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OSBI Mission

The mission of every OSBI member is to insure the safety and security of the citizens of Oklahoma.

OSBI Vision

The OSBI will continue to be the professional law enforcement agency for the State of Oklahoma. We provide specialized apprehension and crime detection services through teamwork, training, research, and implementation of innovative technologies. We recruit and retain the expertise required to meet changing responsibilities. We increase public awareness through proactive publicity and education.

Foreword

The Oklahoma State Bureau of Investigation (OSBI) and the Criminalistics Services Division (CSD) are dedicated to provide quality service and results. The OSBI CSD has adopted the standards set forth by the American Society of Crime Laboratory Directors/Laboratory Accreditation Board (ASCLD/LAB) Legacy and International accreditation programs. In order to facilitate accreditation by the International program, this manual has been organized using the same outline structure as the International Organization for Standardization and the International Electrotechnical Commission (ISO/IEC) 17025 standards. Much of the language in this manual is new. However, some language was extracted from the previous CSD Quality Manual.

Quality Policy

The OSBI CSD management is committed to providing quality and professional service to our customers. It is the objective of the OSBI CSD to provide service that meets or exceeds the customer's needs and to ensure that quality and analytical practices meet or exceed the standards required for accreditation. The management system documents are provided to CSD employees to communicate the procedures which must be followed to provide this level of quality and service. As indicated in Section 1 below, all CSD personnel are responsible for knowing and

implementing these policies. This manual and the laboratory practices will be reviewed regularly in order to attain compliance with ISO/IEC 17025 standards and to continually improve the effectiveness of the management system.

1. Scope

- 1.1** This manual sets forth the policies and procedures which govern the work performed by members of the OSBI CSD.
- 1.2** All members of the CSD are responsible for knowing and abiding by all management system policies and procedures.

2. References

The following standards guide the requirements set forth in this policy manual. If the reference listed does not include a date, the most recent revision of the referenced document applies.

ISO/IEC 17025:2005

ASCLD/LAB-*International* Supplemental Requirements – Testing (Effective 04/01/2011)

Quality Assurance Standards for Forensic DNA Testing Laboratories (Effective 07/01/2009)

Quality Assurance Standards for DNA Databasing Laboratories (Effective 07/01/2009)

The FBI Quality Assurance Standards Audit for DNA Databasing Laboratories (Effective 07/01/2009)

The FBI Quality Assurance Standards Audit for Forensic DNA Testing Laboratories (Effective 07/01/2009)

ASCLD/LAB Proficiency Review Program

ASCLD/LAB Guiding Principles located at

http://www.ascl-d-lab.org/about_us/guidingprinciples.html

3. Glossary

In addition to the terms and definitions listed below, any definition provided in one of the documents listed in Section 2 also applies.

ADMINISTRATIVE STAFF: The Criminalistics Administrators, CSD Director and Executive Secretary.

ADMINISTRATIVE SUPERVISION: The authority to monitor the day-to-day activities and perform traditional managerial duties of assigned units or laboratories.

CASE FILE: The file folder containing hard copy documentation relevant to a particular case or the electronic file contained within the BEAST that contains documentation relevant to a particular case.

CASE RECORD: The cumulative records which document the quality, technical, and analytical information relevant to a particular case.

COMPLAINT: The expressed dissatisfaction by a customer with the quality or timeliness of work products or services.

CONVENIENCE PACKAGE: Evidence which is properly sealed and marked for identification may be placed in unsealed containers such as boxes or bags for the purpose of grouping items of evidence or for the convenience of carrying the evidence without that container having to meet the “proper seal requirements,” as long as evidence security requirements are otherwise met. These containers should be marked as a “convenience package.”

CORRECTIVE ACTION: An action or actions implemented to correct circumstances which led to non-conforming work. Successful corrective actions should prevent a reoccurrence of the same type of non-conforming work. This is also referred to as “preventive measures taken” by QAS Standard 14.1.b.5.

CORRECTIVE ACTION PLAN: A plan to resolve a discrepancy identified in casework, database activities, or proficiency testing work which will correct the problem and prevent a future occurrence. (QAS based definition – Std. 14)

CRIMINALISTICS ADMINISTRATOR (CA): Individual who reports directly to the CSD Director and is responsible for supervising Criminalist Supervisors.

CRIMINALIST SUPERVISOR: Individual who reports to a Criminalistics Administrator and supervises criminalists.

CRITICAL REAGENT: A reagent that requires testing on established samples before use on evidentiary samples in order to prevent unnecessary loss of sample.

CUSTOMER (CLIENT): A recipient of the OSBI Criminalistics Services Division reports and/or services. A customer can also be within the Criminalistics or Investigative Services Division.

DERIVATIVE EVIDENCE: Any tangible material removed or derived from an evidence item already having an assigned item number. Examples are cuttings, debris collections, latent lifts, and retained stain samples. Derivative evidence or containers will be marked with appropriate case number, item or sub-item numbers, analyst's initials and date, and listed in the case file.

EVIDENCE: For the purposes of this directive, evidence shall mean all materials submitted for scientific analysis during the course of an official criminal investigation.

EVIDENCE DESTRUCTION FORM: A form used to document permission for the destruction of evidence.

EVIDENCE RELEASE FORM: A form used to document the return or release of evidence to the courts, OSBI employees, or submitting agencies.

EVIDENCE TAPE: Tamper proof tape used in sealing evidence containers.

FUNCTION VERIFICATION: A check to determine if a piece of equipment or instrumentation is working correctly within specified parameters.

MAJOR DEVIATION: A planned and approved modification to current policy or protocol which will apply for a set period of time or to a defined grouping of cases or samples.

MINOR DEVIATION: A planned and approved modification which will be applied to a single case, sample, or single batch of samples/cases.

NO ANALYSIS CASE: Evidence in cases submitted to the laboratory where charges have been dismissed or for some other reason no analysis is required can be returned to the submitting agency.

NON-CONFORMING WORK: Work that does not meet the standards set forth in policy, procedure, protocol, or does not meet the needs of the customer. This may occur due to protocol drift or due to a quality or technical problem with a reagent, supply, or instrument.

ORIGINAL REQUESTING AGENCY: The agency having jurisdiction in the case that made the request for services. Evidence will be returned after analysis to the original requesting agency unless specified otherwise in this policy.

PERFORMANCE CHECK: Actions taken to ensure analysis methods still perform as intended. Performance checks are similar to validations, but more limited in scope.

PHYSICAL EVIDENCE TECHNICIAN: Individual responsible for the reception, storage, documentation, and handling of the physical evidence submitted to an Oklahoma State Bureau of Investigation Laboratory.

PREVENTIVE ACTION: Actions taken to improve circumstances which could lead to non-conforming work. (ISO/IEC 17025:2005 based definition)

PROTOCOL DRIFT: Unintentional and/or unauthorized deviations from current protocol.

PROPER SEAL: An evidence container is “properly sealed” only if its contents cannot readily escape and only if opening the container would result in obvious damage/alteration to the container or its seal. Staples alone cannot provide a sealed condition on evidence packaging. It is acknowledged that not all evidence can be sealed inside a container. A proper seal would constitute tape sealing, heat-sealing, or lock sealing and initialing the seal. A date on the seal is also recommended.

Evidence such as weapons which will require only test firing or serial number restoration may be tagged with an identification tag and do not require a container.

QUALITY: Adhering to generally recognized standards of good laboratory practice.

QUALITY ASSURANCE (QA): Those processes necessary to provide confidence that the results from OSBI Criminalistics Services Division analysis and testing will satisfy given requirements for quality.

QUALITY ASSURANCE AUDIT: A systematic examination and review to determine whether quality processes and related results comply with the protocols, policies, and procedures, and whether these practices are suitable and effective in achieving the quality objectives.

QUALITY ASSURANCE PROGRAM: OSBI Criminalistics Services Division guidelines describing recognized quality assurance requirements for forensic laboratory analysis and reporting.

QUALITY CONTROL (QC): The day-to-day operational techniques and activities used by the laboratory to consistently provide accurate analytical results that fulfill the requirements for quality.

QUALITY IMPROVEMENT COMMITTEE (QIC): The Quality Improvement Committee is an ongoing committee for the purposes of reviewing and implementing ways to improve the

quality of laboratory services. This committee, which meets at least quarterly, is composed of all Regional and Unit Supervisors, discipline Technical Managers and Administrative Staff.

QUALITY MANAGER (QM): The Criminalistics Administrator assigned the responsibility of overseeing quality operations including proficiency testing, auditing, reviewing non-conforming work, etc.

QUALITY RECORDS: Records generated from quality assurance procedures. This includes, but is not limited to, proficiency tests, corrective and preventive actions, audits, training documentation, continuing education, and testimony review.

REFERENCE MATERIAL: A material for which values are certified by a technically valid procedure and accompanied by or traceable to a certificate or other documentation, which is issued by a certifying body. Examples include known drug standards and NIST Standard Reference Materials (SRM's) which can include known values for a variety of substances, including DNA profiles.

REFERENCE STANDARD: A traceable standard, generally having the highest metrological quality available, from which measurements are derived. An example would be NIST traceable weights.

REMEDATION: Steps taken to correct non-conforming work, such as issuing an amended report, re-testing, etc. This is also referred to as "corrective actions taken" by QAS Standard 14.1.b.4.

REQUESTING OFFICER: The individual, authorized by statute, requesting examination of the submitted evidence. Criminalists will not be listed as a requesting officer.

RFLE: Request For Laboratory Examination form.

SAMPLING: The practice of testing a portion of a substance and reporting a conclusion for the whole substance using a statistically based or reasonable assumption of homogeneity of the whole.

SAMPLE SELECTION: The practice of selecting one or more samples from an item for testing based on training and experience. Following analysis, results are reported clearly and unambiguously to indicate that the results reported apply to the sample, not the whole item.

SUBMITTING OFFICER: The person delivering evidence to an OSBI laboratory. Criminalists will be listed as the submitting officer when involved with the collection of evidence.

TECHNICAL MANAGER (TM): The individual assigned the responsibility and authority for the technical operations in a particular discipline.

TECHNICAL PROTOCOLS (PROCEDURES): Technical procedures are a key element in establishing and maintaining quality control within the laboratory. Written procedures will be prepared for those routine tests performed in the OSBI Laboratory. The procedures used may be those developed and adequately validated by an outside agency or laboratory or those developed and validated in-house.

TECHNICAL RECORDS: Documentation generated in the analysis of casework or database samples. This includes reports, examination documentation, quality control results, etc.

TEMPORARY EVIDENCE CLOSURE OR SEAL: Temporary evidence closure consists of a piece of tape across a box, a paper clip on a folded evidence envelope, or some other closure that would not normally constitute a proper seal on evidence. A temporary closure is acceptable when the analyst will be away from the work area for a short period of time or overnight as long as the evidence is secured in a locking drawer or controlled access evidence area.

