined by the individual lab managers. The lab manager has the responsibility to ensure that authorized equipment user list(s) for their laboratory are updated and available.

Recognizing that there may be occasions where an individual scientist may need access to and use of equipment in another Division laboratory, if the scientist has been given authorization to use similar equipment and software in one Division lab, that authorization may automatically apply to any laboratory using comparable equipment and where appropriate, software.

Laboratory technicians, scientists in a training status and interns will be given authorization commensurate with their level of experience. Each lab manager will maintain documentation of persons authorized to operate the laboratory’s equipment used for testing, calibration and sampling. Laboratory technicians and interns must have their training documented and be specifically authorized to operate specified laboratory equipment.

13.2.4 Equipment Identification

Each item of equipment and its software used for testing and calibration and significant to the result shall be uniquely identified.

In laboratories with multiple items of equipment of the same make/model, each item of equipment will be uniquely identified and the identifier will be used in the examination documentation and hard copy equipment data. In a laboratory with only one item of equipment for a specific test or procedure, the equipment identification is documented in the laboratory equipment list.

13.2.5 Equipment Documentation

Up-to-date instructions on the use and maintenance of equipment (including any relevant manuals provided by the manufacturer of the equipment) shall be readily available for use by the appropriate laboratory personnel.
Maintenance procedures will include a maintenance plan that indicates the frequency and type of maintenance to be performed (i.e., annual, as needed or by manufacturer). Scheduled manufacturer maintenance contract information (if applicable) will also be retained by the laboratory/discipline.

Each item of equipment and any associated software significant to the tests and/or calibrations performed will have records that are maintained. The records shall include at least the following:

- Equipment identity: type, manufacturer, model, serial number or unique name, and current location
- Identity of the item of equipment’s software and firmware, if applicable
- Original equipment paperwork provided with equipment installation
- Manufacturer’s instructions, if available, or reference to their location
- Maintenance plans, where appropriate
- Maintenance procedures and records of maintenance performed
- Date of maintenance, initials of the person doing the maintenance, activity conducted, including damage, malfunction, modification to, or repair of, the equipment
- Performance verification procedures, or reference to appropriate technical procedures manual
- Documentation of performance verification
- Calibration procedures (as required), including procedures for when intermediate checks are needed to maintain confidence in the calibration status of the equipment
- Scheduled calibration (if required) including dates, results, reports and certificates, adjustments, acceptance criteria and the due date of next calibration or the calibration interval
- Any damage, malfunction, modification or repair to the equipment
- Internal validation procedure, data and documentation
- List of authorized equipment users

Equipment maintenance, calibrations, and results of performance verifications that are performed will be documented and maintained in an equipment maintenance log. This log will be kept in close proximity to the equipment whenever possible. An electronic log is an acceptable alternative or complement to a written log. Because equipment logs are part of the case record, the equipment log retention time will be the same as for case files. Maintenance/verification logs will be kept with the equipment if the equipment is transferred to another laboratory.

13.2.6 Methods Used on Equipment

The laboratory or discipline will ensure that all methods used on analytical equipment, either newly purchased, or existing equipment that is significantly modified such that the change(s) affects the outcome of the test, or are to be used for new analytical applications,
are properly validated prior to being placed in service for casework in a CLD laboratory. Refer to the section on Method Validation for more details.

A laboratory may adopt a validated method which, e.g. has been published as a standard, or purchase from a qualified vendor a complete measuring system to be used for a specific application. In both these cases, basic validation work has already been carried out. However, the laboratory must confirm its ability to apply the method. This verification requires that some experimental work be completed in order to demonstrate that the method works in the CLD laboratory.

13.2.7 Equipment Data

When computers or automated equipment are used for the acquisition, processing, recording, reporting, storage or retrieval of test or calibration data, the laboratory shall ensure that:

- Computer software developed by the user is documented in sufficient detail and is suitably validated as being adequate for use;
- Procedures are established and implemented for protecting the data; such procedures shall include, but not be limited to, integrity and confidentiality of data entry or collection, data storage, data transmission and data processing;
- Computers and automated equipment are maintained to ensure proper functioning and are provided with the environmental and operating conditions necessary to maintain the integrity of test and calibration data.

Laboratory-developed software, commercial software being used outside its intended application, and custom software created by a third party specifically for the laboratory, shall be validated and records of the validation maintained. A validation plan will be developed: the method validation section can be used as a guideline.

Commercial off-the-shelf software (e.g. word processing, database and statistical programs) in general use within their designed application range may be considered to be sufficiently validated. However, laboratory software configuration/modifications shall be authorized, validated and documented as being adequate for use prior to implementation.

13.3 Equipment Performance Verification

Equipment to be used for existing applications and methods must be performance verified before use. The laboratory or discipline will ensure that all analytical equipment has their performance verified prior to use. The process will be as extensive as is necessary to meet the needs of the given application or field of application. All performance verifications will be performed by qualified personnel with adequate resources.

Performance verification procedures will include (see also section on Performance Validation for Methods):
• the principle of the verification;
• verification frequency;
• verification tolerances, acceptance criteria
• specific step-by-step verification instructions including the use of any reference standards or reference materials. When possible, all verification will be completed with traceable reference standards or materials.

Performance verification procedures that are completed will be documented in the equipment maintenance log and will include the following:

• Verification date
• Initials of the person performing the verification
• Type of verification performed (e.g. internal diagnostic or comparison to a reference standard)
• Verification results (pass or fail?);
• Identification of reference standard or reference material used
• Any comments regarding performance checks

Upon verifying the performance of a new item of equipment, the Technical Lead or for equipment in DNA, the DNA Technical Leader, will ensure that the performance verification is completed successfully. Upon completion, documentation of the performance verification will be retained in the laboratory. The Technical Lead will prepare an IOC directed to the section supervisor and/or Laboratory Manager, as appropriate, advising them that the equipment has been performance verified and is ready for use in casework. By signing and acknowledging the IOC, the Laboratory Manager authorizes the placing of the equipment into service.

Items of equipment which have previously had their performance verified do not require further authorization by the Lab Manager or the DNA Technical Leader.

13.4 Equipment Calibration

Equipment requiring calibration will be calibrated prior to being placed into service in a CLD laboratory. Functional areas will have a list of the equipment used in testing that has a significant effect on sampling, the test result, or the total uncertainty of the test result, which require calibration. Functional areas will also describe specific requirements for the calibrations. In situations where the calibration of equipment does not have a significant effect on sampling, the test result, or the total uncertainty of the test result, the laboratory shall have objective evidence to demonstrate the insignificant contribution.

Calibration status will be checked after any unexpected shutdown or removal of the equipment from service and following service or other substantial maintenance.
Calibration checking procedures will be described, where appropriate, in the functional area technical manuals. Calibration procedures shall be established for key quantities or values of the equipment where these properties have a significant effect on the results.

Equipment requiring calibration will have a documented calibration schedule. The recalibration schedule will include the frequency of calibration required, the status of calibration and the next calibration due date. Calibration check intervals will not be less stringent than that recommended by the manufacturer. If a laboratory determines that intermediate checks of the calibration status are needed, the procedure shall define the frequency of the checks. Once established, any extension in the interval of intermediate checks shall be based on documented empirical data and an evaluation of risk. Calibration/recalibration documentation and calibration certifications will be maintained in the Equipment Maintenance log.

Whenever practicable, all equipment under the control of the laboratory and requiring calibration shall be labelled, coded or otherwise identified to indicate the status of calibration, including the date when last calibrated and the date or expiration criteria when recalibration is due.

Where calibrations give rise to a set of correction factors, the functional area procedures manuals shall have procedures to ensure that copies (e.g. in computer software) are correctly updated.

In situations where the measurements require measurement traceability and calibration of equipment does not have a significant effect on sampling, the test result, or the total uncertainty of the test result, the laboratory shall have objective evidence to demonstrate the insignificant contribution.

When external calibration services are used, traceability of measurement will be assured by the use of calibration services that can demonstrate competence, measurement capability and traceability. The calibration certificates issued by these services will contain the measurement results, including the measurement uncertainty and/or a statement of compliance with an identified metrological specification.

If available, suppliers of external calibration services for reference standards requiring calibration (see section below on Traceability of Measurement Standards) and for calibration equipment where the calibration of the equipment has a significant effect on the accuracy or validity of sampling or a test result, or the total uncertainty of the test result, shall be either:

- a National Metrology Institute that is a signatory to the International Bureau of Weights and Measures (BIPM) - CIPM Mutual Recognition Arrangement with the calibration to be performed or the certified reference material listed to be purchased in Appendix C of the BIPM key comparison database (KCDB)2; or
- a service supplier accredited to ISO/IEC 17025 by an accrediting body that is a signatory to the International Laboratory Accreditation Cooperation (ILAC): Mutual
Recognition Arrangement, with the calibration of measuring equipment and/or reference standard to be performed listed in a scope of accreditation.

In situations where a supplier of external calibration services that meets the above listed criteria is not available, the laboratory shall confirm competence, measurement capability, and measurement traceability for the supplier and the service being purchased. Objective evidence of the confirmation shall be available for review. Documentation of vendor competence, capability and traceability will be maintained as described in the section on Vendor Evaluation in this manual.

13.5 **Equipment Calibration/Certification/Maintenance Schedules**

Calibration/Recertification and maintenance for the following general laboratory equipment used in case work will be conducted and scheduled accordingly:
- Scales/Balances will be calibrated at least once per year by a qualified external agency.
- NIST traceable rulers, rods, micrometers, calipers, tape measures and other measuring devices will be recertified before certification expires.
- If calibration is required for pH meters, it will be performed prior to use.
- NIST traceable thermometers and temperature probes will be recertified before their certification expires. For non-NIST traceable thermometers, each functional area that uses them will establish protocols and documentation for temperature monitoring, with periodic checks of the thermometer against a NIST traceable probe.
- Pipettes will have the calibration checked at least once yearly by a qualified external agency.
- Microscopes will be serviced as necessary by qualified microscope technicians. It is recommended that the service interval not be longer than three years. Each laboratory will maintain microscope service and repair records for microscopes in their inventory.

Calibration/recertification and maintenance schedules and procedures for discipline specific equipment will be outlined in the respective functional area technical procedures manuals.

13.6 **Equipment Out of Service**

Equipment that has been subjected to mishandling, gives questionable results, or has been shown to be defective or outside specified limits, shall be taken out of service. It shall be isolated to prevent its use or clearly labelled or marked as being out of service until it has been repaired and shown by calibration or performance verification checks to perform correctly. In addition, the removal of the equipment from service, date of removal, why the equipment was removed from service, and the date the equipment was placed back in service will be documented in the equipment log.
The laboratory shall examine the effect of the defect or departure from specified limits on previous tests and/or calibrations. If the nature of the malfunction is such that the accuracy of previous test results are not suspect and did not impact casework, the corrective steps mentioned above will be completed and documented in the equipment log.

If the nature of the malfunction is such that the accuracy of previous test results are suspect, the situation shall be immediately brought to the attention of the appropriate Technical Lead, DNA Technical Leader if DNA or CODIS, Section Supervisor and Lab Manager. The Lab Manager will inform the Standards and Accountability Section, and corrective action shall be performed. The laboratory will follow the corrective action procedure for nonconforming work.

### 13.7 Equipment Responsibilities

- Forensic Scientists/Lab Technicians assigned equipment responsibilities are responsible for performing assigned equipment validation, performance verification and maintenance and will document all necessary information concerning verification and maintenance activities in the Equipment log.
- Forensic Scientists/Lab Technicians are responsible for ensuring that the equipment in use has been properly calibrated or verified prior to use.

Technical Leads are responsible for:
- Preparing and organizing validation studies as needed
- Ensuring that calibration/verification and maintenance procedures are in place for each item of equipment and software determined to require verification and maintenance in their discipline
- Writing and modifying the verification procedures for each item of equipment in their discipline
- Monitoring compliance with calibration/verification and maintenance procedures through periodic spot checks
- Addressing problems concerning verification according to WSP CLD Policy
- Annually verifying external calibration companies are ISO compliant

Lab Managers/Supervisors are responsible for:
- Ensuring that all users are properly trained and authorized prior to equipment use
- Periodic review of all calibration/verification and maintenance records and activities
- If a problem with an item of equipment is identified, such that the accuracy of previously reported test results is suspect, the Lab Manager/Supervisor shall immediately alert the Standards and Accountability Section and the Technical Lead for that discipline. Lab Managers and Supervisors will ensure that corrective actions take place.

Standards and Accountability Section is responsible for:
• Monitoring compliance with calibration/verification and maintenance procedures through annual audits of logs.

13.8 **EQUIPMENT USE OUTSIDE THE LABORATORY**

When, for whatever reason, equipment goes outside the direct control of the laboratory, the laboratory shall ensure that the function and calibration status of the equipment are checked and shown to be satisfactory before the equipment is returned to service.

When equipment that is outside of the laboratory’s permanent control is used by CLD analysts for reporting results, the analyst must document that the equipment has been properly maintained and controlled.

If analysis is performed by CLD personnel at a laboratory not regulated by the WSP CLD, the following restrictions apply:

• The analysis may be conducted if the outside laboratory can demonstrate that it meets the requirements of ISO. The use of equipment from another laboratory and the name of the laboratory must be documented in the examination documentation and in the lab report. The outside laboratory must provide validation documentation to demonstrate that the instrument meets the needs of analysis.

• Analysis may be contracted to another laboratory. The contract laboratory will be solely responsible for the content of its own casework report. The referral of evidence to an outside or contract laboratory must have the approval of the submitting agency and the Prosecuting Attorney’s office where applicable. Discussions with the submitting agency and Prosecuting Attorney’s office should include the limitations and expenses of outside analysis. The CLD lab reports must reflect the referral of the evidence to the other laboratory.

• Convicted offender DNA samples may be contracted out without notification to the collecting agency.