TESTING: Testing refers to analysis conducted at the request of OSBI CSD customers. This may include casework analysis, database sample analysis, or other work mandated for the OSBI CSD. This does not apply to training, research, etc.

TRACEABILITY: The property of a result of a measurement whereby it can be related to appropriate standards, generally international or national standards, through an unbroken chain of comparisons.

VALID COMPLAINT: A complaint that has been verified and that warrants action.

VERIFICATION: Determining whether or not a stated complaint is well founded, and indisputable.

4. Management Requirements

4.1 Organization

4.1.1 Legal Responsibility

The OSBI CSD is a division of the Oklahoma State Bureau of Investigation, which has been granted legal authority by state statute. Refer to O.S. Title 74, Section 150.2. The CSD Director is appointed by the Director of the OSBI and has the responsibility and authority for all laboratory functions and personnel.

4.1.2 Operating Guidelines

The OSBI CSD will provide forensic science service which meets or exceeds the needs of customers. OSBI CSD service will also meet or exceed the applicable standards set forth by ASCLD/LAB.

4.1.3 Management System Scope

The policies and procedures set forth in the management system apply to work performed by CSD personnel in any temporary, mobile, or permanent facility.

4.1.4 Interrelation of the OSBI CSD

In addition to serving customers outside the OSBI, the CSD also provides services to the Investigative Division of the OSBI. An agency organizational chart is located on the OSBI Intranet at http://128.1.2.243:7001/hr_master/faces/orgchart.jspx?_adf.ctrl-state=1475ubhsll_14. The responsibilities of the agency director, deputy director, and investigative personnel are located in OSBI Policy 103.

4.1.4.1 The responsibilities and authority of the CSD Director are defined in [Quality Procedure \(QP\) 1](#).

4.1.4.1.1 The OSBI CSD Director shall have sufficient authority to make and enforce decisions.

4.1.5 Management Requirements

The OSBI CSD ensures the effectiveness of the management system through the following steps.

- a) Personnel are provided sufficient authority and resources to complete their duties, including implementing the management system and identifying, correcting, and minimizing deviations from policies, procedures, and protocols.
- b) Personnel are protected from influences which could adversely affect the quality of work performed. See OSBI Policy 105 and O.S. 257-20-1-9.
- c) Confidential case information, including electronically stored and distributed reports and documentation, is protected. Refer to Oklahoma Statute Title 74, Section 150.5.
- d) Activities which would bring question to the competence or integrity of the agency and its employees are prohibited. Refer to the OSBI Code of Ethics and OSBI Policy 105.
- e) Organizational structure, including relationships between management, technical, and support personnel is defined. Refer to the attached organizational chart ([OSBI CSD QMA 1](#)).
- f) The authority, responsibilities, and interrelations for any position which impacts the quality of work performed are specified. Refer to [OSBI CSD QMA 1](#) and [QP 1](#). Current job descriptions are located at http://www.ok.gov/opm/jfd/jfd_g-page.htm.
 - 1. No employee is accountable to more than one supervisor per function.
- g) Testing staff, including trainees, will be supervised by individuals who are familiar with the methods and procedures used. This may be accomplished through the Supervisor's own experience in the methods and procedures used by staff or through the Supervisor's coordination with Technical Managers and/or Criminalistics Administrators familiar with the methods used. Refer to [QP 19](#) for additional information on training.
- h) Each discipline has a technical manager who has the authority, responsibility, and resources required to ensure the appropriate quality of work. Refer to [QP 1](#) for additional information regarding responsibilities and authority.
- i) One CA will be appointed as the QM for the CSD. Refer to the [OSBI CSD QMA 1](#) and [QP 1](#). Current job descriptions are located at http://www.ok.gov/opm/jfd/jfd_g-page.htm.
- j) Key managerial personnel (as defined in section 4.1.8 below) are responsible for naming a designee and notifying employees during planned absences. If a designee is not named, or there is an unplanned absence, the individual's

supervisor will be responsible for appointing a designee and notifying employees. Deputies for key managerial personnel are responsible at a minimum, for the critical duties of the position which cannot be delayed until the individual returns.

k) Through routine unit and discipline meetings, all employees are informed of the importance of their activities and how those activities help ensure that the CSD meets the objectives of the management system.

4.1.6 Effectiveness of the Management System

Administrative staff meets regularly and during meetings discusses the effectiveness of the management system and reviews the communication processes used in the laboratory to ensure they are appropriate.

4.1.7 Safety Coordinator

The individual assigned as the CSD Safety Coordinator (refer to OSBI CSD QMA 1) has the responsibility and authority for implementing, updating, and ensuring compliance with the health and safety program.

4.1.8 Key and Top Management

Key management personnel includes the following positions:

- CSD Director
- Quality Manager
- Safety Coordinator
- LIMS Administrator
- FSC Building Manager
- Technical Managers
- All Supervisors and Administrators

Top management is the CSD Director.

4.2 Management System

4.2.1 Management System Documents

The OSBI CSD management system documents the policies and procedures to be followed in order to ensure the quality of laboratory services provided. The OSBI CSD management system consists of the quality policy manual, the quality procedure manual, and discipline quality and protocol manuals. The documents of the management system are available to all CSD employees on the OSBI Intranet, <http://osbinet/main/>. Refer to [QP 2](#) for distribution procedures.

4.2.2 Quality Manual

This document in its entirety is the quality manual for the OSBI CSD. The [Quality Policy Statement](#) is located following the Foreword.

4.2.2.1 The ASCLD/LAB Guiding Principles of Professional Responsibility for Crime Laboratories and Forensic Scientists have been included as a reference to this manual (See Section 2 – References).

4.2.2.2 The CSD Director ensures that these Guiding Principles are reviewed by all CSD personnel annually and maintains a record of that review, in accordance with [QP 18](#).

4.2.3 Management Commitment

Management is committed to developing, implementing, and continually improving the effectiveness of the management system. This is evident through management's involvement in quality procedures including audits, proficiency testing, management system review, etc.

4.2.4 Management Communication

Management communicates the importance of meeting customer, statutory, and regulatory requirements during regular meetings of the Quality Improvement Committee (QIC).

4.2.5 Supporting Procedures

Quality policies are included in the quality manual, which follows the same outline as the ISO/IEC 17025 standards. Procedures governing the implementation of these policies which apply to multiple disciplines are included in the Quality Procedures. Quality policies and technical procedures which apply to a single discipline are included in the discipline quality and protocol manuals. Discipline specific manuals should not contradict the CSD Quality Manual or Quality Procedures.

4.2.6 Roles and Responsibilities

The roles and responsibilities of technical management and the quality manager are provided in section 4.1.5 above and the referenced attachments.

4.2.7 Management System Integrity

CSD Management will preserve the integrity of the management system anytime changes to the system are planned and implemented.

4.3 Document Control

4.3.1 General

The OSBI CSD controls all documents included in the management system to ensure the documents are appropriate to the work conducted. The management system consists of internally and externally generated documents. Documents as referenced in this policy include policies, procedures, regulations, standards, software, manuals, etc. Refer to [QP 2](#) for document control procedures.

4.3.2 Approval and Issue

4.3.2.1 Any technical protocol or discipline quality manual documents will be reviewed by the technical manager or his/her designee. Technical protocols and discipline quality manuals will be approved by the technical manager and the Criminalistics Division Director or designee, in his/her absence.

Management system documents including quality policies and quality procedures will be reviewed by the Quality Manager and will be approved by the Quality Manager and the Criminalistics Division Director or designee, in his/her absence.

4.3.2.2 [QP 2](#) describes the steps taken to ensure that:

- a) The current authorized version of management system documents is available at all OSBI CSD facilities.
- b) Management system documents are periodically reviewed and revised as appropriate.
- c) Documents which are no longer valid are removed from use promptly.
- d) Retired documents that are retained for legal or knowledge preservation purposes are marked appropriately to prevent unintended use.

4.3.2.3 Each internally issued document will be identified with the information specified in [QP 2](#).

4.3.3 Document Changes

4.3.3.1 Changes to documents can be made in two ways. Documents are revised following [QP 2](#). In addition, changes to documents can be documented using a major deviation, as described in [QP 3](#). Both methods follow the same review and approval method.

4.3.3.2 Each internally issued document will include an attached document history page or section. Insertions or alterations made to the document with each revision will be noted in this section, whenever practical.

4.3.3.3 Documents will only be amended as indicated under section 4.3.3.1 above. Amendments may not be made by hand writing on documents.

4.3.3.4 [QP 2](#) details how changes are made and controlled for documents issued through the OSBI Intranet.

4.4 Review of Requests, Tenders, and Contracts

4.4.1 General

[QP 4](#) establishes the procedures that will be followed for the review of requests, tenders, and contracts. This procedure ensures that:

- a) The customer's requirements, which include the type of analysis or methods to be used, are well defined, documented, and understood.
- b) The OSBI CSD is capable of meeting the customer's needs.
- c) The appropriate test method is selected.

Any differences between the request, tender, or contract will be resolved before work commences. Each contract should be satisfactory to both the OSBI CSD and the customer.

4.4.2 Records

An electronic or hard copy of the RFLE will be maintained with the case file as a record of the request, review, and contract. In addition, any significant changes will be recorded in a conversation log, e-mail, or equivalent document. All records of changes to the contract will also be maintained with the case file (either electronic or hard copy).

4.4.3 Subcontracted Work

This review process shall apply to work performed by any OSBI laboratory, regardless of which laboratory received the evidence and performed the review. It shall also apply to analysis that is subcontracted to a non-OSBI laboratory.

4.4.4 Deviations

The customer will be informed of deviations from the contract. If an analyst determines that requested analysis is not appropriate or recommends alternate or additional analysis, the customer will be contacted prior to modifying the contract.

4.4.5 Amendments

Any amendment or modification to the contract after analysis begins will be reviewed in the same manner listed under [QP 4](#). The person making the amendment will notify the affected personnel.

4.5 Subcontracting of Tests

In order to provide the best service possible, OSBI laboratories may choose to transfer work to another OSBI laboratory or subcontract to an outside vendor. However, when a customer (investigating agency, prosecuting agency or defense attorney) requests evidence be sent out for further testing and specifies where to send the evidence it is not considered subcontracting.

4.5.1 Qualification of Subcontract Laboratory

All OSBI laboratories and any laboratory performing work for the OSBI must be accredited through the ASCLD/LAB Legacy or ASCLD/LAB International program.

4.5.2 Customer Notification

The OSBI CSD shall notify customers in writing when subcontracting work to an outside vendor. When appropriate, the OSBI CSD will also obtain approval from the customer, preferably in writing.

4.5.3 Review of Subcontracted Work

The OSBI CSD maintains responsibility for subcontracted work, unless the customer or a regulatory authority specifies which subcontractor will be used.

4.5.4 Records of Subcontractors

The QM will receive and maintain a copy of the accreditation certificate for any laboratory which performs analysis on behalf of the OSBI.

4.6 Purchasing Services and Supplies

4.6.1 General

Selection and purchasing of services and supplies will be made according to OSBI Policy 208. The purchase, receipt, and storage of reagents and consumable materials used for analysis will be conducted according to [QP 8](#).

4.6.2 Verification of Reagents, Supplies, and Consumable Materials

Any supply, reagent, or consumable item that will affect the quality of analysis will not be used until inspected and/or verified according to [QP 8](#).

4.6.3 Descriptions of Items Affecting Quality

Items that affect the quality of analysis will be identified on the Internal Purchase Request (IPR) with a description specific enough to ensure the appropriate quality of item is purchased. This description may be a product number, catalog number, a reference to a particular grade or purity, or other technical description. The description provided will be reviewed and approved with the IPR.

4.6.4 Evaluation of Suppliers

The technical manager of each discipline will determine which reagents, consumables, supplies, and services are critical and affect the quality of testing. The technical managers will also oversee the evaluation of suppliers and maintain a list of approved suppliers, as described in [QP 9](#).

4.7 Service to the Customer

4.7.1 Assisting the Customer

The OSBI CSD will cooperate with OSBI customers to ensure that service provided meets customers' needs. This includes clarifying requests for analysis and monitoring the laboratory's work performance. However, the OSBI CSD will ensure that cooperation with one customer does not compromise confidentiality of other customers. Refer to [QP 10](#) for procedures on customer assistance.

4.7.2 Soliciting Feedback from Customers

The OSBI CSD will seek feedback from customers, primarily through the use of surveys. Feedback will be utilized to improve the management system, analytical procedures, and customer service. [QP 11](#) details the procedure for soliciting general customer feedback. [QP 32](#) details the procedure for soliciting feedback specific to testimony provided.

4.8 Complaints

Complaints will be resolved and documented according to [QP 12](#).

4.8.1 Quality Complaints

[QP 12](#) will also be used to resolve and document complaints submitted by employees regarding quality aspects of the management system.

4.9 Control of Nonconforming Work

4.9.1 Policy

Any work that does not conform to the requirements set forth in this manual, the Quality Procedures, or in OSBI technical protocols shall be addressed according to [QP 13](#). By following the procedure detailed in [QP 13](#), OSBI CSD shall ensure that:

- a) Responsibilities and authorities for managing nonconforming work are specified and appropriate actions are defined and taken when nonconforming work is identified.
- b) The nonconforming work is evaluated to determine the significance.
- c) A decision regarding the acceptability of nonconforming work is made and correction is done immediately.
- d) The customer is notified and work is recalled when necessary.
- e) The responsibility for authorizing work to resume is defined.

4.9.2 Implementation of Corrective Action

If the evaluation of the nonconforming work indicates a significant possibility that the problem could recur, or there is an indication that lab operations do not comply with

OSBI policy and procedures, then corrective action procedures outlined in [QP 14](#) will be followed.

4.10 Improvement

The management system will be continually improved using information gained during audits, analysis of statistical data, corrective and preventive actions taken, management review, etc.

4.11 Corrective Action

4.11.1 General

When nonconforming work is identified, it will be addressed according to [QP 13](#). This procedure details the appropriate authorities for implementing corrective actions.

4.11.2 Cause Analysis

As indicated in [QP 14](#), the first step of corrective action will be to investigate the root cause of nonconforming work.

4.11.3 Selection of Corrective Action

After the completion of the root cause analysis, potential corrective actions will be evaluated. The goal of the corrective action is to correct the problem and prevent the problem from recurring. The corrective action plan will also be appropriate to the magnitude and risk of the problem. The corrective action plan most likely to succeed in these areas will be selected and implemented. Any changes necessary as a result of the corrective action investigation will be implemented and documented.

4.11.4 Monitoring Corrective Actions

For each corrective action plan, the results of the corrective action will be monitored to determine effectiveness.

4.11.5 Additional Audits

When the nonconforming work indicates that there is a failure to comply with ISO/IEC 17025 standards or CSD policies and procedures, an audit of the areas of activity in question will be conducted as soon as possible. In addition, an audit may be used following the implementation of a corrective action plan in order to assess the effectiveness of the corrective action.

4.12 Preventive Action

4.12.1 General

Needed improvements or potential sources of nonconformity will be identified and routed as indicated in [QP 15](#). Preventive action plans will be developed, implemented, and monitored for effectiveness in order to ensure that opportunities for improvement are exploited and nonconforming work is prevented.

4.12.2 Procedure

[QP 15](#) details how to initiate preventive actions and how to utilize controls or other measures to ensure the preventive action is effective.

4.13 Control of Records

4.13.1 General

4.13.1.1 [QP 16.1](#) describes the procedure for maintaining quality and technical records. Quality records and technical records are defined in the glossary.

4.13.1.2 Records will be legible and stored in a manner that they are readily retrievable and protected from damage and loss. Retention times for records are also reflected in [QP 16.1](#).

4.13.1.3 Records will be kept in secure locations and are confidential.

4.13.1.4 Procedures for records stored electronically are detailed in [QP 16.1](#).

4.13.2 Technical Records

4.13.2.1 The OSBI CSD will retain records of examination documentation and supporting documentation, such as quality assurance/quality control documentation, and copies of reports for the period of time defined in [QP 16.1](#). Each case record will contain enough information to identify factors affecting uncertainty of measurement, if possible and applicable, and to enable re-analysis to be conducted under conditions as close to the original as possible. The identity of the individuals who sampled evidence, conducted testing, and/or verified results will be reflected in the case record.

4.13.2.2 Observations, data, and calculations must be recorded at the time they are made and must be identifiable to the specific case involved.

4.13.2.2.1 Examination documentation must include at a minimum, the start and end dates of examination.

4.13.2.3 Mistakes in examination documentation will be crossed out with a single line, initialed, and the correct value added alongside. Erasing, obliterating, or otherwise making the original data illegible is not permitted. Similar measures must be taken with records stored electronically to avoid losing or altering original data.

4.13.2.3.1 Any change made to existing hard copy examination documentation will also be initialed by the person making the addition.

4.13.2.3.2 Examination documentation is considered complete when it is submitted for administrative and/or technical review. Any change made to completed examination documentation shall be tracked.

4.13.2.4 Documents maintained as part of the case record are identified in [QP 16.2](#).

4.13.2.5 Examination and supporting documentation must be sufficient for another examiner to determine what was done and to independently interpret the data.

4.13.2.5.1 Latent print documentation shall meet all requirements listed in Appendix C of the ASCLD/LAB Supplemental Requirements.

4.13.2.5.2 Operating parameters used during instrumental analysis shall be recorded in the examination documentation, protocol, or another suitable and appropriate location.

4.13.2.6 Each page of examination documentation will bear the case number and examiner's handwritten initials (or secure electronic equivalent of initials or signature).

4.13.2.7 If a technician or other individual prepares examination documentation which another analyst interprets, reports, or testifies to, the person who prepares the examination documentation must initial the page(s) he/she prepares.

4.13.2.8 All administrative documentation, received or generated by the OSBI CSD, must be labeled with the laboratory case number.

4.13.2.9 When multiple cases are analyzed simultaneously, the case number of each case must be appropriately recorded on the printout if the data is recorded on a single printout.

4.13.2.10 Examination documentation should be one-sided. Each side of any two-sided examination documentation will be treated as a separate page (initialed and case numbered).

4.13.2.11 Examination documentation will be permanent in nature.

4.13.2.12 Verifications of analytical findings, such as latent print or firearms identifications, will be conducted by qualified examiners. Verifications will be documented to include what was verified, whether the second examiner agreed, and when the verification was conducted.

4.13.2.13 The meaning of any abbreviations or symbols specific to the OSBI CSD will be documented either in the case record or in discipline quality manuals or protocols.

4.14 Internal Audits

4.14.1

The OSBI CSD shall conduct internal audits as described in [QP 17](#).

4.14.1.1 Internal audits will be conducted annually.

4.14.1.2 Documentation of internal audits will be retained as quality records according to [QP 16.1](#).

4.14.2

If audit findings identify nonconforming work or indicate that the effectiveness of operations or validity of test results may be questionable, then procedures outlined in [QP 13](#), if applicable, and [QP 14](#) will be promptly followed.

4.14.3

An audit report will be completed according to [QP 17](#).

4.14.4

Implementation and effectiveness of any corrective actions generated as a result of an internal audit will be verified and recorded according to [QP 14](#).

4.14.5

Each OSBI laboratory will submit an Annual Accreditation Audit Report to ASCLD/LAB according to the deadline in [QP 17](#).

4.15 Management Reviews

4.15.1

OSBI CSD management will conduct a review of the management system and casework activities, at least annually, to ensure their continued effectiveness and to introduce changes or improvements as needed. The procedure for management system reviews is detailed in [QP 18](#). Records of management system reviews will be retained as a quality record according to [QP 16.1](#). Management system reviews will include the following topics:

- a) suitability of policies and procedures
- b) reports from managerial and supervisory personnel
- c) outcome of recent internal audits
- d) corrective and preventive actions
- e) external audits
- f) proficiency test results
- g) changes in volume and type of analysis
- h) customer feedback
- i) complaints
- j) recommendations for improvement
- k) any other relevant factors

4.15.2

Findings from management reviews and the actions taken will be recorded according to [QP 18](#). CSD management will ensure that actions are carried out according to an appropriate timetable.

5. Technical Requirements

5.1 General

5.1.1

Several factors impact the reliability of analysis conducted by the OSBI CSD. These may include the following:

- human factors (5.2)
- environmental conditions (5.3)
- protocols and method validation (5.4)
- equipment/instrumentation (5.5)
- measurement traceability (5.6)
- sampling (5.7)
- evidence handling (5.8)

5.1.2

The OSBI CSD shall take account of the factors listed above when developing and validating procedures, training and qualifying personnel, and in the selection and calibration of instrumentation.

5.1.3

The reliability of reagents will be verified according to [QP 8](#).

5.1.3.1 Reagents prepared in-house will be labeled with the identity of the reagent and the lot number or date of preparation at a minimum. Records identifying who prepared the reagent and documenting the function verification will be maintained.

5.2 Personnel

5.2.1

OSBI CSD Management shall ensure the competence of any individual who performs analysis, operates instrumentation, evaluates results, or signs reports. Work conducted by trainees shall be properly supervised. The education, training, experience, and/or demonstrated skill of an employee shall be used to qualify the individual.

5.2.1.1 Each OSBI CSD discipline shall have a documented training program which will be used to train employees in the knowledge, skills, and abilities necessary to perform analysis. Requirements for discipline training manuals are outlined in [QP 19](#).

5.2.1.2 Where applicable, training programs shall also address courtroom testimony.

5.2.2

OSBI CSD Management has established the goals for education, training, and skills of employees. These goals and the procedure for identifying training and conducting training are outlined in each discipline training manual. [OP 19](#) details how the effectiveness of the training program will be evaluated.