13.9 **MEASURING OR CALIBRATION EQUIPMENT**

Functional area technical procedure manuals will include procedures, or make reference to procedures, to ensure safe handling, transport, storage, use and planned maintenance of measuring equipment (i.e., balances, pipettes, calipers, trigger pulls, rulers) to ensure proper functioning and in order to prevent contamination or deterioration.

Test and calibration equipment, including both hardware and software, shall be safeguarded from adjustments which would invalidate the test and/or calibration results.

Should the measuring equipment be used outside the permanent laboratory for tests, calibrations or sampling, additional procedures should be considered and included in the procedures as needed.
13.10 Traceability of Measurement Standards

13.10.1 Policy

All test equipment used in CLD laboratories that has a significant effect on the measurement result and their associated uncertainties of measurement, will be traceable to national and/or international standards of measurement. This will be done through the use of a measurement standard. The CLD will safely handle, transport and store these measurement standards in order to prevent contamination or deterioration and in order to protect their integrity.

13.10.2 Procedure

Measurement standards or materials (e.g., thermometers, weights) used to check accuracy of other equipment or instruments shall not be used for other purposes.

All calibrations and adjustments using these standard materials will be documented.

All in-house NIST traceable CRMs must be periodically checked and recertified by an external agency to maintain their NIST traceability. Recalibration or recertification of these materials will take place before their certification expires.

Vendors used for calibration or recertification of these standards shall provide documentation of NIST traceability.

If available, suppliers of certified reference materials used to establish or maintain measurement traceability shall be either:

- a National Metrology Institute that is a signatory to the BIPM - CIPM Mutual Recognition Arrangement with the certified reference material listed in the BIPM key comparison database (KCDB), or
- an accredited reference material producer that is accredited to ISO Guide 17034 (or to ISO Guide 34:2009 until November 1, 2019) by an accrediting body that is a signatory to a mutual or multilateral recognition arrangement in an ILAC recognized regional accreditation cooperation or the ILAC Mutual Recognition Arrangement, with a scope of accreditation covering the certified reference material.

In situations where a certified reference material producer that meets the above criteria is not available, the laboratory must confirm competence, measurement capability and measurement traceability for the supplier and product being purchased. Objective evidence of the confirmation shall be available for review. Documentation of vendor competence, capability and traceability will be maintained as described in the section on Vendor Evaluation in this manual.

When using external calibration services, traceability of measurement shall be assured by the use of calibration services from laboratories that can demonstrate competence,
measurement capability and traceability. The calibration certificates issued by these laboratories shall contain the measurement results, including the measurement uncertainty and/or a statement of compliance with an identified metrological specification. Following service, maintenance and recalibration by such vendors, the certification or documentation provided by them will be maintained in the laboratory.

If mishandling of standard materials brings accuracy into question, the standard materials shall be taken out of service and recalibrated.

To the extent possible, the CLD shall utilize measurement standards that are traceable to SI units of measurement, to certified reference materials, or to other applicable verification sources.

When traceability of measurements cannot be made in or is not relevant to SI units, then measurement standards will establish traceability by one of the following:

- The use of certified reference materials and/or values of certified reference materials from a competent supplier
- The use of specified methods, published standards, and/or consensus standards that are clearly described and accepted as providing measurement results fit for their intended use and ensured by suitable comparison
- Participation in inter-laboratory comparisons

Documentation of this traceability to SI units or CRMs and the recalibration/recertification information shall be maintained by the appropriate Laboratory, Laboratory section or the Standards and Accountability Section.

Internally developed measurement standards must be checked against published references or certified reference materials, identified and controlled. Documentation of traceability must be maintained in the laboratory.

13.11 Traceability of Reference Standards and Materials

13.11.1 Policy
Reference standards and materials that are maintained and used in casework for identification, comparison or interpretation purposes shall be fully documented, uniquely identified and properly controlled. In addition, CLD employees will safely handle, transport, store and use these reference materials and standards in such a manner as to prevent contamination and deterioration and to protect their integrity.

13.11.2 Procedure
Each laboratory is responsible for ensuring that their reference standards and materials are fully documented, uniquely identified and properly controlled.
If mishandling of reference standards or materials brings their accuracy into question, the standards or materials shall be taken out of service until accuracy is verified.

Reference materials and standards having an expiration date must be periodically checked and renewed prior to their expiration dates. Expired reference materials may be used as secondary standards, but may not be used for calibration purposes. Expired standards or materials may also be used for non-casework related functions such as validation, training, performance checks, and competencies at the supervisor’s discretion. Expired standards or reference materials used for these purposes must be clearly identified as such.

For many types of analysis, calibration and instrument performance checks may be carried out using laboratory-made standards containing the analytes under test, prepared from chemicals of known purity and composition, or matrix matched standards. Alternatively, standard solutions may be purchased. Many chemicals can be purchased with the manufacturer’s statements or certificates. Wherever possible, laboratories should obtain supplies of chemical reference standards from ISO compliant suppliers.

Records will be retained for reference standards and materials to include where applicable: verifications, results of verifications or periodic checks, acceptance criteria, relevant dates and the period of validity.

Examples of WSP CLD Reference Standards include:
- FTIR NIST traceable polystyrene Reference Standards
- Mass Spectrometer perfluorotributylamine (PFTBA) Reference Standard
- Glass Reference Standards for Refractive Index Measurements

Examples of WSP CLD Reference Materials include:
- Ignitable Liquid Reference Materials
- Drug Reference Materials
- Fiber Reference Materials
- Polymer Reference Materials
- Firearms Reference Collection

13.12 DRUG REFERENCE MATERIALS


13.13 INDIVIDUAL CHARACTERISTIC DATABASES

13.13.1 Policy

All individual characteristic database samples under the control of the CLD will be treated as reference materials. Individual characteristic database samples under the control of the CLD include known biological samples from convicted offenders (CODIS). The local CODIS
DNA database (LDIS) will be administered by the local CODIS administrators. The State DNA database (SDIS) will be administered by the CLD CODIS Laboratory Manager.

13.13.2 Procedure
Whenever possible, all samples will be treated in a manner that reasonably ensures their utility as reference materials. In general this will consist of:

- Maintaining control of samples utilizing a unique identifier. Agencies contributing to individual characteristic databases may use various methods to accomplish uniquely identifying database samples;
- Protecting the samples from loss, cross transfer, contamination and/or deleterious change. Individual characteristic database samples must be treated in a manner that maintains their use as reference materials;
- Restricting access to those persons authorized by the Laboratory Manager. The laboratory manager shall authorize access to those individuals having a legitimate purpose and maintain that list in the laboratory for review by auditors. Such persons include, but are not limited to, individuals responsible for database maintenance and administration and equipment repair. These individuals may or may not be under the control of the laboratory. Documentation of access to the CODIS database will be maintained by the state CODIS administrator;
- Each unit that utilizes an individual characteristic database will be responsible for preparing and using a protocol for handling and storing such reference material as it relates to individual characteristic databases, taking into account the special needs of the unit. These protocols must be described in the functional area technical manuals.

13.14 Other Databases

13.14.1 Policy and Procedure
The CLD maintains other databases for the cataloging, storing and retrieval of quality and technical information. Access to these databases will be limited to authorized staff. Administration and changes to these databases will be by designated individuals only. Laboratory Managers will designate an individual or individuals for the management and administration of the laboratory specific databases.

The CLD databases will be managed by the Quality Process Manager or designee. The CLD LIMS database will be administered by the FLSB IT Manager.

Other databases include but are not limited to:

- Laboratory Chemical Inventory Database (All laboratories)
- Laboratory training samples collections
- Laboratory Key Inventory Database (All laboratories)
13.15 Measurement Uncertainty

13.15.1 Policy
The CLD will have and apply appropriate procedures for estimating the measurement uncertainty for all reported quantitative test results, where required. The discipline procedures will identify the contributions to measurement uncertainty and make a reasonable estimation to ensure that the form of reporting the result takes into consideration any applicable measurement uncertainty. When evaluating measurement uncertainty, all contributions that are of significance, including those arising from sampling, shall be taken into account using appropriate methods of analysis. Where the test method precludes rigorous evaluation of measurement uncertainty, an estimation shall be made based on an understanding of the theoretical principles or practical experience of the performance of the method.

13.15.2 Definitions

13.15.2.1 Significant Figures
Significant figures are those digits between and including the least and most significant digits in a number. The leftmost nonzero number is the most significant. The rightmost nonzero number is the least significant digit. If a decimal point is in the number, the rightmost digit is the least significant even if it is a zero.

13.15.2.2 Bias
The difference between a measurement result and the true or target value of the property being measured. The bias can be absolute or relative. The bias quantifies the accuracy of the measurement.

13.15.2.3 Coefficient of Variation (cv)
The relative standard deviation expressed as a percentage; another way to quantify the precision of measurement.

13.15.2.4 Decision Rule
A rule that describes how measurement uncertainty is accounted for when stating conformity with a specified requirement

13.15.2.5 Measurement Uncertainty
The property associated with a measurement result that characterizes the dispersion of the values that could reasonably be attributed to the true value being measured.

13.15.2.6 Standard Uncertainty
The uncertainty of a measurement result expressed as a standard deviation.

13.15.2.7 Expanded Uncertainty
A multiple of the standard uncertainty which provides an interval within which the true quantitative result is expected to lie with a stated level of confidence. For a multiple of
k=2, the interval will yield approximately 95% confidence that it contains the true property being measured.

**13.15.2.8 Confidence Interval**

A confidence interval gives an estimated range of values which is likely to include an unknown population parameter, the estimated range being calculated from a given set of sample data (Definition take from *Valerie J. Easton and Johns H. McColl’s Statistics Glossary v1.1*).

**13.15.3 Procedure**

Measurement Uncertainty is a parameter associated with a measured result that characterizes the possible range of values that could, under a specified level of confidence, be attributed to the result or method. In other words, the Measurement Uncertainty is used to indicate the degree of variability, at a specified level of confidence that can be expected for that particular measurement or method.

Measurement uncertainty takes into consideration all the potential variables that contribute to the measured result. Sources contributing to the uncertainty may include, but are not limited to, the reference standards or materials used, the procedure or equipment used, the environmental conditions, the properties or condition of the item being tested and the analyst performing the test. All components that may contribute to the measured uncertainty will be taken into consideration when estimating the measurement uncertainty.

Quantitative measurements: the measurement uncertainty shall be determined for quantitative measurements, such as weights of controlled substances.

Qualitative procedures such as identifying the presence or absence of a controlled substance or biological fluid, firearm operability and firearm/tool mark identifications (bullet or cartridge comparisons, etc.) do not require an estimate of measurement uncertainty.

The following test procedures have been identified as requiring an estimate of measurement uncertainty:

- Firearm barrel length determinations
- Marijuana weights
- THC concentration
- Measured trajectory angles

The affected functional area technical manuals will detail the procedures describing how the measurement uncertainty is calculated and how it must be applied when reporting the result. The procedures for evaluation of measurement uncertainty shall:
require the specific measuring device or instrument used for a reported result to have been included in or evaluated against the estimation of measurement uncertainty for that method;
include the process of rounding the expanded uncertainty;
require the coverage probability of the expanded uncertainty to be a minimum of 95.45% (often referred to as approximately 95%); and
specify the schedule to review and/or recalculate the measurement uncertainty.

13.15.4 Significant Figures and Truncating Values
The number of significant figures must, at a minimum, correspond to the uncertainty in the measurement and must not be more than the precision of the measuring device. If truncation is required for reporting purposes, truncation will occur after calculation to the appropriate significant figures and calculation of the measurement uncertainty.

13.15.5 Reporting Measurement Uncertainty
When measurement uncertainty is required, the examination documentation must contain the uncertainty of measurement or a reference to it. When this measurement uncertainty is of significance to the requestor or if it impacts the evaluation of a specification limit stated by a regulatory body, statute, case law, or other legal requirement, the range of values and the attendant uncertainty will be reported with specific confidence limits. Reports of analysis shall not overstate certainty of findings. The reported measurement uncertainty statement shall:

- include the measured quantity value, y, along with the associated expanded uncertainty, U, and the coverage probability;
- be in the format of y ± U with the units of y and U being consistent;
- limit the rounded expanded uncertainty to at most two significant digits, unless the laboratory has a documented rationale for reporting additional significant digits; and;
- require the rounded expanded uncertainty be reported to the same level of significance as the measurement result.

13.15.6 Measurement Uncertainty Records
The following records shall be maintained for each evaluation and estimation of measurement uncertainty:

- statement defining the measurand;
- statement of how traceability is established for the measurement;
- the equipment (e.g., measuring device[s] or instrument[s]) used;
- all uncertainty components considered;
- all uncertainty components of significance and how they were evaluated;
• data used to estimate repeatability, intermediate precision, and/or reproducibility;
• all calculations performed; and
• the combined standard uncertainty, the coverage factor, the coverage probability, and the resulting expanded uncertainty.
14 ACQUISITION OF SERVICES, SUPPLIES AND EQUIPMENT

14.1 POLICY

All purchasing, ordering and payment procedures will comply with WSP Budget and Fiscal Services (BFS) requirements. Such requirements are set forth in the Budget and Fiscal Services SOP and are found on the BFS Intranet website. The CLD is responsible for the acquisition, custody and disposal of all property within its control; therefore the CLD should only acquire property necessary to fulfill its mission. Division equipment and property will not be used for personal purposes.

Supplies and services that affect the quality of tests shall be selected and purchased at a quality appropriate for the analysis. Equipment will be selected on the basis of its appropriateness for specific functions, initial cost, ongoing support costs, and the availability of funds for equipment purchases and maintenance.

Each functional area or discipline shall maintain specifications and procedures for selection, evaluation and verification of supplies, services and equipment that affect laboratory testing within their technical procedures. Reference standards, reference materials, and calibrations of equipment/reference standards used to establish and/or maintain measurement traceability shall be viewed as critical, affecting the quality of the tests.

The laboratory shall communicate its requirements to external providers, as applicable, for:

- the products and services to be provided;
- the acceptance criteria;
- competence, including any required qualification of personnel;
- activities that the laboratory, or its customer, intends to perform at the external provider’s premises

Records shall be kept that demonstrates the receipt of supplies to include the ordering and acquisition date, and the receiver. Each laboratory shall ensure that standards, controls and reagents used in technical procedures are inspected or otherwise verified as complying with standard specifications or requirements defined in the methods for the tests concerned, or tested prior to use. Other recognized standards, e.g., testing of the delivered product or service, may be used at the discretion of the laboratory.

The laboratory shall evaluate all suppliers of materials (reagents and supplies), services and equipment affecting the quality of tests ensuring that specific requirements and standards of quality are met. A list of evaluated and suitable approved suppliers shall be maintained by the QP Manager, along with their record of compliance with established specifications.
### 14.2 Procedure

#### 14.2.1 Ordering and Purchase Approval

Data describing the type, class, grade, precise identification, specifications or other technical data including quality required of supplies, services and equipment to be ordered will be reviewed prior to purchase to ensure that the quality of the supplies, services and equipment are appropriate for the analysis.

Only CLD approved vendors for the purchase of supplies and services that affect laboratory testing will be used.

An order will be placed with a supplier only after the supervisor, laboratory manager or their designee has authorized the order in writing or by email. Prior to placing an order, it will be assigned a purchase order number or other approved means of payment to be provided to the vendor if needed, and then used for tracking the order. A system shall be used for monitoring supply orders.

While the CLD Commander is responsible for the CLD budget, laboratory managers are authorized to approve purchases totaling up to $5,000 per order, and supervisors are authorized to approve purchases totaling up to $1,500 per order. (Shipping, handling, and taxes are not included in the limit). Persons who are designated to be in charge may sign for the manager by using the following:

_________________________ by _________________________

Manager’s Name                       Signature of Person in Charge

Laboratory Managers and/or office managers will designate:

- Person(s) responsible for placing orders;
- Person(s) responsible for receiving orders and verifying that they are complete and correct;
- Person(s) responsible for tracking orders from time of placement through preparation of payment vouchers.

Laboratory managers must ensure that payment documents are prepared, and that any other purchasing-related responsibilities are fulfilled. Detailed instructions for preparing the various payment documents are included in the Budget and Fiscal Services Procedures Manual.

#### 14.2.2 Receiving Supplies and Services

Upon receipt, supplies, reagents or services will be checked or verified as complying with the purchase request. This can be done by checking the packing slip against the purchase request and against what was actually received to ensure all are in agreement.
If the shipping documents or labels do not match, the supplies or material will not be placed into service until the problem is resolved. Any discrepancies in the order will be recorded on the order documents and resolution sought. In addition, if the resolution includes returning the item, this will be noted on the shipping documents.

The person receiving the material will indicate the following information on the packing slip or receipt:

- The date received (Example: “Rcv’d 1/10/17”)
- An indication the appropriate item and quantity were shipped
- Approval will be indicated by the receiver’s initials
- The packing slip or receipt will be attached to the order document. Both will be retained for a minimum of one year in the laboratory for future reference.

Purchased supplies will not be placed into service or directly provided to the customer until they have been verified as meeting requirements, and documented as verified, per procedures described here or in technical procedures.

On receipt of vendor equipment calibration certificates, a scientist or technician will review and verify that the calibration certificate meets the specifications required. This review will include a check of the recalibration date and equipment identification. Unless specified otherwise in technical procedures, the first page of the calibration certificate will indicate by signature or initials and date that the calibration certificate meets the specifications. These records will be retained by the respective laboratories.

All equipment will be kept secure from damage, misuse, misappropriation, and theft. All equipment must be maintained in proper working condition. Equipment needing repair must be brought to the attention of the section supervisor who will inform management as necessary.

If a product or service that affects the quality of their tests has been put in use is found to be defective (e.g., not the expected quality), resolution shall be sought and the following shall occur:

- The supervisor or technical leader will assess the item/product for suitability.
- If the product/item has or may damage instrumentation or a process, then the supervisor/technical lead will immediately contact the QP Manager who will alert all possible users.
- A supervisor/technical lead will assess the damage and contact the responsible company for replacement of the product/item and/or possible reimbursement for damages.
- Review of any cases that may have been affected will be conducted. See the section on Nonconforming Work and Corrective Actions.
- The laboratory manager will keep a record of any defective products and take this into account when preparing their review of suppliers as well as considering any future purchases.
• CLD personnel have the responsibility to inform their immediate supervisor of a problem with product or services received from a vendor.
• Documentation of contact with the vendor will be maintained as appropriate.

14.2.3 Storage of Reagents and Laboratory Consumable Supplies
At a minimum, reagents and lab consumable supplies should be stored according to manufacturer/vendor recommendations. See the CLD Safety Manual for guidelines.

Each Laboratory Manager or their designee shall maintain a computerized inventory of all chemicals kept in the laboratory. That inventory shall be reviewed and/or updated once per year. Safety Data Sheets (SDS) shall be readily available to all personnel.

14.2.4 Vendor Evaluation
The list of approved suppliers of reagents, supplies and services that affect the quality of testing is on the FLSB Portal. A vendor evaluation for new suppliers of reagents, supplies and services, including proficiency test providers, shall be conducted by the purchaser with information provided to the QP Manager or designee. The vendor evaluation should be based on the following criteria:
• The vendor is currently an ISO certified supplier, is ISO registered or can demonstrate ISO compliance.
• Quality of product/service provided by vendor as related to documented requirements in discipline-specific technical procedures or quality manuals and the Quality Operations Manual.
• Conforms to recognized standards for providing quality goods and services to the State of Washington.

This conformance can be determined by examining the record for past successful performance for the individual laboratory, the laboratory system, or to other government entities. A record of successful past performance meets the requirement for conformance to recognized standards for providing products and services of acceptable quality to the CLD.

A Vendor Approval request, copies of national accreditation documents or a memo covering these points for each vendor shall be prepared by the purchaser and forwarded to the QP Manager. This information may be transmitted electronically.

Vendors providing Calibration Services or Certified Reference Materials will be evaluated annually to ensure current accreditation or certification and appropriate Scope of Accreditation. The Quality Process Manager will be responsible for these annual evaluations.
14.2.5 Acquisition and Retention of Donated Items

Donated items, which may include items that were previously evidence, may be acquired for laboratory purposes but must comply with the following rules:

- Evidence donated to the CLD must be accompanied by written disposition from the donating agency;
- The documentation will include the following information:
  - Date of transfer
  - Name of person releasing property,
  - Name of employee acquiring property,
  - Description of the item(s) including identifiers (serial number, etc.) if applicable,
  - Signature of person acquiring property,
  - Purpose for acquiring the property
- This documentation must be retained until the item is disposed of.

Copies of this receipt will be made available to all parties in the acquisition. Receipts will be retained until the item is disposed of or consumed.

14.3 Secondary Drug Reference Materials

The retention of secondary drug reference materials from casework samples or other non-certified sources will be allowed when there is no primary source reference material commercially available, or when there is a sufficient quantity of material in the case item such that taking a small sample will not consume more than half the original sample amount. These drug samples will be used primarily for training purposes.

To retain a sample from casework, the following procedure must be followed:

- The section supervisor must approve taking a portion of the casework material and the requesting scientist must obtain written permission from the submitting agency prior to removal of the sample. The written permission becomes a permanent part of the case file;
- The sampling must be witnessed by another scientist and will be documented in the examination documentation which will show the initials of the scientist and witnessing scientist, date, item number, and amount of sample removed. A notation will be made in LIMS regarding the sample having been removed from the drug item;
- The secondary drug reference material must be documented in the drug database and must be verified in the same manner as primary drug reference materials before being used in casework. The verification procedure is detailed in the Drug Reference Materials section of the Materials Analysis Technical Procedures Manual. A copy of the written permission from the submitting agency to remove the material from the case will be retained with the verification data.
14.4 **TRANSFER AND DISPOSAL OF PROPERTY/EQUIPMENT**

Transfer and/or disposal of property/equipment will follow policies established by the WSP Asset Inventory Control Officer and WSP Regulation Manual. Retention of any equipment no longer serviceable (such as instruments retained for parts) must be approved by the Laboratory Manager.
15 INVENTORIES AND REFERENCE COLLECTIONS

In order to facilitate daily operations and ensure quality compliance certain inventories and reference collections need to be maintained.

Inventories are subject to audits and as such need to be documented and controlled.

Reference collections shall be maintained, fully documented, uniquely identified and protected from unauthorized access, tampering and loss.

15.1 INVENTORIES

15.1.1 Inventory of Keys/Key Log

Each Laboratory Manager, or designee, shall conduct an inventory of the keys to the laboratory to verify the accuracy of the key log records and correct any discrepancies. The inventory will be conducted annually. A copy of the verified inventory shall be retained for review during the annual audit of that laboratory.

15.1.2 Inventory of Equipment and Instruments

The accountability for control of equipment and supplies in a laboratory lies with the Laboratory Manager. Inventories of laboratory equipment are maintained by WSP Supply and periodic inventory audits will be required per Supply’s schedule. A copy of the verified inventory shall be retained for review during the annual quality audit for that laboratory.

Each Laboratory Manager shall insure that State Identification Number Tags appear on fixed asset items which require such tags, as set forth by the WSP Supply Section. If a laboratory receives such an asset but it does not have a tag, the Laboratory Manager will request a tag from the WSP Supply Section. All fixed assets are subject to inventory.

15.1.3 Inventory of Library Materials

The accountability for control of library materials rests with the FLSB Librarian. The FLSB Librarian will maintain a current, updated inventory housed within the laboratory. Library material information will be entered in the Bureau Library Database in the FLSB Portal when received.

15.1.4 Chemical Inventory and SDS Check

The accountability for control of chemicals in a laboratory lies with the Laboratory Manager. To that end, Laboratory Managers, or their designees, will maintain a current database of all chemicals in the laboratory. The database will include approximate quantities of chemicals present along with their location. See also section on Purchasing Critical Services and Supplies.

Each Laboratory Manager, or designee, shall conduct an annual inspection of the database to verify all chemicals have been entered into the database. A copy of the verified
inspection, including the date of inspection, shall be retained for review during the annual audit for that laboratory.

Chemicals no longer needed will be discarded. The section supervisor, in conjunction with the laboratory safety officer, will be responsible for proper disposal of all chemicals. SDSs will be retained for all chemicals on site, and chemicals will be stored according to their safety requirements.

15.1.5 Vehicles

The accountability for control of vehicles assigned to a laboratory lies with the Laboratory Manager. To that end, each Laboratory Manager, or their designee, will regularly maintain the lab vehicles according to the WSP Vehicle Maintenance Schedule or according to the manufacturer’s service recommendations, to ensure the estimated life expectancy of the fleet. Regulations pertaining to fleet vehicles assigned to the laboratory are found in the WSP Regulation Manual Chapter 17 and on InsideWSP under the Fleet Section.

Vehicles are tracked and monitored by the Fleet Section. Maintenance records will be retained in each laboratory as needed. Managers shall be responsible for monitoring vehicle repairs and costs by periodically reviewing maintenance files and ensuring accuracy is maintained. A copy of all invoices and purchasing card receipts for vehicle maintenance and repairs will be entered into BMC Remedy in the Vehicle File system as described in Fleet Section procedures.

15.1.6 Special Reference Collections

The CLD maintains several reference collections, including ignitable liquids, minerals, paints and fibers. Specific collections requiring annual audits are described below. Discrepancies found during audits will be documented and addressed with appropriate correction and/or corrective actions without undue delay.

15.1.6.1 Drug Reference Materials Collection

The policies and procedures regarding the Drug Reference Materials collection including annual inventories are described in the Drug Materials Standards section of the Materials Analysis Technical Procedures Manual.

15.1.6.2 Firearms Reference Collections

The policies and procedures regarding the Firearms Reference Collections, including annual inventories, are described in the Firearms/Toolmarks Technical Procedures Manual.

15.2 CONTROL OF OTHER VALUED GOODS

15.2.1 Travel and Purchasing Credit Cards

All Travel and Purchasing credit cards shall be under the control of Laboratory Managers or their designees, and kept in secure locations within the laboratory. Credit card use will be documented, with the receipts for all purchases signed by the user pursuant to WSP
Budget and Fiscal Services SOP guidelines. Travel cards may be assigned to specific individuals when appropriate.
16 AUDIT PROGRAM AND MANAGEMENT SYSTEM REVIEW

16.1 POLICY

The audit program for the CLD shall be implemented in order to determine whether the management system:

- conforms to:
  - the laboratory’s own requirements for its management system, including laboratory testing;
  - the requirements of ISO/IEC 17025:2017 and ANAB AR 3125;
- is effectively implemented and maintained.

The audit program for the CLD encompasses a variety of quality measures. All laboratories will be audited annually to verify that operations are in conformance with established CLD policies, accrediting body requirements, any supplemental document requirements, the FBI Quality Assurance Standards (QAS) for Forensic DNA Testing Laboratories, FBI QAS for DNA Databasing Laboratories, and applicable WSP policies, rules and regulations. Internal and external audits will be documented and documentation will be retained for at least one cycle of accreditation (four years).

In addition to the annual internal quality audit, an annual Management System Review of the CLD management system’s operations for the previous year will be conducted to verify that the management system conforms with established CLD policies, accreditation requirements, and applicable WSP policies, rules and regulations. The management system and testing activities will be reviewed to ensure their continuing suitability, adequacy and effectiveness and to introduce necessary changes or improvements.

The audit program also includes:

- Court testimony monitoring
- Review of completed annual requirements
- Evidence Audits
- Drug Reference Materials Collection audit
- Firearms Reference Collection audit
- Inventory of Keys/Key Log
- Chemical Inventories
- Safety audits

This program takes into consideration the importance of the laboratory activities concerned, changes affecting the laboratory, and the results of previous audits.