5.2.3

OSBI CSD shall use personnel employed by or under contract to the OSBI. If contract or additional support personnel are used, OSBI CSD will ensure that these personnel are also appropriately supervised and competent for the work they perform. Their work shall also be in accordance with the OSBI CSD Management System.

5.2.4

Current job descriptions for managerial, technical, and key support personnel involved in analysis are located at http://www.ok.gov/opm/jfd/jfd_g-page.htm. More specific job descriptions may also be located in each discipline's quality manual.

5.2.5

OSBI CSD Management shall indicate which procedures and work each employee is authorized to perform by providing a written memo detailing the task(s) and the date the individual is authorized to perform the work. Records of these authorizations shall be maintained in an electronic or hard-copy training notebook for each individual.

5.2.6

OSBI CSD personnel shall meet the education and competency requirements detailed in the ASCLD/LAB Supplemental Requirements. Technicians will meet the educational requirements established in the applicable written job description. If there is not an applicable job description available through the Office of Personnel Management, the Supervisor will be responsible for developing a written job description for the technician position(s) in his/her unit.

5.2.7

The OSBI CSD provides access to current literature sources by ordering journals and by providing internet access and on-line subscriptions to employees.

5.3 Facilities

5.3.1

OSBI CSD shall provide laboratory facilities with proper energy sources, lighting, temperature, and other environmental conditions to ensure correct performance of tests and procedures. Employees should exercise caution when conducting sampling or analysis in a location other than a permanent facility, such as a crime scene, to ensure that environmental conditions do not negatively impact the integrity of evidence or results. Accommodations and environmental conditions which would impact results shall be documented in the technical protocols.

5.3.2

When specific environmental conditions are required by the technical procedure or could impact the quality of results, the OSBI CSD shall monitor, control, and record the appropriate environmental conditions. Testing shall be stopped if the environmental conditions would negatively impact test results.

5.3.3

Incompatible testing activities shall be separated by time or space in order to prevent cross-contamination.

5.3.4

Access to laboratories and evidence rooms will be limited to employees assigned to the laboratory unit, evidence technicians assigned to the physical facility, Criminalistics Administrative personnel, and other personnel as authorized by the Criminalistics Services Division Director on a limited or permanent basis.

5.3.4.1 Laboratory security procedures are located in [QP 20](#).

5.3.5

Good housekeeping shall be maintained in OSBI CSD facilities. If necessary, technical protocols will be prepared for cleaning/sterilization procedures.

5.3.6

OSBI CSD follows the health and safety program detailed in OSBI Policies 121.0 through 121.5.

5.4 Test Methods and Validation

5.4.1 General

OSBI CSD uses appropriate methods for all testing and evidence handling. Evidence handling procedures are included in [QP 6](#) and [QP 7](#). Technical procedures, estimations for uncertainty of measurement, and any statistical techniques for analysis of testing data are included or referenced in discipline specific quality manuals and/or protocols. Instructions on the operation of instrumentation, sample handling and preparation will also be included or referenced in the discipline specific quality manuals and/or protocols, if written instructions are necessary to ensure the quality of test results. Any deviations to these procedures occur only as outlined in [QP 3](#).

5.4.1.1 All analytical protocols shall be documented and issued according to [QP 2](#).

5.4.1.2 Appropriate controls and standards shall be specified in the analytical records and the results of controls and standards tested shall be documented in the case record.

5.4.2 Selection of Methods

The OSBI CSD shall use analysis methods which meet the needs of the customer and which are appropriate for the testing conducted.

5.4.2.1 The reliability of any new method will be internally validated and the results of the validation study documented prior to implementing the procedure for use in casework. The procedure for suggesting, conducting, documenting, and maintaining records of a validation study are outlined in [QP 21.2](#).

5.4.3 Laboratory-developed Methods

Validation of new methods developed by the OSBI CSD shall be planned and conducted by qualified personnel who have the necessary resources. Effective communication shall be maintained and the validation plan shall be updated as the method development proceeds.

5.4.4 Non-standard Methods

Only approved technical procedures will be used in the analysis of casework. If a non-standard method is necessary, the method shall be subject to the agreement of the customer. The agreement with the customer shall include a clear specification of the

customer's requirements. The method must be validated prior to use on evidence samples.

5.4.5 Validation of Methods

5.4.5.1 Validation of a method shall provide objective evidence that the method meets the particular requirements for a specific intended use.

5.4.5.2 All methods used by the OSBI CSD shall be validated to ensure that the methods are fit for the intended use. Documentation of validation studies shall record the results obtained, the procedure used, and a conclusion indicating whether the method is fit for the intended use.

5.4.5.3 In order for a method to be determined fit for an intended use, the range and accuracy of the values obtained from the method must be relevant to the customer's needs.

5.4.5.4 Before implementing a validated method new to the OSBI CSD, the reliability of the method will be demonstrated against any documented performance characteristics (such as sensitivity or specificity) of the method. Records of the performance check shall be retained.

5.4.6 Estimation of Uncertainty of Measurement

The procedure for estimating the uncertainty of measurement is located in [QP 22](#).

5.4.7 Control of Data

5.4.7.1 Calculations and data transfers shall be subject to appropriate checks in a systematic manner, such as the administrative and technical review process. If an additional check is required, it should be included in the appropriate discipline protocol(s).

5.4.7.2 When computers or automated equipment are used for the acquisition, processing, recording, reporting, storage, or retrieval of test data, OSBI CSD Management shall ensure that:

- a) Computer software is documented in sufficient detail and suitably validated.
- b) Procedures are used to protect the data; such procedures shall include, but are not limited to, integrity and confidentiality of data entry or collection, data storage, data transmission, and data processing.

- c) Computers and equipment are maintained to ensure proper functioning and are provided with environmental and operating conditions necessary to maintain the integrity of the data.

5.5 Equipment

5.5.1

OSBI CSD Management shall furnish each laboratory and/or unit with the equipment necessary to ensure the correct performance of analytical tests conducted. If equipment outside of the immediate control of the OSBI CSD is used for analysis of evidence samples, OSBI CSD Management shall ensure that the equipment meets the standards outlined in ISO/IEC 17025.

5.5.2

Equipment and software used to perform analysis, calibration, or sampling will comply with specifications relevant to the test and shall be adequate to achieve the required accuracy. Calibration programs/procedures will be established as outlined in [QP 24](#).

5.5.3

Instruments will be operated by authorized CSD personnel. Authorization for unsupervised use of instruments will be documented in the authorization to work memo addressed in section 5.2.5. OSBI CSD trainees and technicians, practicum students, and interns are authorized to use equipment/instruments in the unit(s) they are assigned to **under the supervision of authorized CSD personnel**. Current instructions for use and maintenance will be included in discipline protocols so that they are readily available for use by the appropriate CSD personnel. Alternately, manufacturers' manuals or use and maintenance instructions may be referenced in the protocol and placed in a designated location for easy access by authorized personnel.

5.5.4

Whenever possible, each instrument used for testing and its software significant to the test result will be uniquely identified. At a minimum, unique asset numbers will be assigned in accordance with OSBI Policy 209.

5.5.5

For each instrument and its software significant to the analysis performed, the following records will be maintained according to [QP 24](#):

- a) the identity of the instrument and software
- b) the manufacturer's name, model number, and serial number and/or asset number
- c) documentation of function verification
- d) the current location, if appropriate
- e) the instruction manual, if available, or a reference to the location of the manual
- f) copies of all calibration and adjustment reports/certificates, including the date, result of calibration/adjustment, acceptance criteria, and the due date for the next calibration
- g) the maintenance plan, if appropriate, and records of maintenance performed to date
- h) description of any damage, malfunction, modification, or repair

5.5.6

Procedures for safe handling, transport, storage, use, and maintenance of measuring equipment are located in [QP 24](#).

5.5.7

Instruments and equipment which have been mishandled or have been shown to be outside acceptable limits will be taken out of service. Out of service instruments and equipment will be clearly labeled as out of service until repairs are made and the instrument/equipment is placed back in service following a successful function verification/calibration. The impact of the defect or departure from acceptable limits will be evaluated and procedures outlined in [QP 13](#) will be initiated.

5.5.8

Whenever possible, equipment which requires calibration will be labeled to show the status of the calibration, including the date of the last calibration and when recalibration is due.

5.5.9

When equipment goes outside of the laboratory, whether for repair or another purpose, the function and calibration status will be checked and shown to be satisfactory before the equipment is returned to service.

5.5.10

If intermediate calibration checks are needed to maintain confidence in the calibration status of equipment, the checks will be done according to a written protocol approved by the appropriate Technical Manager.

5.5.11

Where calibrations result in correction factors that must be used, the discipline shall implement a procedure to ensure that any copies (e.g. in computer software) are correctly updated.

5.5.12

Applicable controls, defined in protocol, will be used to safeguard test and calibration equipment including hardware and software from adjustments which would invalidate the test and/or calibration results.

5.6 Measurement Traceability

5.6.1 General

Any equipment used for testing or calibration which has a significant impact on the accuracy or validity of the test, calibration, or sampling shall be calibrated before being placed in service. This includes any equipment used for subsidiary measurements such as environmental conditions, if it would have a significant impact on the validity or accuracy of results.

The procedures for calibration of equipment are outlined in [QP 24](#).

5.6.1.1 As specified in [QP 24](#), the procedures for checking the calibration of equipment are established based on the specific requirements of the tests being conducted. Under normal circumstances, a check of calibration will be conducted after any shut down and following service or other substantial maintenance. Calibration check intervals will not be less stringent than the manufacturer's recommendations.

5.6.2 Specific Requirements

5.6.2.1 Calibration The OSBI does not provide calibration services as defined by ISO/IEC 17025.

5.6.2.2 Testing

5.6.2.2.1 As indicated in [QP 23](#) and [QP 24](#), the calibration program for equipment is designed to ensure that calibrations and measurements are traceable to the International System of Units (SI), if possible. This is not required if the associated contribution of the calibration to the total uncertainty of the test results is negligible. In this situation, the OSBI CSD shall ensure that the equipment used provides the uncertainty of measurement needed.

5.6.2.2.2 Where traceability of measurements to SI is not possible and/or relevant, the OSBI CSD shall provide confidence in measurements by establishing traceability to appropriate standards such as certified reference materials, specified methods, and/or consensus standards.

5.6.3 Reference Standards and Materials

5.6.3.1 Reference Standards [QP 25](#) outlines the procedures for the calibration of reference standards. Reference standards will be calibrated by an organization capable of providing traceability to SI units as described in ISO/IEC 17025 standard 5.6.2.1. Reference standards will only be used for calibration unless it can be demonstrated that other use will not invalidate their performance as a reference standard. Reference standards will be calibrated before and after adjustments.