Audits and the Management System Review afford opportunities for the CLD to consider the risks and opportunities associated with the laboratory activities in order to:
• give assurance that the management system achieves its intended results
• enhance opportunities to achieve the purpose and objectives of the CLD
• prevent, or minimize adverse impacts and potential failures in the laboratory activities, including as it relates to personnel health and safety
achieve improvement

The Management System Review, internal and external audit reports along with any nonconformance and corrective action plans will be documented and retained at least through one cycle of accreditation (four years).

Additional audits, such as a focus review, may be requested by the Laboratory Manager, SAS Manager, Quality Assurance Manager, CLD Commander or the FLSB Director at any time. The DNA Technical Leader or discipline Technical Leads with SAS Manager’s approval may conduct on-site technical audits to make recommendations for improvement in a specific discipline.

16.2 Procedures

16.2.1 Internal Audits
Each laboratory will undergo an annual internal audit. The scope focus areas of the annual internal audits vary over a four year audit cycle to ensure that all aspects of the management system are audited. SAS staff will lead the internal audit with designated lab managers as assistant lead auditors as needed. Together or as assigned, they have the responsibility to:

• provide an audit plan to the lab being audited
• organize and lead the internal audit
• ensure the audited lab manager is kept informed throughout the entire internal audit process and that a common understanding of the final audit report IOC is reached
• submit an audit report IOC addressed to the SAS Manager with copies to the CLD Commander, Quality Assurance Manager and lab manager of the audited lab.
• provide supporting audit documentation to SAS
• initiate nonconforming work corrective action requests in order to allow appropriate correction and corrective actions without undue delay

The Quality Assurance Manager will submit internal audit reports to the accrediting body by the due date established by the accrediting body.

Audits will include on-site inspections of laboratory facilities. Audit teams will use interviews, observations, direct witnessing, and documentation and technical file reviews in order to confirm the laboratory’s conformance to CLD and accreditation requirements. This includes an evaluation of the staff’s technical competence to conduct activities under the lab’s scope of accreditation. Audits will include a review of the previous year’s
corrective actions to confirm implementation, effectiveness and continued conformance. Each internal audit shall include case file review and direct observation of a sample of accredited services within each discipline accredited within the laboratory, and for which the lab is seeking accreditation, by technically qualified auditors. Technical auditors shall have experience in the discipline being audited to include successful proficiency testing within that discipline within a 4 year period. Labs with multiple categories of testing within an accredited discipline shall have each category audited over a four year internal audit cycle.

Audits will be conducted by trained, qualified personnel who are, wherever resources permit, independent of the activity to be audited. Auditors may come from the CLD or from outside the Division.

The SAS Manager is responsible for assuring internal audits and other audits are planned, organized and completed as required by the schedule. The SAS will define the audit criteria and scope for each audit. The DNA Technical Leader organizes and directs the DNA internal QAS audits. The Quality Assurance Manager will have oversight of nonconformances, CARs and follow-up activities.

A summary report of internal or external audits of the DNA functional area and CODIS will be prepared by the DNA Technical Leader and submitted to the CLD Commander.

16.2.2 Management System Review
An annual management system review of the CLD management system’s operations for the previous year will be conducted in the first quarter of the calendar year.

Fifteen Elements of the Management System Review (MSR)
The annual MSR will address the following points:

1. The suitability of policies, procedures and management system
2. Reports from managerial and/or supervisory personnel
3. A review of the annual internal laboratory audits
4. Corrective and preventive actions taken in the last year
5. Assessments performed by external organizations
6. The proficiency testing program and its outcomes
7. Changes in internal and external issues that are relevant to the CLD
8. Changes in the volume and type of work
9. Customer and personnel feedback
10. Quality system complaints
11. Status of actions from previous management reviews and results of risk identification
12. Effectiveness of implemented improvements
13. Other relevant factors such as quality control activities, adequacy of resources, and staff training, health and safety
14. A review of the fulfillment of and overall objectives of management system policies

15. Identification of risks and opportunities

The results of the MSR will be considered by the CLD Commander for risks and opportunities and for planning purposes. Items from the MSR used for planning purposes will have goals, objectives, and action plans.

A summary report IOC of the annual MSR will be prepared by the SAS Manager and directed through the CLD Commander to the FLSB Director with courtesy copies to all CLD lab managers. The summary report IOC will include all decisions and actions in regard to:

- Each of the Fifteen Elements of the MSR
- Evaluation of the effectiveness of the management system and its processes
- Risks and opportunities identified during the MSR
- Recommendations for improvement in laboratory operations
- Provisions of required resources
- Any other changes needed

The SAS Manager will follow-up on the status and outcomes of the summary report prior to the next year’s MSR with an IOC directed through the CLD Commander to the FLSB Director with courtesy copies to all CLD lab managers.

Implementation of corrective actions initiated from the MSR will be followed up to monitor effectiveness by the SAS Manager within an appropriate timeframe determined by the lab managers during the MSR and also reviewed at the following year’s MSR.

16.2.2.1 Audit and MSR Corrective Actions

Any findings of nonconformance from the annual MSR and external or internal audits will be addressed through the Corrective Action Process. Documentation of the finding(s) and corrective action plans are recorded in QPIT, unless otherwise required by an accrediting body.
17 ASSURING THE QUALITY OF TEST RESULTS

17.1 POLICY

The CLD is committed to providing the best quality service available to all members of the criminal justice system. A key component to providing high quality service is through a documented proficiency testing program. While proficiency testing is an integral part of an effective quality assurance program, it is not the sole indicator of satisfactory performance or delivery of a quality product. Proficiency testing does not replace high-quality work, standards, controls, and other conventional quality assurance practices.

The CLD may use, but is not limited to the following for monitoring the validity of tests performed:

- Use of certified or secondary reference materials and collections
- Positive and negative controls
- Orthogonal methods, such as use of alternative instrumentation that has been calibrated to provide traceable results
- Functional check(s) of measuring and testing equipment
- Use of check or working standards with control charts, where applicable
- Intermediate checks on measuring equipment
- Replicate tests or calibrations using the same or different methods
- Correlation of results for different characteristics of an item
- Intralaboratory comparisons (internal proficiency tests)
- Repeat testing (re-examination)
- Technical reviews, including inter-lab technical reviews
- A documented proficiency testing program (which may include testing of blind samples)
- Internal audits

The monitoring will be planned and any resulting data will be recorded and reviewed.

Other sections of this manual cover the variety of means to monitor and assure the quality of the test results: this chapter addresses performance monitoring by internal and external proficiency testing (intralaboratory and interlaboratory comparison).

17.2 PROFICIENCY TESTING

17.2.1 Definitions

17.2.1.1 Approved Proficiency Test Provider

An individual, organization or company that is accredited to ISO/IEC 17043 by an accreditation body that is a signatory to the APLAC MRA or IAAC MLA and has the applicable proficiency test(s) on its scope of accreditation, or where not available or not
appropriate for the testing conducted, gain approval from ANAB for alternative means by which the laboratory’s performance can be assessed.

17.2.1.2 External proficiency testing

Interlaboratory comparison: Organization, performance and evaluation of measurements or tests on the same or similar items by two or more laboratories in accordance with predetermined conditions.

17.2.1.3 Internal proficiency testing

Intralaboratory comparison: Organization, performance and evaluation of measurements or tests on the same or similar items within the CLD in accordance with predetermined conditions.

17.2.1.4 Proficiency testing program

Program by which performance monitoring by internal and external proficiency testing is implemented.

17.2.1.5 Proficiency Test

A proficiency test is an internal or external test that is provided to evaluate the capability of analysts, technical support personnel and the quality performance of a laboratory.

17.2.1.6 Proficiency Review Committee (PRC)

A committee of individuals appointed by ANAB, because of their experience and expertise, to provide oversight for ANAB in the proficiency testing program for specific forensic disciplines.

17.2.1.7 Proficiency Test Evaluation Form

The form used to provide comments on an individual’s internal or external proficiency test.

17.3 PROFICIENCY TESTING PROGRAM

The CLD proficiency testing program will be directed by the Quality Process Manager and shall be in conformance with the CLD accrediting body’s performance monitoring by comparison program. Appropriately accredited approved proficiency test providers will be used where available. Before ordering proficiency tests, the Quality Process Manager will confer with the laboratory managers and supervisors of each laboratory to determine the numbers and types of tests needed.

The CLD proficiency testing program requires that expected internal and external proficiency test results are not known or readily available to the test taker. Each laboratory must successfully complete, per calendar year, at least one external proficiency test with authorized release of the test results to ANAB from the test provider for each discipline in which it provides accredited services. If application for accreditation has been made for a discipline (for example, scope expansion), the laboratory will successfully complete at least
one external proficiency test for each discipline in which application for accreditation has been made.

Each scientist/technician within the CLD will successfully complete at least one internal or external proficiency test per calendar year in each discipline (as listed on the scope of accreditation) in which the scientist performs casework. Scientists in the DNA and CODIS functional areas and other personnel designated by the DNA Technical Leader will successfully complete two external proficiency tests each year. Individuals involved only in serology screening for biological evidence will undergo annual external proficiency testing. Individuals involved only in outsourced DNA analysis technical review will complete two external proficiency tests each year. Additional information for DNA proficiency testing is described in the DNA Quality Assurance Manual.

Additionally, each person conducting testing must have all aspects of casework on their laboratory’s scope of accreditation in which they perform testing monitored at least once in a four year accreditation cycle. In the event that interlaboratory or intralaboratory proficiency testing options are not available or appropriate, documented observation-based performance monitoring by a technically competent individual is acceptable.

DNA analysts and technical support personnel performing DNA analysis shall comply with proficiency test requirements of the Quality Assurance Standards for Forensic DNA Testing Laboratories and Quality Assurance Standards for DNA Databasing Laboratories, as appropriate.

Each laboratory shall have a proficiency testing plan, overseen by the Quality Process Manager, that demonstrates conformance with the proficiency testing requirements stated above and ensures inclusion of a representative sample of the types of tests within each discipline listed on the laboratory’s scope of accreditation.

The objectives of the proficiency testing program are to:

- Demonstrate the current competence of the scientist and technical support personnel
- Demonstrate the current competence of the laboratory
- Ensure that quality work is being maintained
- Identify areas where additional training or resources would be beneficial
- Verify the validity of technical procedures

17.4 Procedure

17.4.1 Proficiency Test Samples

Proficiency test samples will be handled in the same manner as case evidence until the Quality Process Manager determines that all proficiency test requirements have been satisfied and the sample is no longer needed for that purpose. Proficiency test samples received from an external proficiency test provider will be deemed sealed. The sample
may then be kept as a training sample, or it may be destroyed as determined by the supervisor or Technical Lead. The final disposition of the sample must be documented by the supervisor in LIMS. An exception to the documentation of final disposition would be those proficiencies in which a physical sample does not exist, such as data found in a website link.

For internal proficiency tests samples, the CLD may use internally created practical tests, previously worked or older unworked commercially provided practical tests, testing reanalysis, external tests whose results are not submitted to the test provider or not authorized for release to ANAB and when appropriate, observation-based tests. When an internal proficiency test sample, including an observation-based test, is internally created or is a previously used proficiency test sample being used a second time, the quality of the test sample must be ensured prior to issuing the test. The quality of a previously used test sample is assessed by the test sample provider to ensure the used test is in good condition, has not exhibited deleterious change, and is of sufficient quantity. Results of the first and second testing will be compared and evaluated as described below: if there is a significant difference between the analyst’s proficiency test results, the reason will be investigated with appropriate follow-up action (such as another test) taken. For test samples internally created but not previously tested or used as part of a testing process, the quality of the test sample shall be confirmed by examination of the test sample by a scientist authorized to perform the testing.

17.4.2 Proficiency Testing Process

The Quality Process Manager will keep records regarding how the test samples are obtained or prepared, as well as completion dates and results of the testing. As tests are received, the Quality Process Manager will disperse the necessary tests to appropriate labs. The laboratory manager and/or supervisor are responsible for assigning proficiency tests to their scientists as needed. The Quality Process Manager will ensure that appropriate proficiency test samples are obtained, assigned, and provided to each scientist with enough advance notice to allow completion prior to deadlines.

Proficiency tests must be completed and the results submitted to the test provider within the timeframe imposed by the provider. This requirement is essential to the overall success of the CLD Proficiency Testing Program; therefore it is the responsibility of the scientist assigned the test to ensure that this requirement is met.

Management has the responsibility to see that the proficiency test is assigned to and received by the scientist in a reasonable time frame. The scientist must perform the testing so that there is sufficient time to accomplish appropriate reviews for the test results to be sent to the test provider by the due date.

The scientist will document and report to their supervisor, laboratory manager, and the Quality Process Manager if a proficiency cannot be completed by the deadline. It is the
responsibility of the laboratory managers and supervisors to ensure that proficiency results are completed and returned to the test provider.

Proficiency tests must be completed in the same manner as casework using approved test methods and be consistent with the respective analysis and reporting procedures of the category of testing. The completed proficiency test provider answer sheet serves in lieu of a laboratory report and no additional laboratory report or draft report is necessary. The proficiency test results will undergo technical review and administrative review before results are sent to the test provider. The technical review will be documented in LIMS and by the reviewer’s initials and date on a copy of the answer sheet or in the electronic case file.

The administrative review will be documented in LIMS and in the case file. The administrative review documents that the answer sheet for the proficiency test has been fully completed and is free of errors.

Proficiency test case files contain a copy of the answer sheet with the assigned scientist identified, case numbers, date of completion, and documentation of technical review.

Copies of the answer sheets with the assigned analyst’s identity and other necessary paperwork, including confirmation of submission to the provider, for the proficiency test will be sent to the Quality Process Manager.

The Date of Completion for the proficiency test will be the date when the results/answer sheets are submitted to the proficiency provider. If the responder is not submitting to the proficiency provider, the completion date will be the receipt of the answer sheet by the Quality Process Manager.