5.6.3.2 Reference Materials As specified in [QP 26](#), reference materials will be traceable to SI units of measurement, or to certified reference materials, whenever possible. Accuracy of internal reference materials will be checked as far as is technically and economically practical.

5.6.3.2.1 Reference collections of data or items encountered in casework that are maintained for identification, comparison, or interpretation purposes shall be fully documented, uniquely identified, and properly controlled.

5.6.3.3 Intermediate Checks When checks are needed to ensure confidence in the calibration status of reference, primary, transfer or working standards and reference materials, these checks will be carried out according to defined procedures and schedules.

5.6.3.4 Transport and Storage [QP 26](#) establishes the procedures for safe handling, transport, storage, and use of reference standards. These procedures prevent contamination and deterioration of the standards and protect their integrity.

5.7 Sampling

The OSBI CSD will not report results based on a statistical sampling method (see [glossary](#)). The OSBI CSD may report results for a whole based on testing a portion in limited circumstances which include toxicology analysis and the identification and quantitation of controlled substances. Discipline quality manuals and protocols will specify the necessary steps to ensure homogeneity of toxicology and drug quantitation samples and the amount of sample to be used for analysis. State statute establishes a legal basis for homogeneity for the identification of controlled substances and the amount of sample to be tested will be based on the analyst's training and experience.

5.8 Evidence Handling

5.8.1

The procedures for transportation, receipt, handling, protection, storage, retention and/or disposal of evidence items are included in [QP 5](#) through [QP 7](#). These procedures include all provisions necessary to protect the integrity of evidence and the interests of the OSBI CSD and our customers.

5.8.1.1 Through compliance with the evidence handling procedures outlined in [QP 5](#) through [QP 7](#), the OSBI CSD documents the chain of custody for evidence received and analyzed by the laboratory. The minimum components of a chain of custody record include the person (by signature or electronic equivalent) or location receiving evidence, the date of receipt or transfer, and a description or unique identifier of the evidence.

In order to ensure a complete and accurate chain of custody, all employees will document evidence transactions in the LIMS at the time evidence is physically moved from one location to another, unless exceptions are provided for in evidence handling procedures. In addition, employees shall not share LIMS passwords with anyone. Failure to comply with this policy will result in progressive discipline.

Failure to comply with evidence handling procedures may also result in progressive discipline.

5.8.1.1.1 As detailed in [QP 6.1](#), when evidence is subdivided in the laboratory, the OSBI CSD requires the same chain of custody documentation for any sub- items created.

5.8.1.1.2 As described in [QP 5](#), evidence accepted and stored by the OSBI CSD will be properly sealed.

5.8.2

The OSBI CSD utilizes the “BEAST” Laboratory Information Management System (LIMS) to identify evidence items while they are in OSBI CSD custody. This system, in conjunction with the evidence handling procedures, ensures that evidence cannot be confused physically or when referred to in the case record or other documentation. The system allows for sub-dividing groups of evidence items, transfer of evidence within the laboratory, and receipt and return of evidence.

5.8.3

When evidence is received, any abnormalities regarding the packaging or condition of evidence will be recorded. If there is doubt whether the item is suitable for testing or if the item does not match the description provided, the customer will be consulted for clarification and the conversation recorded using the “Narrative” button on the “Case Info” tab in the LIMS before proceeding.

5.8.4

[QP 6.1](#) details the procedures for preventing loss, deterioration, or damage to evidence items during storage and handling. This includes ensuring the security and proper environmental conditions of evidence storage locations.

5.8.4.1 All evidence will be stored in a secured, limited access storage area when not in the process of examination.

5.8.4.2 [QP 6.1](#) details how to secure unattended evidence in the process of examination.

5.8.4.2.1 [QP 6.1](#) also clearly defines when evidence is considered to be in the process of examination.

5.8.4.3 Each item of evidence shall be marked with the case number and item number. If it is not possible to mark the evidence or if marking the evidence with the item number could affect the integrity of the evidence, then the proximal container or tag shall be labeled.

5.8.4.4 When evidence, such as latent prints or impressions, can only be recorded or collected by photography and the image itself is not recoverable, the photograph or negative of the image shall be treated as evidence.

5.8.4.5 OSBI CSD personnel collecting evidence at a crime scene will ensure that the evidence is protected from loss, cross-transfer, contamination, and deleterious change, whether in a sealed or unsealed container, during transport to the laboratory. Crime scene evidence shall be appropriately identified, packaged, and entered into the LIMS as soon as practical.

5.8.4.6 The OSBI CSD maintains individual characteristic databases in the Latent Evidence Unit, CODIS Unit, and Firearms Unit. Procedures for the operation of individual characteristic databases are located in the appropriate discipline/unit quality manuals and/or protocols. These procedures include a designation of how database samples will be treated (as evidence, reference sample, etc.), how samples will be identified, how samples will be protected from loss, cross-transfer, contamination, and deleterious change, and how access to the databases will be restricted.

5.9 Assuring Quality of Test Results

5.9.1

The OSBI CSD procedures for monitoring the validity of tests are located in technical protocols as appropriate. In addition, procedures for proficiency tests, re-examination, and reviews are referenced below. Quality control data will be recorded in a way to allow trends to be detected and whenever practical, statistical techniques will be used to review the data. OSBI CSD quality control monitoring is planned and reviewed according to the procedures referenced. Monitoring includes the following:

- a) use of appropriate controls and standards, which are specified in protocols and recorded in case records
- b) regular use of certified or secondary reference materials, as appropriate
- c) internal and external proficiency testing
- d) re-analysis of casework

5.9.2

Quality control data will be analyzed and planned action will be taken to correct the problem if the quality control data is outside the predefined window for acceptability.

5.9.3

The OSBI CSD proficiency testing program is located in [QP 30](#).

5.9.4

Technical review of casework will be conducted according to [QP 31](#) in order to routinely verify that conclusions reported are accurate and supported by the examination documentation. [QP 31](#) further defines the scope of a technical review, how technical reviews will be documented, and what actions will take place if a discrepancy is noted.

5.9.5

Administrative review of casework will be conducted according to [QP 31](#) to ensure that reports and case records are accurate and complete. All OSBI CSD reports, with the exception of no analysis reports, will be administratively reviewed prior to release.

5.9.6

Testimony provided by OSBI CSD analysts will be monitored according to [QP 32](#).

5.10 Reporting Results

5.10.1 General

The results of all analyses and examinations conducted by OSBI CSD personnel will be reported accurately, clearly, unambiguously, and objectively in a Criminalistics Examination Report.

5.10.1.1 In the event that a request for analysis is cancelled, no-analysis or partial analytical reports will be issued according to [QP 28](#).

5.10.2 Test Reports

Analytical reports will be prepared and issued according to [QP 28](#).

5.10.3 Test Reports – Additional Requirements

5.10.3.1 OSBI CSD reports and/or case records will include the following information:

- a) Deviations from, additions to, or exclusions from the protocol and specific test conditions as necessary for interpretation of the test results shall be recorded in the case record.
- b) When relevant, a statement of compliance with requirements or specifications should be included in the case record.
- c) Where applicable, a statement on the estimated uncertainty of measurement should be included in the test report. Under most circumstances, records for uncertainty of measurement will be maintained by the laboratory and available on request. A statement should be included in the report when it is relevant to the validity of the test result, the customer requests the statement, or if the uncertainty affects compliance to a specification limit.
- d) Opinions and interpretations shall be included in the report when necessary. For example, expert opinions regarding comparison of latent prints or interpretations of DNA profiles.
- e) Additional information shall be included in the report and/or case record as required by the method or by the customer.

5.10.3.2 If a sampling plan is used to analyze evidence, the following information shall be included in the case record:

- a) the date of sampling
- b) identification of the item sampled
- c) location of sampling
- d) reference to the plan and procedures used
- e) details of any environmental conditions during sampling that may affect the test results
- f) any standard or other specification for the sampling method and any deviations, additions to, or exclusions from the specification

5.10.3.3 [QP 33](#) describes the procedure used for releasing case information.

5.10.3.4 Any OSBI CSD analyst who issues a report or testifies based on the examination documentation generated by another individual shall complete and document a review of all relevant pages of documentation in the case record.

5.10.3.5 When associations are made, the significance of the association shall be communicated clearly and qualified in the report.

5.10.3.6 If comparisons are performed and result in an elimination, the elimination shall be clearly communicated in the report.

5.10.3.7 If results are inconclusive, the reason why no definitive conclusion could be reached shall be documented in the report.

5.10.4 Calibration Certificates

The OSBI CSD does not issue calibration certificates.

5.10.5 Opinions and Interpretations

The OSBI CSD issues reports including opinions and interpretations only for forensic disciplines which have been appropriately validated and documents the training of each analyst issuing reports with opinions and interpretations. Opinions and interpretations shall be clearly identified in OSBI CSD reports.

5.10.6 Testing Results from Subcontractors

When analysis is subcontracted, the subcontractor shall provide a case record and report which meet the same requirements as OSBI reports and case records. The OSBI shall maintain a copy of the case record, and after reviewing the case record and report, the subcontractor's report will be sent to the customer.

5.10.7 Electronic Transmission of Results

Reports issued electronically must meet the same requirements stated above.

5.10.8 Format of Reports

OSBI CSD reports shall be formatted in a manner to accommodate the types of tests conducted and to minimize the possibility for misunderstanding or misuse.

5.10.9 Amendments to Reports

Modifications to OSBI CSD reports shall be handled according to [QP 28](#). Analysis conducted subsequent to the issuance of a report will be included in a separate, uniquely identified report. Corrections to an issued report will be made by issuing a corrected report and indicating which report it replaces.

I. Scope

This procedure explains the responsibilities and authority of key CSD personnel.

II. Procedure**A. Responsibilities of CSD Personnel****1. CSD Director**

The CSD Director will promote and direct the quality system and ensure that the policies and objectives are documented, as well as communicated to, understood by, and implemented by CSD personnel. The CSD Director serves as the laboratory director for the Forensic Science Center (FSC) and is an ex officio member of all CSD committees.

2. Criminalistics Administrators

Each Criminalistics Administrator (CA) will be assigned the administrative supervision of specific laboratories and/or laboratory units. Each CA will also be assigned additional responsibilities as indicated below.

- a) The CA responsible for the administrative supervision of the Forensic Science Center (FSC) Toxicology Unit, Firearms/Toolmarks Unit, FSC Biology Unit and Biology Technical Manager, Trace Evidence Unit, and the Eastern Regional Laboratory is responsible for the statewide coordination of these forensic disciplines and units/laboratories. This position will also be responsible for overseeing or coordinating CSD activities in the areas of , grants management, fleet management, laboratory surveys, complaints, and drug destruction. All activities will comply with quality standards set forth by the OSBI CSD.
- b) The CA assigned the administrative supervision of the FSC Drug Unit, FSC CODIS Unit, the Northeast Regional Laboratory at Tahlequah, and the FSC Physical Evidence Unit is responsible for the statewide coordination of these disciplines/units/laboratories. This position will also be responsible for coordinating CSD activities in the area of LIMS Administration, serving as a Discovery Assistant in accordance with OSBI Policy 226, and oversight of assigned regional laboratory facilities. All activities will comply with quality standards set forth by the OSBI CSD.

- c) The CA responsible for the administrative supervision of the Southwest Regional Laboratory at Lawton, FSC Latent Evidence Unit, and the Northwest Regional Laboratory at Enid is responsible for the statewide coordination of these disciplines/units/laboratories. This position will also serve as the Division Quality Manager and will be responsible for coordinating CSD activities in the area of quality control/quality assurance. This includes, but is not limited to, proficiency testing, laboratory accreditation, testimony monitoring, and audits. The CA assigned to this position will also be responsible for overseeing or responding to Open Record Requests.

3. Administrative Programs Officer – Evidence Discipline

The administrative programs officer over the evidence unit at FSC shall:

- a) Oversee the storage, maintenance, archival, and destruction of technical records.
- b) Oversee the destruction of evidence samples and prescription drug samples.
- c) Oversee and coordinate statewide activities of the physical evidence technicians/units.
- d) Assist with Laboratory Information Management System (LIMS) administration and generation of statistical reports.

4. Technical Managers

Each Criminalist Supervisor at the FSC, with the exception of the FBU and CODIS Supervisors, also serve as the Technical Manager for his/her discipline. The Biology Technical Manager will be identified on a Biology specific organizational chart. Each OSBI CSD Technical Manager shall:

- a) Assist with management reviews as described in [QP 18](#).
- b) Review and approve all technical procedures within the discipline.
- c) Implement and review quality documentation within the discipline.
- d) Stay abreast of recommendations made by Scientific Working Groups for the discipline and incorporate appropriate recommendations.
- e) Educate all discipline members in the implementation of the quality assurance program and confirm that all members of the discipline understand the importance of the program.
- f) Participate in audits and inspections when requested.

5. Criminalist Supervisors

Each OSBI CSD Criminalist Supervisor shall:

- a) Assist with management reviews as described in [QP 18](#) and disseminate information regularly to members of their unit.
- b) Ensure that members of the unit understand and follow all quality assurance procedures.
- c) Know and follow the CSD Quality Assurance Program.
- d) Make recommendations to improve quality within the discipline and division.
- e) Educate all unit members in the implementation of the quality assurance and safety programs and confirm that all members of the discipline understand the importance of the program.
- f) Serve as laboratory director, if assigned to a regional laboratory.
- g) Criminalist Supervisors are responsible for monitoring administrative review for their lab or functional unit. Supervisors are to be knowledgeable regarding the quality of casework produced by their staff.

6. Criminalists

Each Criminalist shall:

- a) Know, understand and apply quality procedures that pertain to their specific discipline.
- b) Ensure completeness of laboratory reports, notes and essential documentation and make recommendations and suggestions for improvements of procedures used for the examination of forensic evidence.
- c) Advise Technical Manager and/or Supervisor of any technical problems or questionable results and make recommendations for improvements.

7. Physical Evidence Technicians

Each physical evidence technician shall:

- a) Know, understand and apply all quality procedures that apply to proper evidence handling including evidence submission, transfer, return or destruction.
- b) Notify the Technical Manager and/or Supervisor of any concerns relating to the quality assurance program of the Division.

8. Laboratory Technicians

Each laboratory technician or part time employee shall:

- a) Know, understand and apply quality procedures that apply to their specific discipline or job task.
- b) Notify the Technical Manager and/or Supervisor of any concerns relating to the quality assurance program of the Division.

B. Authority of CSD Personnel

1. CSD Director

The CSD Director has the authority to make and enforce decisions impacting any and all work produced by the division.

2. Criminalistics Administrators

Under the administrative direction of the CSD Director, the Criminalistics Administrators have the following authority:

- a) The Quality Manager will have the express authority to immediately halt any laboratory activity that fails to exhibit the required levels of accuracy, specificity, reliability or validity with respect to the CSD Quality Assurance program.
- b) Technical decisions made by each Criminalistics Administrator responsible for the coordination of a forensic discipline will apply to all personnel engaged in any capacity within the affected forensic discipline. These decisions will be made after consultation with the Technical Manager for the discipline.
- c) The Safety Program Coordinator has the express authority to immediately halt any laboratory activity which is determined to fall outside established safety policies and procedures and applicable laws.

- d) Authority of each Criminalistics Administrator shall include but not be limited to the assignment of specific duties or responsibilities to specific personnel and the review of the activity of those personnel engaged in these duties including all quality practices adopted by the OSBI CSD.

3. Technical Managers

The technical manager of each discipline has the following authority:

- a) Technical managers will assign and approve forensic procedures. All procedures will address and include practices consistent with the quality standards.
- b) Technical managers have the authority to suspend any work which does not comply with the OSBI CSD Quality System or any applicable quality standards.

4. Criminalist Supervisors

Criminalist supervisors have the authority to suspend any work which does not comply with the OSBI CSD Quality System or any other applicable quality standards.

III. Attachments

None

I. Scope

All management system documents will be approved, issued, modified, and controlled according to this procedure. Management system documents include policies/procedures developed internally, externally prepared documents or standards which are referenced or used (user's manuals, applicable standards, etc.), and software (internally or externally developed) used for testing purposes.

II. Procedure**A. Control**

1. All approved, internally generated CSD management system documents (policies, procedures, protocols, forms, etc.) will be placed on the OSBI Intranet. Any hard copy or other electronic copy is considered an uncontrolled document.
2. Uncontrolled copies may be made, if necessary, to reference at a work area that doesn't have easy access to the OSBI Intranet. However, CSD employees creating or using uncontrolled copies must verify the uncontrolled copy is still current before each use and immediately dispose of any uncontrolled copy that is not current.
3. Uncontrolled copies may also be made for the purpose of responding to discovery requests/orders.
4. External documents, software, and any other management system documents which are not distributed through the OSBI Intranet will be referenced in the CSD or appropriate discipline quality manual, protocol, or an attachment to the appropriate document. The reference must identify the current revision/version approved for use and the distribution or location of the document.
5. The individual responsible for the initial approval of internally generated documents (Quality Manager (QM) or appropriate Technical Manager) should maintain a copy of the current version that can be edited when the document requires revision.

B. Approval

1. Technical protocols/procedures, discipline quality manuals, and related attachments and references will be approved by the appropriate Technical Manager and the CSD Director or designee.
2. The CSD Quality Manual, Quality Procedures, and related attachments and references will be approved by the QM and the CSD Director or designee.

3. CSD documents distributed through the OSBI Intranet will include the signature of the individuals who have approved the document..

C. Issue

Once approved, the document(s) shall be distributed to the designated point(s) of issue. When a document is replaced or rescinded, it shall be removed from the point(s) of issue at the time it is replaced or no longer effective.

1. Approved CSD documents distributed through the OSBI Intranet, with the exception of forms, will be scanned or otherwise converted to pdf format prior to placing the documents on the Intranet.
2. Documents may be added to and removed from the OSBI Intranet by the Quality Manager, appropriate Technical Manager, FSC Executive Secretary or designee.
3. Documents referenced by the CSD or discipline quality manuals will be added to or removed from the designated point(s) of issue by the QM, appropriate Technical Manager, or designee.
4. Externally generated management documents will be available at each location where related work is conducted. For example, any externally generated management documents referenced by analysts conducting drug analysis will be located at each regional laboratory providing drug analysis.

D. Notification

For internally generated management documents, an e-mail will be sent to the appropriate individuals indicating that the document has been issued, revised, or rescinded. The e-mail will be sent by the Quality Manager, appropriate Technical Manager, FSC Executive Secretary, or designee.

E. Archiving

Obsolete versions of management system documents should be retained indefinitely. All retained, obsolete documents must be moved to a specified archive location and/or marked as obsolete or out of date.

Whenever possible, archived documents will be maintained electronically in a secure OSBI network folder. Any hardcopy archived management system documents which are retained should be maintained, at a minimum, in the FSC administration area.

F. Identification

Internally generated CSD management system documents will be uniquely identified and include the following:

1. document number or designation
2. title
3. date of issue and/or revision number
4. page numbering
5. total number of pages or mark indicating the end of the document
6. issuing authority

Forms or other attachments to management system documents will be identified in the following manner:

1. unique form number or designation
2. date of issue and/or revision number
3. page number and total number of pages (e.g. page X of Y) or the designator “AΩ” may be used to indicate a one page form

Revision numbers for forms and attachments may be tracked independent of the document revision number. The current attachment revision number (if applicable), changes made to attachments, and approval of attachments will be included in the attachment, history, and approval section of the document it is attached to.

G. Review and Revision

- I. Management system documents will be reviewed annually.
- II. Internally generated CSD documents will include a history section or attachment which will be used to document the completion of revisions and, when possible, identify modifications made during revision. Management system documents which are reviewed and found not to need revision may be documented in the history section/attachment and/or in the management system review memo submitted according to [QP 18](#).
- III. Temporary deviations or modifications implemented between annual review/revision will be documented and issued according to [QP 3](#).

III. Attachments

None

I. Scope

This procedure explains the process to follow when a CSD employee believes that a deviation from a current CSD-authored, controlled document is necessary. This procedure does not apply to any policies or procedures issued from outside the CSD. Any deviations from CSD policy, procedure, or protocol which do not comply with this procedure are considered protocol drift and must be evaluated as potential non-conforming work according to [QP 13](#).

II. Procedure

A. Requirements and recommendations

In order to ensure the quality of analysis conducted and services provided, written policies, procedures, and protocols have been established and issued to all appropriate CSD personnel. However, due to the variability of evidence and circumstances encountered, many protocols, procedures, and policies are worded to include **recommendations** (indicated by “should”) instead of **requirements** (indicated by “shall”, “will”, or “must”). All CSD personnel are expected to follow both requirements and recommendations set forth in CSD policies, procedures, and protocols, with the following exceptions:

1. Planned deviations from **requirements** can be requested and conducted following approval of a minor or major deviation as indicated below.
2. CSD employees may deviate from **recommendations** stated in protocol, procedure, or policy, provided the employee can articulate a legitimate reason which warrants the deviation.
3. CSD employees should make a notation explaining the deviation from **recommended** procedure.
4. If an employee is not certain whether circumstances warrant a deviation from **recommended** procedure, he/she should consult the Supervisor for assistance.

B. Minor deviations

CSD employees will complete the following steps to request, approve, and document authorization to deviate from current policy, procedure, or protocol for an individual sample, case, or batch of samples/cases.

1. The employee will describe the proposed deviation to his or her immediate Supervisor or designee and obtain approval before implementing the deviation.
2. The Supervisor or designee will evaluate the benefits and risks of the proposed

deviation to determine if the circumstances warrant the deviation. The Supervisor or designee will consult with the discipline Technical Manager (TM) or CSD Quality Manager (QM), if necessary, to thoroughly evaluate the benefits and risks of the deviation.