The intent of the proficiency testing program is to identify individual technical issues and also systemic issues. Therefore, if a technical reviewer disagrees with the conclusions reached by a scientist, then it is incumbent upon the reviewer to bring the problem to the attention of the scientist and their supervisor. The procedure used to resolve technical review conflicts in actual casework as outlined in this manual will be followed.

When the results of the proficiency tests are received, the Quality Process Manager or designee will review the scientist’s and the provider’s results. The Technical Lead/DNA Technical Leader of each discipline will be involved, as much as possible, in the evaluation of the proficiency answer sheets for technical accuracy. The scientist’s results will be compared to the proficiency test provider’s results (manufacturer’s specification and answer information) and evaluated for technical accuracy and agreement. The scientist’s supporting documentation (notes) may additionally be compared and evaluated. The criteria for determining successful completion is whether the scientist’s results are technically accurate and consistent with the proficiency test provider’s results. For proficiency tests involving quantitative analyses, such as THC quantitation, results will be acceptable if the reported results are within two standard deviations of the assigned value.
of the analyte or within the satisfactory range as defined by the statistical analysis reported in the test result summary.

Proficiency test records will be maintained at CLD Headquarters. Proficiency test records include:

- Proficiency test unique identifier
- How tests were obtained or created (external, internal, observation-based)
- Discipline tested
- Written instructions for completion
- Identity of person taking the test
- Location where the proficiency test was taken
- Due date and completion date
- Copy of the proficiency test report (answer sheet(s)) and records submitted to the test provider
- Expected proficiency test results
- Copy of the proficiency test evaluation form
- Documentation that feedback was provided to the analyst
- Any discrepancies noted
- Details of corrective actions taken (when necessary)

The proficiency test records maintained with the case file shall also include all examination documentation supporting the results and conclusions.

Proficiency test records will be retained at CLD Headquarters for at least one full accreditation cycle. After this time they may be archived with the same agency retention schedule as case files.

Test items, except for controlled substances, may be stored in the appropriate functional area with other training materials. This will be documented in LIMS by releasing the item to a location outside of the evidence vault, within the laboratory.

Completed drug analysis proficiency samples, when stored outside of the vault, must be stored using the same precautions and levels of security used for other controlled substances such as drug reference materials. This will be documented in LIMS by releasing the item to a controlled storage location.

If the drug analysis proficiency samples are to be destroyed, the destruction of the samples must be documented using a WSP Property/Evidence Report form 3000-110-096. The item to be destroyed will be sealed in an appropriate package, assigned a WSP Property/Evidence number, and signed over to a WSP District PEC. Copies of the WSP Property/Evidence Report that contain the signature of the releasing scientist and the WSP District PEC will be retained for at least one accreditation cycle and the destruction of the samples documented in LIMS by transferring the evidence to the “destroyed” location.
17.4.3 Satisfactory Proficiency Test Results

If the test results are satisfactory, the Quality Process Manager will complete documentation of the satisfactory result in the records. Documentation of satisfactory completion will be issued to scientists, their laboratory managers, and supervisors. The documentation shall be initialed and dated by the analyst performing the proficiency test before inclusion in the case file.

17.4.4 Proficiency Test Discrepancies

If there is a discrepancy between the scientist’s test results and the provider’s results, the Quality Process Manager will enter into QPIT a Proficiency Test Inquiry (PTI) and ensure notification of the laboratory manager, supervisor, Technical Lead or DNA Technical Leader as applicable, and scientist who performed the test. The CODIS Lab Manager will additionally be notified if related to a DNA PTI. The laboratory manager may subsequently receive a Letter of Inquiry from the ANAB Proficiency Review Committee (PRC). The Letter of Inquiry shall immediately be forwarded to the Quality Process Manager. The SAS, in collaboration with the supervisor, Technical Lead/DNA Technical Leader, and/or laboratory manager will determine a course of action. The Management Liaison may also be involved if necessary.

Responses are required on all Letters of Inquiry and should be directed to the PRC contact person identified in the letters. A Request for Additional Information and the Referral Letter are communications that could be received after the PRC has reviewed the laboratory’s response to a Letter of Inquiry.

If there is a test response different from the expected and consensus response, the laboratory manager may receive a “Notification” from the ANAB PRC. The Notification does not require a response to the PRC. The purpose of the Notification is to ensure that the laboratory is made aware of the test response difference. The SAS Manager, in collaboration with the supervisor, Technical Lead/DNA Technical Leader, Quality Assurance Manager, and/or Laboratory Manager will determine a course of action. The Management Liaison may also be involved if necessary. There is no need for a laboratory to respond to a Notification unless the test results were not correctly transcribed and summarized by the proficiency test provider. The laboratory shall retain the Notification with the proficiency test file, the DNA Technical Leader (only if DNA related), and the SAS Manager.

If other quality/procedural concerns are raised during the proficiency process, the Quality Process Manager may enter and track a Proficiency Test Inquiry in QPIT for additional information and follow-up. The request for information will be provided to the scientist, supervisor, lab manager, appropriate Technical Lead/DNA Technical Leader, QAM and the SAS Manager. The Management Liaison may also be involved if necessary. The SAS will make the determination if a nonconformance has occurred and if a corrective action investigation is necessary. This approach is similar to the Letter of Inquiry or Notification Letter issued by the ANAB Proficiency Review Committee.
If a scientist’s performance on a proficiency test requires further development to meet quality standards, the SAS Manager, in collaboration with the scientist, supervisor, Technical Lead/DNA Technical Leader, and/or laboratory manager will determine a plan of action which shall include removal of the scientist from casework in the area in which the proficiency was performed and remedial training. The Management Liaison may also be involved if necessary. The SAS Manager will prepare a report to the CLD Commander outlining the issues and the actions taken. The SAS Manager or designee will then draft a response from the division to the PRC indicating the actions taken. The CLD Commander will review and approve the final draft. Once approved, the response will be sent to the PRC.

The proficiency test case file will contain a record of the discrepancy between the scientist’s test results and those of the test provider. The Quality Process Manager will retain complete records for the CLD.

17.4.5 Proficiency Testing and Job Performance

Satisfactory performance on a proficiency test should be documented in the employee’s supervisory desk file. Likewise, any problems identified from the review of a proficiency test, if reflective of difficulties with a scientist’s individual work performance, will be addressed by the supervisor and documented in the scientist’s supervisory desk file. The supervisor may enlist input and assistance from the Technical Lead or DNA Technical Leader, the SAS Manager, the Quality Assurance Manager, the Laboratory Manager, the Division Commander, and other appropriate individuals. See sections in this manual on Nonconforming Work and Corrective Actions and Documenting Job Performance.
18 TECHNICAL PROCEDURES AND METHODS

18.1 POLICY

The CLD will use appropriate technical procedures and methods that have been scientifically validated and accepted for use in the field of forensic science. This includes methods and procedures for the sampling, handling, transport, storage and preparation of evidence items, the operation of all relevant equipment having a significant effect on the accuracy or validity of the result of the test, calibration, or sampling, and an estimate of the measurement uncertainty where appropriate. All methods and procedures will be kept up to date, documented and readily available for review by laboratory personnel. If an issuing body revises a method, verification shall be repeated to the extent necessary. Any deviation from a standard technical procedure or method will require that the details of the modification as well as the justification and the authorization be documented in the examination documentation and maintained as a permanent part of the case file.

18.2 TECHNICAL PROCEDURES

Technical procedures must be based upon sound scientific principles and as effective and efficient as possible. Every procedure used must be generally accepted in the relevant scientific field. All test methods that involve the comparison of an unknown to a known shall require the evaluation of the unknown item(s) to identify characteristics suitable for comparison and, if applicable, characteristics suitable for statistical rarity calculations (DNA), prior to comparison to one or more known item(s). Characteristics and their comparison criteria are described in the discipline technical procedure manuals. This requirement is not focused on the process of screening an unknown in order to identify evidence or characteristics that may be the subject of further comparison. In these cases, it may be appropriate to perform a preliminary characterization of the known sample prior to the assessment of the unknown.

An additional criterion for the selection of a technical procedure is general acceptance by the appropriate functional area group within the division. Even though a technical procedure may have gained general acceptance within the relevant forensic science community, it must also be understood, supported and accepted by those who must employ that technical procedure in cases submitted to the division. Supervisors and Technical Leads will communicate the development plan and progress to the members of their functional area.

Technical procedures must be as well documented as possible. Documentation includes specific literature articles, texts, reviews, and data compilations. A Reference List may be included in either the technical procedures or the training manuals. The procedure should include:

- Definition of terms
- Literature references
- Scope of the analysis conducted
- Standards for notes, interpretation of results and reporting
- Minimum examination requirements
- Equipment/instrument specifications required
- Equipment/instrument operation, maintenance and verification procedures
- QA statement
- Safety statement
- Documented validation studies

Note: Methods currently in use that have been validated by the laboratory, that are of a long standing, historical nature, long predating this ISO-based version of manual, may not have validation documentation on file. The validity of these methods has been repeatedly demonstrated by past and present analytical results and accurate proficiency test results.

All technical procedures shall include provisions for the following:

- Quality control and quality assurance: this includes guidelines for negative controls, knowns, and calibrations, and how they should be reported.
- Test data interpretation
- Be applicable to the submitted item in order to conduct requested analysis
- Defining of ranges of the conclusions that can be drawn from the data
- For test methods involving the comparison of an unknown to a known requiring identifying characteristics for comparison, criteria for the evaluation of the characteristics
- If applicable, evaluation criteria for characteristics suitable for statistical rarity calculations
- Where safety issues exist, safety precautions specific to each technical procedure shall be included in the documentation and incorporated into the technical procedure. Safety should also be a major consideration in the development and acceptance of a technical procedure. Any precautions and limitations of the technical procedure must be documented in the technical procedures manual.

Procedural manuals will include procedures, or make reference to procedures, to ensure safe handling, transport, storage, use and planned maintenance of measuring equipment (i.e., balances, pipettes, calipers, trigger pulls, rulers) to ensure proper functioning and in order to prevent contamination or deterioration.

18.3 Using Technical Procedures

There are specific technical procedure manuals for each recognized forensic discipline. The official CLD controlled manuals are maintained electronically by the QPM and are readily available to all analysts and staff.
The items submitted to the laboratory and the information needed by the criminal justice system can vary greatly in different cases. If forensic scientists lack knowledge and/or the available procedures to process certain cases, then they will notify their supervisor who will either refer the analysis to another forensic scientist who has the necessary expertise, or consult available literature, colleagues, and other sources (such as manufacturers, universities, and other agencies) in order to obtain the needed data or technical methodology. If this is not successful because of limited resources, then the report should clearly explain the limitations of the work and that the conclusions were drawn from limited analysis.

Although many acceptable procedures may exist to perform a particular examination, considerable variations in case samples require that forensic scientists have the flexibility to exercise discretion in selecting the method most appropriate to the problem at hand.
19 METHOD VALIDATION

19.1 Policy

The laboratory shall use appropriate methods and procedures for all tests and test data interpretation within its scope and which meet the needs of the customer. Validated methods published in international, regional or national standards or by reputable technical organizations, or in relevant scientific texts or journals, or as specified by the manufacturer of the equipment, shall preferably be used. The laboratory shall ensure that it uses the latest valid edition of a standard method unless there is a documented reason not to do so and there are documented steps to ensure consistent application. The laboratory or discipline will ensure that all non-standard methods, laboratory-developed methods, standard methods used outside their intended scope, and modified methods that are significantly modified such that the change(s) affects the outcome of the test are properly validated prior to use. The range and accuracy of the values obtainable from validated methods (e.g. the uncertainty of the results, detection limit, selectivity of the method, linearity, limit of repeatability and/or reproducibility, robustness against external influences and/or cross-sensitivity against interference from the matrix of the sample/test object), as assessed for the intended use, shall be relevant to the customers' needs.

The validation will be as extensive as is necessary to meet the needs of the given application or field of application. All validation studies will be planned and performed with the aid of the Technical Lead or under the direction of the DNA Technical Leader by qualified personnel with adequate resources to perform the validation. Personnel authorized to perform casework are also authorized to develop, modify, verify, and validate methods within their area of authorization. Additional personnel can be authorized to do these functions on a case by case basis. The proper validation of a technical procedure requires an understanding of the theoretical basis for the technical procedure. Such knowledge provides a means of assessing the specificity and limitations of the technical procedure and predicting possible sources of error. Results of the validation will be documented and archived. Archival may be as part of the functional area technical manual or in a separate validation binder. The documentation will include the data, the procedure and controls or standards used, a statement as to whether the method is fit for the intended use, and documentation of approving authority.

Prior to implementation of a validated method that is new to the laboratory, the efficacy and reliability of the method shall be demonstrated in-house against all documented performance characteristics of that method. Records of performance verification shall be maintained for reference.

The guidelines below will be used for development of method validation plans.
19.2 **Procedure**

Prior to beginning any validation study, a validation plan will be prepared by the scientist involved and the discipline Technical Lead/DNA Technical Leader and submitted through the chain of command to the SAS Manager for review and approval. The selection of the appropriate type of validation should be part of the planning process. As the study progresses, the plan will be reviewed and updated as necessary. Any modifications to the plan shall be approved, authorized and documented. Effective communication among all personnel involved including other analysts in the section, the Technical Lead or DNA Technical Leader and the SAS Manager will be accomplished through verbal or written communications.

Method validations shall:

- Be a planned activity that is approved and authorized;
- Be conducted by competent, authorized personnel;
- Be reviewed regularly as the validation is carried out, and updated as needed, with any change in requirements requiring modifications to the development plan being approved and authorized;
- Encompass the test process to include data interpretation;
- Establish the data required to report a test result, opinion, or interpretation;
- Identify limitations of the test method, reported test results, opinions, and interpretations;
- Specify when a currently validated method, including associated data interpretation, needs additional validation; and
- Require a validation plan providing direction for parameter evaluation and parameter acceptance criteria to determine if the method is fit-for-purpose prior to starting a method validation.
- Consider costs, risks and resources.