3. If approved, the deviation will be documented in the case notes when applicable to case work and documented in a relevant location for non-case work activities.
4. The approval will be documented by initialing a short description of the deviation in the case notes or applicable materials in non-case work activities.
5. Alternately, a description of the requested minor deviation may be documented electronically by the analyst in the narrative section of the electronic case file. A Supervisor or designee may document his/her approval electronically by logging into the case and entering a narrative indicating his/her approval.

C. Major deviations

CSD employees will complete the following steps to request, approve, and document authorization to deviate from current policy, procedure, or protocol for a defined period of time or grouping of cases or samples. These steps will also be used to initiate, approve, and document permanent changes to policies, procedures, and protocols in between annual review and revision of controlled documents.

1. The requesting individual will complete Section I of the Deviation Request Form ([CSD QPA 3.1](#)) and specify on the form:
 - a) the applicable protocol, procedure, or policy
 - b) a description of the requested deviation
 - c) the specific instance(s) for which the deviation is requested
 - d) the reason for the deviation
2. The requesting individual will then forward the form for approval as indicated below.
3. Prior to implementation, all major deviations must be approved by the same authorities responsible for approving the document being modified.
 - a) Section II must be completed by the appropriate Technical Manager for any deviation impacting a discipline quality manual or protocol approved by the TM.
 - b) Section III must be completed by the Quality Manager for any deviation impacting the CSD Quality Manual, any Quality Procedure, attachment, or any other management system document initially approved by the QM.

- c) Section IV must be completed by the CSD Director or designee for all major deviation requests.
 - d) Any deviation request for a document which was also originally approved by another individual must be routed to that individual for evaluation and approval. This additional evaluation and approval should be documented on an attached memo or e-mail.
4. Individuals responsible for reviewing a deviation request will evaluate the request in the same fashion as the document being modified. This review includes an evaluation of the merit and risk of the deviation and whether the proposed modification complies with any and all applicable standards.
 5. Major deviations which are approved will be issued in the same fashion as the controlled documents affected by the deviation. The FSC administrative office will retain a copy of approved deviation requests.
 6. Deviation requests which are not approved will not be disseminated or retained. The individual denying the deviation request should notify the requestor of the decision.
 7. Approved deviations may be effective once signed by the CSD Director or designee. When necessary to delay the implementation of a deviation until employees have had an opportunity to be notified of the deviation, the CSD Director, QM, or TM may indicate an effective date on the bottom of the form. If the effective date is left blank, the deviation is effective on the date it was signed by the CSD Director.
 8. Major deviations should be routed and approved/disapproved by the appropriate persons within two weeks of the date of request.

III. Attachments

[OSBI CSD QPA 3.1, Rev 1 Deviation Request Form](#)

I. Scope

This procedure will be used to evaluate all requests for laboratory examination and provide response to the customer requesting analysis. Whenever OSBI CSD personnel accept evidence for analysis, the entry of the evidence into the OSBI CSD system constitutes a contract with the customer.

II. Procedure

A. Review of Requests

The following steps shall be taken to review all requests for analysis. This includes requests made at the time of evidence submittal and subsequent requests or amendments made after evidence has already been received by the OSBI CSD.

1. Verify that the individual requesting the analysis is authorized by statute to request services from the OSBI CSD. The agencies and individuals authorized to request services are listed in Title 74, Sections 150.2 and 150.5.
2. Verify that the type of evidence being submitted falls within the acceptance requirements described in [QMA 2](#).
3. Verify that the OSBI CSD is capable of providing the type and degree of service requested. A list of available services/methods is listed in [QMA 4](#).
4. If necessary, consult the appropriate Supervisor or Criminalistics Administrator to determine that the request meets the criteria listed above.

B. Tenders

Based on the results of the review conducted, one of three responses will be given to the customer.

1. If the request meets the criteria listed in II.A.1 through II.A.3, the analysis will be conducted.
2. If the request does not meet the criteria but can be modified to meet the criteria, the modification will be proposed to the customer. If the customer agrees, then the analysis agreed to in the modification will be conducted. For example, if a customer requested blood typing, DNA analysis could be proposed as an alternative.

3. If the request does not meet the criteria and cannot be modified to meet the criteria, the requested analysis will not be conducted. In this situation, the customer will be notified of the reason that the analysis cannot be conducted.
4. Under most circumstances, the customer will be notified of the inability to conduct analysis in the same fashion that the request was received. For example, if the request is made in person, the customer will be informed in person at the time the request is made.
5. The customer may be notified in a different fashion when the alternate method provides for better or more direct communication or when the customer cannot be reached by the same method that the request was received.

C. Contracts

When the OSBI CSD agrees to conduct analysis for a customer, a contract is established between the OSBI CSD and the customer. This contract is established in the following ways.

1. For requests received for evidence that has not yet been submitted, the contract is established by following the evidence intake procedure in [QP 5](#). The electronic submission record or a printed copy of the submission represents the contract established.
2. Requests received for evidence that has been submitted constitute an amendment to the original contract. Once an amendment has been agreed upon by the OSBI CSD and the customer, it will be documented in the narrative section located on the “Case Info” tab or by placing a copy of the communication (e-mail, letter, etc.) in the case file.
3. The OSBI has published a notice to customers ([OSBI CSD QMA 1.1](#)) regarding some deviations which may be made in the normal course of analysis. When necessary additional notifications may and/or should be made. A record of any notification made will be maintained in the case file.

III. Attachments

None

I. Scope

These procedures shall be used by any OSBI CSD employee when receiving evidence.

II. Procedure**A. Review of the Request**

Conduct a review of the request as described in [QP 4](#). A request for examination should be submitted on the Request for Laboratory Examination (RFLE) form ([OSBI CSD QPA 5.1](#)).

1. If the OSBI CSD does not have the capability to provide the service requested, return the evidence to the submitter. If possible, provide assistance to the customer in locating a laboratory that can provide the services needed.
2. If the OSBI CSD can provide the needed service, accept and log-in the evidence as described below.

B. Evaluation of Evidence Integrity Concerns

Inspect the evidence to ensure that it is packaged in a manner that will preserve the integrity of the evidence.

1. Evaluate each package to ensure that it is appropriate for the type of evidence it contains. For example, arson samples should be in arson cans while evidence with dried biological stains should be in packaging that will prevent mold or bacterial growth.
2. Evaluate the seals on each package to ensure that they protect the evidence from loss, cross-transfer, contamination, and deleterious change. Ensure that the officer's initials or other identifying mark are on each seal. Ask the officer to add his/her initials or identifying mark if initials are not present. Refer to [OMA 3](#) for more information regarding proper seals.
3. Determine whether there are any special storage conditions (e.g. store refrigerated) which need to be observed to protect the evidence.
4. Evaluate the items grouped together in each package and the types of analysis requested. If necessary, have the officer re-package items to ensure more efficient flow of evidence. For example, if a projectile which needs Firearms analysis is packaged with clothing which needs Biology analysis, have the officer repackage the projectile into a new container.

C. Evaluation and Identification of Safety Concerns

Inspect the evidence submitted to ensure that the evidence is packaged and labeled in a manner which ensures the safety of CSD personnel.

1. When firearms are submitted, have the officer indicate on his/her RFLE whether the weapon is unloaded. (Officers should be instructed to indicate unloaded only if they have direct knowledge that the weapon is unloaded.) If not, verify or have a qualified individual verify, that the firearm(s) has/have been unloaded or otherwise rendered safe. Document that the firearm is unloaded/rendered safe on the evidence package.
2. Ensure that chemicals, including any known carcinogens, mutagens, toxic substances, and volatile or foul smelling compounds are properly labeled and packaged according to safety policy and MSDS recommendations.
3. Ensure any sharp item (syringe, knife, glass, etc.) is packaged in a fashion that prevents the item from puncturing the package and potentially injuring CSD personnel.
4. Ensure that proper warning labels are on each package. This includes biohazard, sharps, and any other necessary hazard label.
5. Liquid evidence, with the exception of toxicology blood kits, must be double packaged in such a manner that the outer package would contain liquid in the event that the inner package was broken or leaked. For example, a properly sealed evidence package that contains liquid evidence could be placed into a bucket for storage and transport to prevent breaks and contain any liquid in the event of a break or leak. The bucket can be a convenience container and does not need to be properly sealed.

D. Data Entry

Enter information provided into the BEAST by creating a new case using manual creation or pre-log or by creating a new submission on an existing case. If it is not certain whether the submission is the first submission in a case, a search should be done with the information provided to ensure there will not be duplicate case numbers assigned to the same case.

1. In the lab case information field on the top half of the screen, enter the appropriate department case number or “NONE*” if an agency case number is not provided.
2. Seal numbers should be listed on the “Reference” line located in the middle, on the left side of the window/screen.

3. In the “Name ID” column on the “Names” tab, hit F4 to list any identifying numbers provided such as social security numbers (SSN), OSBI numbers, and Department of Corrections (DOC) numbers. Driver’s license numbers or any other numbers not in the F4 field may be listed by typing them directly into the Name ID field and adding an abbreviation to identify the type of number listed. For example, DL 1233425 would indicate driver’s license number 1233425.
4. For a rush request, select “1” for the case priority. For a routine/normal request, select “2” as the case priority.
5. For items received by mail, enter the shipping or mailing tracking number in the “Tracking Number” field by scanning the barcode or typing the number. If the tracking number is entered manually or if there is an agency barcode number/label on the mailing container, do not discard the container.
6. Enter each outer package of evidence on the “Containers” tab.
 - a) Enter a letter designation for each outer package in the “Cont. #” column.
 - b) Select a description of the evidence package that most closely describes the package from the drop down list in the “Package” column. If an appropriate description is not in the list, select “miscellaneous.”
 - c) Assign one item number to each container of evidence.
 - d) Select the analysis needed from the “Service Req(s) F4” column.
7. Enter item specific information on the “Items” tab.
 - a) In the “Pkg.” column, enter ITEM or the type of evidence collection kit (DUI, SAKIT, etc).
 - b) If the item is a blood alcohol kit, change the value in the “Qty.” column to 4.
 - c) In the “Type” column, select “EVIDENCE” or the appropriate evidence code from the drop down list. For evidence collection kits, such as blood alcohol, sexual assault, or gun-shot residue kits, select the appropriate kit type. The sexual assault kit type should reflect the agency that provided the kit (OCPD, OSBI, or TPD).
 - d) For blood alcohol kits, highlight the “Attribute F4” column, press the F4 key, and enter the blood kit and citation number in the window that opens.

- e) In the “Description F7” column, enter the description of what is in each package based on the information provided by the officer and/or the labeling on the package.

E. Finalize Case Creation

Finish creating the case by completing the following steps.