Laboratory personnel wishing to introduce and validate a new technical procedure, or significantly modify an existing procedure such that the change(s) affects the outcome of the test, shall seek initial approval for development through their supervisor. When drafting a proposal the scientist, Technical Lead/DNA Technical Leader will include the following:

- The basis of the recommendation, including the objective, benefits to the laboratory system and relevance to the customer
- Who is expected to conduct the development and validation
- Where the work will occur
- Compatibility with evidence handling procedures
- Ability to quickly provide accurate data
- Ability of the procedure to produce the most discriminating information
- What equipment will be used
• Compatibility with available equipment, facilities and personnel
• The extent that the procedure destroys, consumes or alters evidence, including compatibility with other procedures which might be used before or after the method for confirmation or other purposes
• The validation plan for the method (see details below), test samples, standards, certified reference materials and/or reference materials to be used
• How each performance characteristic of the validation that is applicable is going to be evaluated and met
• The estimated costs associated with the development and validation, including supplies, materials and implementation
• Literature research: Review of publications, academic materials, safety procedures, protocols and manufacturer’s specifications, etc. involving the technique or procedure being validated
• Source of the technical procedure, if scientific in nature
• Safety Data Sheets (SDS) regarding any chemicals required for the technical procedure
• Projected schedule of periodic reviews to assess fulfillment of the objective
• Timeline for implementation

The proposal will be evaluated based on the above elements, customer needs, and availability of appropriate staff and adequate resources. If the proposal is authorized upon review by the SAS Manager, the validating laboratory will document the validation process. If the validation is successful, the documentation will be sent to the relevant Technical Lead or DNA Technical Leader for review. The Technical Lead or DNA Technical Leader will determine if the new technical procedure has been sufficiently validated and if it should be included in the procedures manual. The procedure will be formatted in preparation for inclusion in the technical manual following the document control process. A copy of the completed validation will be submitted to the SAS Manager.

For equipment method validation, a copy of the validation documentation and data must be kept with the equipment records. The Technical Lead or DNA Technical Leader will prepare an IOC directed to the section supervisor and/or Laboratory Manager, as appropriate, advising them that the equipment method has been validated and is ready for use in casework. By signing and acknowledging the IOC, the Laboratory Manager authorizes the placing of the equipment into service.

Method validation records shall be retained to include:
• the validation procedure used;
• specification of the requirements;
• determination of the performance characteristics of the method;
• results obtained; and
• a statement on the validity of the method, detailing its fitness for the intended use.

**19.2.1 Method Validation Plans**

In addition to the above considerations, method validation plans will include:

- Objective of the Method
- Equipment Used
- Testing Samples
- Reference Standards and casework samples: The samples used for validation should be representative of the types of casework samples and blanks routinely analyzed using the technique or procedure, and include standards, reference materials and/or certified reference materials. The technical procedure must be tested using known samples. If a new technical procedure is intended to supersede an existing procedure or if it parallels an existing procedure, then the results of both procedures should be compared. The known samples should be designed to resemble actual evidence materials as closely as possible, taking into account the following (as applicable) matrix, sample age, environmental effects, sample homogeneity and other factors as appropriate to the functional area. The performance characteristics of validated methods, as assessed for the intended use, shall be relevant to the customers’ needs and consistent with specified requirements.

- Applicable performance characteristics
  - **Accuracy**: The quality or state of being correct; the degree to which the result of a measurement, calculation or specification conforms to the correct value or a standard. Labs should test accuracy by using samples of known values or quality or by comparing results to references of known values or quality.
  - **Repeatability Precision**: A measure of the variability in results when a measurement is performed by a single analyst using the same equipment over a relatively short timescale (for example, about 6-15 repeats). Repeatability is expected to give the smallest amount of variation in results.
  - **Reproducibility Precision**: A measure of the variability in results between laboratories using the same method. Reproducibility is expected to give the largest variation in results. The method must be reproducible by another individual using the original test documentation.
  - **Specificity**: The extent to which the method can be used to determine particular analytes in mixtures or matrices without interferences from other components of similar behavior. Does the method provide results specific to the substrate tested (i.e., What is the occurrence of false positives?).
  - **Robustness**: A measure of the method’s capacity to remain unaffected by small, but deliberate variations in method parameters.
  - **Sensitivity or Linearity Studies**: This performance characteristic informs the working range of your instrument or test, the limits of detection and range of analysis (lower to upper limits). The goal is to determine a range of which a reliable result can be obtained that is above baseline and interpretable. This is
an important characteristic for monitoring quality control of an instrument or test. Is the sensitivity so great that many false positives occur? Is the sensitivity so low that many false negatives occur? Are you working within the linear range of detection? What are the detection limits of the method or instrument?

- **Ruggedness**: Assesses the factors external to the method, such as environmental temperature, lighting, humidity and human error. When applicable, evaluate the method using known samples under different environmental conditions.

- **Measurement Uncertainty**: This is a requirement for validations where the method requires measurements that will affect the test result. If the method provides reported quantitative data, the validation study must include an estimate of accuracy and precision at concentrations which are representative of casework samples expected to be encountered when using the method.

- **Sources of Error**: It is important to be clear about any sources of error that occur during the validation. These sources of error will each have to be addressed in the validation report. Known sources of error can come from scientific literature references used for the method that you are validating such as relevant journal articles

  - Timeline for completion
  - Projected schedule of periodic reviews to assess fulfillment of the objective and customer relevance
  - Documented, approved and authorized modifications to the plan
  - Documented technical review
  - Final report IOC summarizing the validation to include literature references, any sample selection plans and a statement that the method is fit for use.

### 19.3 PERFORMANCE VERIFICATION

Before being placed into service, equipment (including that used for sampling) shall be calibrated or checked to establish that it meets the laboratory's specification requirements and complies with the relevant standard specifications. It shall be checked and/or calibrated before use in the laboratory by authorized personnel. A performance verification tends to mimic an abbreviated validation and is meant to check the accuracy and reliability of equipment and methods. Functional area technical procedure manuals may have additional details on performance verification specific to their testing.

Performance verifications must be completed for the following:

- Methods that have undergone validation and implemented in the CLD laboratory;
- Newly purchased equipment of similar make and model or operating on the same principles or basic technologies as other equipment already implemented and used in the CLD;
- Equipment transferred from one lab to another;
- Regular verification of equipment currently in use to ensure that the instrument/equipment continues to function to manufacturer’s specifications and to in-house procedure specifications
- Previously validated methods (e.g. published standard methods) new to a laboratory;
- To the extent necessary, methods in use that have been revised by the issuing body;
- Other circumstances deemed appropriate by the Technical Lead, DNA Technical Leader or functional area technical procedure manuals.

**Performance verification procedures:**

- All verification results must be documented, maintained onsite, and will be subject to review during the annual internal audit.
- Verification must demonstrate that a representative set of reference materials has been carried through the process and yielded the expected results.
- The performance verification must have demonstrated in-house the reliability of the method against all documented performance characteristics of the method and shall be maintained for reference.
- The samples used for verification should be representative of the type of standards routinely used for controls and specimens routinely analyzed using the technique or procedure.
- Accuracy and precision studies to verify that the equipment or procedure is within previously established manufacturers or procedure specifications.
- Include estimates of measurement uncertainty for quantitative methods, as applicable.
- For equipment, a copy of the verification results and data must be kept in the equipment log.

### 19.4 Deviation from Technical Procedure

Deviations from the CLD Technical procedures may occasionally be justified.

#### 19.4.1 Deviation

Deviation is a pre-planned change or variation in a technical procedure.

#### 19.4.2 Policy

Any deviations from Division technical procedures must be approved in writing by the SAS Manager and functional area Technical Lead. If the functional area Technical Lead is not available, the appropriate management liaison may approve a deviation in consultation with the SAS Manager. In the DNA functional area, only the DNA Technical Leader’s approval (or those designated in the DNA Quality Assurance Manual) is required. The approval must be technically justified and documented in the case file. Deviation from test methods shall occur only if the deviation has been documented, technically justified, authorized, and accepted by the customer.
20 SAMPLING AND SAMPLE SELECTION

20.1 POLICY

The CLD laboratories typically do not engage in statistical sampling of evidence. Instead, analysts select samples for analysis, which may not be a true statistical representation of the whole. Where required, the Technical Lead, or the DNA Technical Leader where appropriate, with input from the supervisors and the Functional Area, will define a sampling plan and procedures for when a discipline must carry out a true sampling of substances, materials or products for subsequent testing. The sampling plan and procedures for sampling will be documented in the specific functional area procedure manual.

20.2 DEFINITIONS

20.2.1 Sampling
Taking a part of a substance, material or product for testing in order to reach a conclusion, make an inference about, and report on the whole

20.2.2 Sampling Plan
A statistically valid approach to determine the number of sub-items that must be tested in order to make an inference about the whole population.

20.2.3 Sample Selection
The practice of selecting a sample(s) of the whole based upon training, experience and competence, to draw conclusions and report only on the sample(s) tested.

20.3 PROCEDURE

As applicable, each discipline shall document in its technical procedures manual a sampling plan and procedure for items that require sampling prior to testing. Sampling plans shall, whenever reasonable, be based on appropriate statistical methods. The sampling procedure shall address the factors to be controlled to ensure the validity of the test results.

Proper sample selection of analytical materials from evidence will be a result of the individual scientist’s training and experience. Sample selection will also be addressed in the training manuals.

Sampling procedures should describe the sampling plan, selection, withdrawal, preparation and treatment of a sample or samples from a substance, material or product to yield the required information. The sampling plan and procedure(s) shall:

- Require an evaluation of the selected population for homogeneity;
- Require the population to have a reasonable expectation of homogeneity to use a sampling plan;
- Require that the sampling plan makes use of probability and provides an opinion or interpretation with a minimum confidence level of 95.45% (often referred to as approximately 95%);
- Require each item selected to meet the sampling plan level of confidence to be tested completely; and
- Provide instruction regarding the course of action to take if one or more selected items demonstrate a lack of homogeneity.

It is appropriate to choose a limited number of samples for analysis from multiple samples if those chosen are representative of the larger sample. Care must be exercised that the analytical results reported are only for the items analyzed and not representative of the whole. No conclusions may be drawn for the whole. For example, if two bags of white powder are analyzed from a submitted sample of 20 bags of white powder of similar appearance, then the report must only reference the results of the analysis of the two bags chosen. Otherwise, a sampling of the whole must be conducted and the sampling method documented, and the results may be applied to the entire quantity of material under investigation.

If the functional area utilizes more than one sampling procedure, the procedure used must be documented in the case file. Deviations from the sampling plan may be requested by the external customer or be deemed appropriate by the analyst. If the supervisor approves a deviation from the sampling plan then this shall be recorded in the case file and communicated to the DNA Technical Leader or applicable technical lead.

As appropriate, documentation of sampling shall include the following:
- Date and time of sampling
- Identification of the personnel performing sampling;
- Identification of the equipment used;
- Unambiguous description and identification of the substance, material or product sampled (such as number, amount or name)
- Location of sampling, including any diagrams, sketches or photographs
- Reference to the sampling plan and procedures used
- Details of any environmental or transport conditions during sampling that may affect the interpretation of the test results
- Any standard or other specification for the sampling method or procedure, and deviations, additions to, or exclusions from the specification concerned
- If necessary for the interpretation of test results and conclusions, the sampling plan used will be included in laboratory reports (see also the section on Laboratory Reporting of Sampling)
21 SUBCONTRACTING OF TESTS

21.1 POLICY

Each contractor or contract laboratory that is employed by the CLD for providing testing services or analysis or technical review will establish competency to perform the contracted work, including provisions for evidence security. A documented technical assessment will be performed, to include a review of their procedures to determine if they meet CLD standards by having the same or equivalent procedures before a contract can be approved. Contractors will review the WSP Regulation Manual and the CLD Quality Operations Manual and relevant Technical Manuals prior to beginning work. Documentation of competency shall be obtained or provided to the CLD prior to submitting samples to a contract laboratory. Contractors and contract laboratory employees designated by the CLD shall be participants in an on-going proficiency testing program provided by an approved outside provider.

All CLD laboratories that receive transfer or referral casework from another CLD laboratory are considered contract labs. All CLD forensic laboratories are considered competent labs and customer requests can be transferred to any WSP lab at the discretion of the individual laboratories.

21.2 DEFINITIONS

21.2.1 Competent Subcontractor

A competent subcontractor is one that is accredited to an appropriate international standard by an accrediting body with a scope of accreditation that covers the services being subcontracted and complies with CLD criteria for the work in question.

21.3 PROCEDURES

The CLD will advise the customer of the subcontracting arrangement in writing and gain approval of the customer, preferably in writing. Convicted offender DNA samples may be contracted out without notification to the collecting agency. DNA sample analysis contracted out by the customer can only be accepted for CODIS consideration if the contract between the customer and the external laboratory has been approved by the DNA Technical Leader prior to the start of analysis. Procedures for outsourcing of DNA analysis are described in the DNA Quality Assurance manual. Communication to the customer regarding the transfer or referral of casework to another CLD laboratory is not required but may be appropriate. If available, a subcontractor shall be accredited to ISO/IEC 17025 by an accrediting body that is a signatory to the ILAC Mutual Recognition Arrangement with a scope of accreditation that covers the testing services being subcontracted. Any specific requirements for establishing subcontractor competency are found in the procedures or quality manuals of the discipline requiring the subcontractor. Subcontractor performance will be evaluated and documented at least annually. Should performance or requirements
not be met, communication between the CLD and subcontractor will occur and appropriate measures, such as initiation of the nonconforming work process, taken.

The SAS will maintain a list of contract laboratories being used and their documentation of competency.

21.4 **USE OF OUTSIDE EXPERTS**

Under certain circumstances the CLD may employ an outside expert. These circumstances include, but are not limited to, consultation in the resolution of technical disagreements, technical reviews of casework, proficiency test preparation, audits, training or other consultations. Use of an outside expert may only be done with the approval of the CLD Commander.

To request the use of an outside expert, a written request must be sent to the CLD Commander to include:
- The scope of work to be performed by the expert
- A CV or resume outlining the expert’s qualifications
22 RESEARCH PROJECTS, PUBLICATIONS AND PRESENTATIONS

22.1 POLICY FOR RESEARCH PROJECTS

All research projects employing the use of laboratory resources will be reviewed and approved by the analyst’s Laboratory Manager and the Standards and Accountability Section prior to the initiation of the project. This includes research projects for the investigation of new methodology or technology, measurement uncertainty studies, or additional studies on currently used methods.

22.2 PROCEDURE FOR RESEARCH PROJECTS

Prior to beginning any research study, a research plan, including experimental design, will be prepared for review and approval by the discipline Technical Lead or DNA Technical Leader, and then submitted up the chain of command through the Lab Manager, to the management liaison, and to the Standards and Accountability Manager or designee for approval. Research projects drafted as part of new employee training plans do not need approval by the management liaison or Standards and Accountability Manager provided associated costs are reasonable and justified. The selection of the appropriate type of equipment, standards, controls, and reagents should be part of the plan as well as a budget estimate. As the research progresses, the plan will be updated as necessary. Effective communication among all personnel involved, including other analysts in the section, the discipline Technical Lead or DNA Technical Leader, the supervisors, the management liaison and the SAS Manager will be accomplished through verbal or written communications.

The research plan shall follow the same criteria as those listed for developmental or non-standard method validation.

22.3 POLICY FOR MANUSCRIPTS AND PRESENTATIONS

An individual shall obtain approval from their supervisor prior to working on a manuscript for publication or a presentation of original research. A technically reviewed draft of the manuscripts or presentation of original research must be submitted to the SAS Manager for approval prior to submission of the manuscript to a journal or prior to the presentation. This policy applies specifically to research where the WSP CLD is mentioned in manuscripts for publication or presentations of research, when the author is a representative of the WSP CLD, or when the research or preparation for the presentation occurred on duty time. Final approval shall come from the CLD Commander.

Other types of presentations shall be approved as follows, and do not apply to the Procedures for Manuscripts and Presentations below:

- Criminal justice personnel training (e.g. law enforcement, prosecutors): presenter’s supervisor and the applicable Forensic Scientist 4 and management liaison (copy the lab manager)
• Informational presentations to the public (e.g. schools, Rotary): presenter’s supervisor (copy the lab manager)
• Internal technical training: Functional Area’s Forensic Scientist 4, or for DNA related training, the DNA Technical Leader
• Internal informational (non-training) presentations or manuscripts do not need prior approval (e.g. functional area meeting briefs or those required as part of a training plan).

22.4 Procedure for Manuscripts and Presentations

Technical reviewer(s) do not need to be Division personnel, but may be anyone qualified by training and experience to give a proper review. The technical reviewer’s approval shall be documented on the final draft of the manuscript/research presentation.

The final technically reviewed draft of the manuscript/research presentation must be submitted to the SAS Manager for approval at least 14 days prior to the time the manuscript or presentation is sent to a journal or the analyst is making the research presentation. The analyst’s supervisor and laboratory manager shall be copied on the communication to the SAS Manager.

At a minimum, the review for approval by the SAS Manager, or designee, will consist of:
• Technical quality and completeness of data: Is there sufficient data of adequate quality to justify inferences?
• Accuracy of the conclusions: Does the data in the manuscript/presentation support the conclusions?
• Proofing of mathematics, spelling, grammar and punctuation;
• Feedback should not consist of the reviewer’s perception of the style.

The author must address the reviewer’s comments and any differences of opinion will be resolved by consensus.

The approval documentation for a manuscript or research presentation from the technical reviewer, SAS Manager and the CLD Commander is the responsibility of the author.

Approval documentation for slide presentations may be noted on the first slide of the presentation by adding a “New Comment” and entering the electronic signature and date of the approver. Approved slide presentations and handouts may be placed on the FLSB Portal (Crime Lab Division/CLD Documents/Approved Presentation Materials) for use by other CLD staff. Minor modifications (e.g. statistics updates, lab specific information) to approved presentation materials may be approved by the presenter’s supervisor.
23 LABORATORY FACILITIES AND SECURITY

23.1 Facilities

In order for the laboratory to efficiently carry out its goals and objectives, adequate and proper space should be allocated for each laboratory activity and function.

The laboratory will have space designated for the safekeeping of official laboratory records and reports as well as space for reference material, books, and other documents necessary for carrying out the functions of the laboratory.

23.2 Building Maintenance and Environmental Conditions

The laboratory shall ensure that the environmental conditions do not invalidate the results or adversely affect the required quality of any measurement.

The laboratory shall monitor, control and record environmental conditions as required by the relevant specifications, methods and procedures or where they adversely influence the quality of the results. Tests and calibrations shall be stopped when the environmental conditions jeopardize the results of the tests and/or calibrations. The specific environmental requirements are addressed in the functional area technical manuals.

Regularly scheduled housecleaning will take place in each laboratory. Each CLD employee will be responsible for the general cleanliness of their own work area. The laboratory managers will be responsible for the overall cleanliness of their facility.

23.3 Security

23.3.1 Division Policy

The laboratory will be designed and equipped to assure the safety of laboratory personnel, the preservation and integrity of physical evidence, and the protection of the laboratory’s assets and records (including computer data). The laboratory will have an alarm system as outlined under “Intrusion Alarms” below. Access to the operational areas of the laboratory will be controlled and limited.

23.3.2 Procedures

Each laboratory facility shall define their areas of accessibility and have guidelines that govern accessibility to those areas. Laboratories differ in design, consequently some areas may, out of necessity, be used for several purposes. The laboratory’s security measures must account for multiuse areas and develop procedures to ensure proper security and access. In general, guidelines should consider the following types of areas:

23.3.2.1 Public Area

An area such as a lobby, common hallway, conference room, or restroom which may be accessed by members of the public during business hours without escort.
23.3.2.2 Evidence Delivery Area
An area designated for members of law enforcement to submit or receive evidence during business hours.

23.3.2.3 Evidence Vault
An area specifically designated for the storage of evidence accessible only by authorized laboratory personnel and escorted departmental auditors and authorized escorted individuals.

23.3.2.4 Evidence Examination Area
An area designated for the examination of evidence which is accessible only to laboratory personnel and authorized escorted visitors.

23.4 Securing the Laboratory

The specific opening and closing procedures for each laboratory will be documented in writing by the Laboratory Manager. Laboratory entry doors and the outside perimeter of the laboratory will be kept secure at all times.

23.5 Keys, Proximity Cards, and Combinations (Access Control Devices)

Laboratory managers or their designees will issue to employees laboratory door and alarm keys or proximity cards, and combinations or codes, as appropriate. Key and proximity card logs will be maintained in accordance with departmental regulations by laboratory managers or their designees, and combinations will be changed as needed to ensure that only authorized individuals have laboratory access. Keys and proximity cards may not be duplicated or loaned, and combinations or codes may not be divulged to unauthorized personnel.

Entrance/exit points and internal areas requiring limited/controlled access will have a lock system. Keys (magnetic cards, etc.) to these locks will be distributed only to employees, or, on a restricted basis, to visitors as noted below (see section on Visitors below). The distribution will be accounted for by the Laboratory Manager or designee in a log. The log may be paper and/or electronic (see WSP Regulation Manual, Section 10.17.010 Access Control Devices (ACD) Accountability).

It is recognized that each laboratory presents a unique security challenge because of its physical location, size, building design, proximity to other state offices and complexity of the security system. At a minimum, the following security criteria will apply:

- The laboratory will be secured during the hours it is vacant by an intrusion alarm
- If a laboratory employee leaves employment in a laboratory equipped with a single laboratory wide alarm code, that alarm code will be changed immediately; if an individual access card had been issued, this card will be immediately deactivated
• If keys or proximity cards are lost, the Laboratory Manager will be immediately notified and appropriate actions to maintain security will be taken
• All evidence will be kept in the evidence vault or a secure storage area during the hours the laboratory is vacant
• Personal firearms are not permitted on WSP sites.

23.6 **INTRUSION ALARMS**

Each laboratory will have an intrusion alarm system that is monitored at all times. Laboratory managers will create written alarm and emergency response procedures. Such procedures will include:

- The name and location of the alarm monitoring company or agency
- Instructions to the monitor regarding the notification of laboratory, police, and fire personnel
- Procedures to follow in the event of a false alarm or alarm malfunction
- Relevant information regarding the availability of laboratory keys, combinations, or codes that might be used in emergencies (e.g. a key in a sealed envelope). The security of any such key, combination, or code must be verified by laboratory managers at least once per year.

Each Laboratory Manager, or designee, shall conduct a test of the alarm system annually to verify that the alarm system is working. This test can be conducted at any time in the calendar year, but must be at least six months after the prior test. Acceptable methods of performing a test of the alarm system include:

- Intentional test: the system is intentionally ‘tripped’ by passing a motion detector, etc.
- Accidental test: an employee accidentally ‘trips’ the system
- Vendor test: a security system vendor is hired to test the system

A log will record all code changes and testing. The log will have the following information:

- Date alarm tested/alarm code changed
- Indication if this is an alarm code change or alarm test
- Zone or area tested (i.e., entry door(s), motion detector, etc.)
- If this is a test, an indication of whether the system passed
- The person who tested/changed alarm

23.7 **FIRE ALARMS**

Each laboratory will have smoke and fire detection systems. Evacuation drills will occur at least once per year and will be documented by the laboratory facility safety officer.
23.8 VAULT ENTRY AND SECURE AREAS

Entry into the vault shall be limited to designated laboratory personnel only, except for any unusual situations which must be preauthorized by laboratory managers or their designees. All visitors to the laboratory evidence vault will sign an entry log and will be escorted at all times. Laboratory managers will be responsible for controlling keys and combinations to vault-locking mechanisms.

Evidence lockers for the short-term storage of evidence will be made available to each scientist. Access to keys for short-term storage lockers will be limited to the section scientists and the laboratory manager or their designee. If not in a short-term storage area, any evidence retained outside the laboratory vault must be in a secure area.

23.9 VISITORS

Employees of the Forensic Laboratory Services Bureau who visit another FLSB laboratory do not need to sign the visitors log but will have limited access and may need to be escorted in certain areas of the laboratory based upon local laboratory policy. FLSB identification will be displayed at all times.

Non-FLSB employees will be required to sign the visitor log, will have limited access, and may need to be escorted in certain areas of the laboratory, based upon local laboratory policy. Visitor badges will be assigned or law enforcement identification displayed while in the laboratory.

At no time will visitors be given access to evidence vault areas or case files without an escort. Laboratory Managers may limit or allow access to the laboratory in special situations that do not meet the examples listed above.

23.10 DNA ELIMINATION SAMPLE FROM VISITORS

Visitors to analytical areas of a laboratory (instrument technicians, building trades workers, etc.) should be made aware of the following:

- A buccal swab sample is required from visitors to laboratory analytical areas prior to entry. This procedure is to monitor potential DNA contamination which could impact the interpretation of DNA test results. Managers (or their designee) may make an exception to this policy where appropriate (e.g. a brief walk-through or tour groups).
- Unless previously provided, a buccal swab sample for DNA elimination purposes will be collected by a CLD staff member. The DNA sample shall be sent to the CODIS Laboratory for analysis.
- Visitors may request the removal of their DNA profile from the elimination sample database once six months has elapsed since their last laboratory visit. Removal may be requested by sending an email to confel@wsp.wa.gov, listing the visitor’s name and the date and location of their last crime lab visit.
• A DNA staff member may also require that a face mask, sleeves, and/or gloves be worn. The personal protective equipment will be provided by the crime laboratory.
• Conversation must be kept to a minimum in order to avoid possible DNA contamination due to talking, sneezing, etc.
• Laboratory space must be kept clean and instrumentation protected from debris. If it is anticipated that the necessary work will produce debris, CLD staff will assist the worker to ensure the lab area is protected.
• Workers shall avoid placing tools, notebooks, etc. on laboratory countertops or instruments unless pre-approved by CLD staff.
• DNA post-amplification room work must be completed at the end of any multi-room project. Tools brought into a DNA post-amplification room shall be cleaned prior to re-use in the crime laboratory.

23.11 Observation/Document Review by Outside Experts

Outside experts wishing to observe the examination of evidence or review documents may be allowed on the premises under certain conditions. These conditions are necessary to protect the integrity of the evidence and the work processes carried out by the laboratory.

Visits by outside experts for the purpose of observing evidence examination will be allowed only when there is a limited sample and the analytical procedure will likely consume the entire sample.

23.11.1 First-Time Visit in the WSP Crime Laboratory Division

Experts will be considered for entry onto the premises after they comply with the following, at least 5 working days, dates inclusive, prior to the requested visit:
• a completed “Request for Observation of Examination/Document Review” form is received;
• a completed “Background Qualifications on Outside Expert” form is received;
• a current Curriculum Vitae is received.

For any request to observe laboratory examination, a DNA profile from the outside expert is required for purposes of DNA elimination. At the time of the first visit, a buccal swab sample will be collected unless the Crime Laboratory Division already has a DNA profile from the individual. The sample will be typed to obtain an STR profile with, at minimum, the CODIS core loci and added to the elimination database.

23.11.2 Repeated Visits in the WSP Crime Laboratory Division

An expert who has been previously granted permission to observe the examination of evidence or review documents on site in a specific discipline is only required to fill out the “Request for Observation of Examination/Document Review” form. The completed form must be received at least 5 working days prior to a requested visit, dates inclusive. The
visit must be in the same discipline as previously granted permission, but can be at different laboratories within the crime laboratory division.

Outside experts may request the removal of their DNA profile from the elimination sample database once six months has elapsed since their last laboratory visit. Removal may be requested by sending an email to confel@wsp.wa.gov, listing the outside expert’s name and the date and location of their last crime lab visit.

The Laboratory Manager will assess each request and can deny the request based on information provided at the time the request is made or based on information provided from previous denials.

While making an authorized visit, outside experts are not allowed to use state-owned equipment to conduct their own independent testing, but may use state-owned equipment insofar as it is necessary to observe the testing being conducted or to view the data collected.

Outside experts may be permitted to photograph, or otherwise record only images of evidence items related to the case, but are not permitted to record images of laboratory personnel or evidence related to a different case. The time allowed for recording evidence images is limited to no more than 15 minutes per item. No videotaping will be permitted.

**23.11.3 Interviewing Employees**

Interviews of employees by media, defense attorneys, or others as deemed appropriate, are allowed only insofar as the employee agrees to be interviewed and the interview process does not have a deleterious effect on the laboratory’s efficiency and resources. Interviews will conform to the following standards:

- Interviews of employees will be prescheduled and conducted with minimum impact to employees’ work assignments;
- All interviews will be conducted in a courteous and professional manner;
- A maximum of two hours will be allowed for any interview. If additional time is needed, the employee may opt to extend the interview or a second interview may be scheduled;
- Employees have the authority to stop or pause an interview for a rest break, or if they become uncomfortable for any other reason;
- Employees may consult with their supervisor or laboratory manager at any time, and may opt to terminate an interview if appropriate. They may also opt to have the supervisor or manager present during the interview.

**23.12 Laptop Security**

Many personnel are issued laptops for the purpose of offering mobility to their work. Because these agency laptops contain sensitive information, it is incumbent upon the custodian of the laptop to exercise due diligence in protecting the device from theft, loss,
or damage. As with all agency computers, when stepping away from the laptop the custodian must, at a minimum, lock the keyboard or log off. If the laptop must be left in a vehicle, it should be locked in a trunk. If this is not possible, as in a station wagon, the laptop will be covered or concealed. Care should be used during travel (airline, train, etc.) that sensitive information cannot be inadvertently viewed by the general public. The same is true for employees who use their laptop at home and reside with people who are not crime lab employees. When staying in a hotel room or in similar situations, if the laptop is left in the room, ensure the laptop is concealed or otherwise not out in the open.

Should the laptop become lost or stolen, the employee will follow the procedure outlined in the WSP Regulation Manual (see section 18.00.060 – Loss or Damage of Department Property and Equipment).
24 HEALTH AND SAFETY

It is important that the CLD establishes and maintains a health and safety program that is
designed to safeguard employees from service-related injury and health problems. The
CLD health and safety program is documented in the CLD Safety Manual.

Ultimate responsibility for the health and safety program lies with the Division
Commander, who along with the Laboratory Managers must provide continuing support
and monitoring. The Division Commander and the Laboratory Managers draw upon the
CLD Safety Officer and CLD Safety Committee personnel for technical support and
assistance. The CLD Safety Committee will be developed along the procedures outlined in
the WSP Safety and Wellness Manual and the Collective Bargaining Agreement.

Laboratory Managers will ensure performance of mandatory safety drills specified in the
Safety Manual. All personnel are required to be aware of the plan and to follow
procedures as situations arise.

It is the responsibility of the Laboratory Manager to ensure conformance with the Safety
Manual. The Safety Committee is responsible for conducting and documenting annual
safety audits. Each Safety Officer serves as a member of the CLD Safety Committee,
chaired by the CLD Safety Officer, which will meet at least annually for the purposes of
reviewing and updating the Safety Manual, discussing division-wide safety issues, and
making recommendations to management for improving the division’s chemical hygiene
and safety goals. Minutes of the Safety Committee meetings will be posted on the FLSB
Portal for review.

Any laboratory staff member has the responsibility to notify the Safety Officer, Section
Supervisor and/or the Lab Manager of any practice felt to be unsafe.

The Laboratory Manager is responsible for advising the Quality Process Manager and the
CL Division Commander of unsafe practices and the corrective measures implemented.
Each laboratory shall have emergency evacuation plans developed and posted in general
areas in the laboratory. The Safety Officer should schedule, implement, and document
annual evacuation drills.

CLD employees will review the Safety Manual on an annual basis. This review will be
documented on the CLD Safety Orientation Checklist (CLD-SAF-15001), to be retained in
each laboratory.

Health and safety issues will be included in each employee’s Position Description Form
(PDF), and they will be evaluated on their performance and their conformance to safety
policy.
24.1 Safety Audit

The Safety Committee will audit each laboratory on a yearly basis, confirming adherence to the Safety Manual. The safety audit report will be retained with other audit reports.

24.2 Safety Training

Each laboratory will be responsible for maintaining documentation of safety training for its employees. (See CLD Safety Manual).
25 APPENDIX 1: ROOT CAUSE ANALYSIS GUIDELINES AND PROCEDURES

Root cause analysis (RCA) is used to define, evaluate and systematically analyze a problem to determine the underlying factor(s) or reason(s) for the problem in order to focus on prevention and continued improvement of the system or process. It is important to realize that a root cause analysis is an event review, not a performance evaluation, and the purpose is learning, not punishment. Accordingly, personnel and disciplinary issues should be handled through a separate process from RCA.

25.1 JUST CULTURE

The CLD embraces a method of root cause analysis that creates a “just culture”. A “Just culture”:

- Is a culture of learning
- Recognizes that competent professionals make mistakes, but holds individuals accountable for reckless behavior. Holding people accountable by punishing them for human error is not going to advance the culture of learning.
- Balances blame-free event reviews with the need for professionals to be personally accountable for adherence to reasonable standards of professional conduct
- Balances an open and honest reporting environment with a quality learning environment and culture
- Fosters learning that will embed knowledge (lessons-learned) that may help prevent similar problems from occurring in the future
- Fosters continuous improvement

Root cause analysis may be the most difficult part of establishing proper corrective actions following the reporting of a nonconformity. By becoming skillful at investigating and solving problems of nonconformity in their work, a laboratory will ultimately need to conduct fewer investigations. But if done inappropriately, a root cause analysis investigation may lead to the inadvertent blame of individuals instead of identifying where a work process has broken down. Such blame will be detrimental to encouraging participation in the root cause analysis process.

The purpose of an RCA is to find out what happened, why it happened, and determine what changes need to be made to mitigate the identified causes of the problem and reduce the likelihood of recurrence.

RCAs are conducted by the individual(s) assigned by the SAS as the investigator. RCAs may be performed by a team, Technical Lead, Supervisor, Lab Manager and/or other subject matter expert. The number of participants conducting the RCA may vary depending on the nature of the nonconformity.
25.2 Procedure

25.2.1 Step 1: Identifying the Problem

The event, or nonconformance, should be clearly defined and analyzed for its causal factors. This entails a detailed review of the event by the investigator. The analysis and review is conducted to identify problems – what went wrong, what is the problem? The investigation normally begins with the objective stating of the problem. The problem statement is a concise, complete, and accurate single sentence describing what the problem or nonconformity is. Examples include:

- the proficiency test was not passed;
- the case file is missing documentation;
- the wrong individual was identified;
- the sample was contaminated; and
- the results from a different case were reported.

Be sure to start with a problem and not the solution. It is tempting to assume we know what will fix the problem before we’ve thoroughly examined it. Assumptions are often wrong and may hinder complete analysis of the underlying causes.

The investigator should not define the problem as a need for something. The problem statement should objectively state what went wrong, not why, or how. A good problem statement will facilitate a more thorough examination of the problem.

One tactic in formulating a problem statement is to work backwards from the point of not meeting a known policy, procedure, or goal or objective of the organization. When a determination of what objectives were impacted is completed, the problems affecting the objectives may be more discernable and the problem statement more readily drafted.

Collect and organize the facts surrounding the event to understand what happened. It is often helpful to create a detailed timeline of events pertaining to and leading up to the nonconformance. The investigator should consider reviewing equipment logbooks, instrument data, case files, procedures and policies, previous occurrences and any trends.

The investigator should interview personnel involved. It is important to get the perspective of people personally involved in the event since people naturally see and interpret things differently. Bring all parties involved in the problem in early so it fosters the non-punitive and problem solving nature of RCA investigations. Keep it transparent, focused, simple, and engaging. It may be helpful to provide a brief review of the process before starting the interview or discussion.

25.2.2 Step 2: Identify Root Causes

In this step, the investigator determines why something went wrong. The contributing factors, situations, circumstances or conditions that led to or increased the likelihood of
the event are identified and analyzed. In this step, the investigator must be both focused and open-minded.

A thorough analysis of contributing factors leads to identification and understanding of the underlying process and system issues (root causes) of the event. Contributing factors are not necessarily the root causes. The investigator must examine the contributing factors to find the root causes. A timeline should be used whenever possible as the basis for identifying all contributing factors. When identifying contributing factors, be careful to avoid “hindsight bias.” Knowing the eventual outcome of a timeline can influence how the investigator views activities leading up to the event. The investigator should consider only those factors that were actually present and known to those involved at the time, not what was only realized after-the-fact.

The investigator must determine if they’ve truly identified a root cause, versus a contributing factor which would require more digging. Ask the following questions for each potential root cause identified:

- Would the event have occurred if this cause had not been present?
- Will the problem recur if this cause is corrected or eliminated?

If the answer is NO, then the investigator has identified a root cause. If the answer to any question is YES, then the investigator may not have identified a true root cause and needs to ask more “why” questions. Continue asking these questions until you get to root causes. There may be multiple root causes.

The investigator should not make judgments about whether an individual did the right thing. This judgment is to be made by the supervisor and manager responsible for evaluating the employee’s performance.

At least one of the RCA tools mentioned below must be used.

25.2.2.1 RCA Tools

There are various approaches to RCA and some may be more effective than others depending on the nonconformity. Brainstorming and creating a cause and effect diagram are two such tools to determine problem statements and root causes. Using a cause/effect diagram while brainstorming possible causes to a problem helps one to focus on the various possibilities. This “Cause Mapping” can be used as a visual technique for capturing the cause and effect relationships in order to lead one back to the root cause(s). First identify the effect (problem statement), then list all possible causes. Some useful categories of causes include:

- People (health, training/skills, time management, knowledge of policies and procedures, etc.)
- Materials and supplies (lack of correct/complete forms, lack of appropriate containers, improper packaging, etc.)
• Procedures/methods (incorrect order of steps, incorrect application of procedure, etc.)
• Environment (HVAC failure, freezer water pipe burst, etc.)
• Equipment/instruments (ran out of gas, CE shut down, etc.)
• Below is an example of a cause and effect “fishbone” diagram (I = Instrument):
Another common RCA tool is “5 Whys”. Starting with the problem statement, the investigator asks “Why (did this process fail)?” repeatedly until the root cause is identified. This questioning process is continued until all the root causes are found. The “5 Why’s” process can also be used as part of a cause and effect diagram as discussed above. It is common to find the same root cause for two or more contributing factors. For example:

**Problem statement:** Analysis of an evidence item was not completed by the deadline.

1. Why? The instrument failed to complete the run.
2. Why? The instrument ran out of carrier gas.
4. Why? More gas was not ordered.
5. Why? An employee forgot to order more gas.

Three basic types of root causes are:

1. **Physical causes** – Tangible, material items failed in some way (e.g., the GC stopped working).
2. **Human causes** – People did something wrong or did not do something that was needed. Human causes typically lead to physical causes (e.g. the GC ran out of carrier gas).
3. **Organizational causes** – A system, process, or policy that people use to make decisions or do work, is faulty (e.g. the employee did not receive instruction on how to order more carrier gas).

Each root cause must be addressed in the corrective action plan.

### 25.2.3 Step 3: Develop a Corrective Action Plan (CAP)

The RCA is shared with the individual assigned the CAP by SAS, which may or may not be the individual assigned the RCA. The CAP will include the result of the RCA, corrective/preventive actions, and a timeline to implement the plan and report results of the implementation. The investigator should make specific, prioritized recommendations for preventive actions that are intended to prevent occurrences of similar events. These recommendations will be made in writing and submitted to the individual assigned the corrective action plan if different from the investigator.

To create the CAP, prioritize the factors that contributed to the nonconformance, evaluating both their severity and the probability of recurrence. The CAP will describe corrective actions (including preventive) that respond to the prioritization and likelihood of repetition of the root causes. Choose actions that address each root cause. These actions will generally require creating a new procedure or making a change to a current process.

When developing corrective actions, consider questions such as:

- What safeguards are needed to prevent this root cause from happening again?
- What contributing factors might trigger this root cause to reoccur? How can we prevent this from happening?
• How could we change the way we do things to make sure that this root cause never happens?
• If an event like this happened again, how could we stop the accident trajectory (quickly catch and correct the problem) before its severity escalates?

Aim for corrective actions with a stronger or intermediate rating, based on the categories of actions below. Corrective actions that change the system and do not allow the errors to occur are the strongest.

Stronger Actions
• Change physical surroundings
• Testing of equipment before purchasing
• Engineering controls into system (forcing functions which force the user to complete an action)
• Simplify process and remove unnecessary steps
• Standardize equipment or process

Intermediate Actions
• Make software enhancements/modifications
• Eliminate or reduce distractions
• Create checklist or other cognitive aid
• Eliminate look alike and sound alike terms
• “Read back” to assure clear communication
• Enhance documentation/communication

Weaker Actions
• Double checks
• Warnings and labels
• New procedure/policy
• Training
• Additional study/analysis

If a particular action cannot be accomplished due to current constraints (e.g. lack of resources), the RCI or individual assigned the CAP should look for other ways of changing the process to prevent a similar event from occurring in the future. Doing nothing should not be an option.

When developing corrective action plans, clearly state what is to be done, by whom, and when. Satisfactory implementation of the corrective action plan will be monitored so it is important to have clearly defined plans with timelines.
25.2.4 Step 4: Evaluation

Corrective actions will be monitored through annual internal audits or as detailed in the CAP. Was the CAP properly implemented and effective? This evaluation is summarized in the Corrective Action Report.