1. For all cases, once the information has been entered into the BEAST, click the “Quick Create” button.
2. For in person submissions, type the name of the individual submitting the evidence and capture his/her signature using the signature pad. Click on “Save Signature” when he/she has finished signing.
3. For in person submissions, offer the submitting officer a copy of the signed submittal form.

F. Scan Any Necessary Documents

Scan a copy of blood alcohol kits, officer RFLE’s, officer affidavits and any other necessary documentation by completing the following steps.

1. Insert the document into the scanner in the proper orientation, according to the scanner instructions.
2. Open the case within the BEAST.
3. With the “Case Info” tab selected, hit the F11 key or click on the “Documents F11” button.
4. Click on the “Scan_Doc” button in the lower right corner of the window that opens.
5. Select “black and white picture or text” and set the page size to “Letter” in the window that opens and click “Scan.”
6. When prompted for a document description, include the case number and “BA kit” , “affidavit”, or an appropriate description of the document scanned and then click ok.

G. Label Evidence and Folder

Ensure the evidence and case file(s) are properly identified by completing the following steps.

1. Apply the appropriate barcodes to the package(s) and file folder(s).

2. If the county and agency are not displayed on the barcode, label the evidence package(s) with this information.
3. If the evidence packaging does not have the department/agency case number or suspect and victim information, ask the officer to add the appropriate information if his/her agency will need it to identify the evidence when it is returned.

H. Transfer Evidence to Proper Storage Location

Transfer evidence to the appropriate vault by performing the following steps.

1. Scan the barcode representing the location to which the evidence will be transferred.
2. Scan the barcode for each evidence package being transferred.
3. Either scan the barcode representing “Process Transaction” or select enter.

III. Attachments

[OSBI CSD QPA 5.1, Rev. 0](#)

I. Scope

This procedure outlines the process for handling evidence within the OSBI CSD. Evidence handling procedures include uniquely identifying evidence, labeling evidence items and/or packaging, and preventing contamination.

II. Procedure**A. EVIDENCE INVENTORY**

1. An initial inventory of evidence containers (packages) is created during the evidence intake process. Refer to [QP 5](#).
2. During the examination of evidence, analysts will create an inventory of evidence items contained within any package or container of evidence that is opened for analysis.
3. The inventory of evidence should include a unique item number or sub-item number for each evidence item or package (if package not opened) observed by the analyst and a description of the item or package.
4. Analysts creating an inventory of evidence which may be forwarded for latent evidence analysis should be cautious when assigning sub-item numbers, since latent evidence analysis may often be conducted on the packaging of an item tested by another discipline (e.g. the packaging is an item for analysis). Whenever possible, the analyst creating the inventory should communicate with a latent evidence analyst to ensure that the inventory is created in a manner which facilitates analysis and reporting by both disciplines.

B. ITEMIZING EVIDENCE

The following procedure will be used to assign a unique item number or sub-item number to each piece of evidence analyzed.

1. Item numbers will be assigned using the default item numbering system currently in effect in the “BEAST” Laboratory Information Management System (LIMS). Each “parent” container or package is assigned an item number at the time of intake.
2. Parent item numbers will not be added, edited, deleted, or otherwise modified without prior approval of the Administrative Programs Officer responsible for the FSC Evidence Unit, or designee.
3. Sub-item numbers will also be assigned using the default numbering system in effect in the BEAST. Sub-item numbers may be edited, but only under the

following circumstances.

- a. Sub-item numbers may be edited in order to match a sub-item number previously assigned for evidence that has been re-submitted.
 - b. In addition, sub-item numbers may be edited if necessary to maintain consistency with a legacy numbering system. For example, if a case was analyzed prior to the BEAST and additional evidence is submitted, the sub-item numbers can be changed to the item numbers that would have been assigned under the original numbering system.
4. Analysts will use the following method to create, on the “Items” tab, an inventory of evidence items and/or packages contained within each container of evidence that is opened for analysis. Exceptions to this method are covered in section II.B.5.
- a. Upon opening each container of evidence, the analyst will determine how many items or packages are present in the container. Analysts may exercise some discretion to determine what is an item. Under some circumstances multiple pieces of evidence may be considered one item, provided that the remaining applicable requirements of this procedure are followed.
 - b. If only one item is present in the package, analysis should be completed and documented using the parent item number assigned to the package. However, if the item has been previously itemized or was submitted in relation to a case that was previously analyzed using a different numbering method, a sub-item should be created in order to maintain consistent numbering. See II.B.3.
 - c. If more than one item is present in the package, the analyst will use the sample button to create the correct number of sub-items. Alternately, the parent package can be sampled once, and the sub-item can be “duped” to create the correct number of sub-items.
5. Sub-item numbers will also be assigned to derivative evidence as necessary to accurately document the chain of custody and report results.
- a. Sub-items of evidence which are generated as a work product are not required to be created on the “Items” tab, if they will not be retained or transferred for further analysis. Examples may include sperm cell search slides and DNA cuttings, extracts, and dilutions.
 - b. An alternate numbering system may be used to distinguish between multiple stains, swabs, etc., during the testing process. Sub-item numbers will be

assigned to all derivative evidence that will be retained/transferred for further analysis.

- c. Evidence has been sub-divided anytime a portion of an evidence sample is removed and is not consumed in testing or returned to the original package. When evidence is sub-divided, a sub-item number must be assigned and created on the items tab so that chain of custody can be accurately tracked in the same fashion as the parent item.
- d. All assigned sub-item numbers will be recorded/created on the items tab.

C. LABELING EVIDENCE

1. The first Criminalist examining a piece of evidence will mark the item with the laboratory case number, item number, date, and his/her handwritten initials. Any Criminalist who subsequently examines the evidence will mark the item with the date and his/her handwritten initials.
2. If the evidence itself cannot be labeled or labeling the item itself could compromise the integrity of the evidence, the proximal container will be labeled.
3. All evidence items and/or their proximal containers will be legibly marked in such a way that the examiner's identifying marks or entry into the container does not cover, obliterate, or substantially alter another examiner's or officer's seal or markings whenever possible. In this way a traceable chain of seals is maintained.
4. When evidence is re-sealed after analysis, the examining Criminalist will mark the outer container with the item/sub-item number(s) contained within the evidence package. The Criminalist should also confirm that all outer evidence containers bear the case number, examiner's initials, container designator, and bar codes. Additionally, outer evidence containers should bear the county of offense and the submitting agency.

D. DIGITAL EVIDENCE

1. When evidence, such as latent prints and impressions, can only be recorded or collected by photography (including digital images) and the evidence in the image is not recoverable, the photograph or negative of the image must be treated as evidence.
2. Photos such as these, that are treated as evidence, must be handled in the following manner:
 - a) The photographs must be labeled with, or contain in the image, the case

number, date taken, analyst initials, and a unique photo number (i.e. Photo 1) or an item/sub-item evidence number. Whenever possible, this information should be included in the image.

- b) The photographs must be listed in the case file, in either hard copy or electronic format.
- c) In latent evidence cases, photographs and negatives can be retained in the case file.

E. EVIDENCE HANDLING

Each analyst is responsible for ensuring that all evidence examined is protected from loss, contamination, cross transfer, and deleterious change.

1. Evidence will be handled taking precautions to prevent any unauthorized alteration, any cross-contamination, or any deleterious changes by the following method or combination of methods:
 - a) Generally, open and examine only one container of evidence at a time.
 - b) Suspect and victim evidence will be searched in separate areas or at different times after decontamination measures are employed to prevent cross-contamination.
 - c) Every reasonable attempt will be made to maintain and preserve informative samples of biological evidence in serology related cases. Analysis of consumption samples will be documented according to [QP 16.2](#). Requests for additional documentation or observation will be handled according to [QP 10](#).
 - d) Representative samples of all informative biological material will be preserved in a manner to minimize degradation of the material and allow for future testing as required. These items may be retained, if necessary, or returned to the submitting agency.
2. All evidence received of insufficient quantity to allow a representative sample after testing should be photographed and documented according to CSD [QP 16.2](#) prior to examination. In addition, any work product of the analysis, such as DNA extracts, that may permit retesting, will be preserved and retained in such a way as to prevent degradation.

F. EVIDENCE STORAGE

1. Temporary closure of evidence is encouraged for evidence in overnight lockup to

prevent the possibility of loss, cross transfer, contamination, or deleterious change. Locking cabinets, drawers, etc. will be provided and used by Criminalists for securing evidence overnight or when the Criminalist will be away from the laboratory. During the process of examining evidence, if an examiner needs to leave for a short period of time, such as for lunch, it is not necessary to pack up the evidence being examined if it is in a secure area (e.g., a limited-access laboratory room).

2. Large items or boxes of evidence in the process of being examined will not be required for lock-up as long as they are closed and/or sealed in a secured restricted access lab area when the Criminalist will be away for short periods of time.
3. Evidence such as fingerprints and/or projectiles in unsolved cases that are subject to frequent requests for comparison may be treated as “evidence in the process of examination.” “Evidence in the process of examination” may be stored unsealed in a secure, limited access area, as long as the evidence is protected from loss, cross-transfer, contamination and/or deleterious change. After 30 consecutive days of no analysis or new requests for comparisons, a case is no longer considered “in the process of examination.” Cases no longer in the process of examination should be closed and the evidence sealed properly until analysis resumes or a new service request is received.
4. Evidence will be stored in conditions which prevent degradation or other deleterious change. Blood and urine samples submitted for Toxicology analysis will be stored refrigerated upon receipt and until they are disposed of according to statute or returned to the appropriate agency.
5. DNA extracts, including those in the process of examination, should be stored refrigerated or frozen.

G. TRANSPORTING EVIDENCE

1. Evidence collected from a crime scene must be protected from loss, cross transfer, contamination and/or deleterious change, whether in a sealed or unsealed container, during transportation to an evidence facility. Where appropriate, further processing to preserve, evaluate, document, or render evidence safe shall be accomplished prior to final packaging. Evidence collected from a crime scene must be appropriately identified, packaged, and entered into the secured electronic evidence tracking system as soon as practical.
2. Evidence that has been received into the custody of the OSBI CSD that must be transported to another facility should be sealed prior to transport.

3. Evidence being transported to another facility should not be left in a unoccupied vehicle overnight.
4. Transportation of evidence will be documented using the evidence transaction procedures outlined in [QP7](#).

III. Attachments

None

I. Scope

When evidence is needed for purposes other than casework analysis, this procedure shall be followed to ensure that the non-casework use of evidence is documented and communicated to OSBI customers.

II. Procedure

A. Approved Non-Casework Use of Evidence

There are several different circumstances other than casework analysis when the use of evidentiary samples is essential to further the mission and goals of the OSBI CSD. These include:

1. Research or validations which improve the quality or types of services the OSBI CSD can provide.
2. Training of CSD employees.
3. Quality control purposes, such as re-analysis casework.

B. Preferred Sources of Evidence for Non-Casework Use

In order to ensure that non-casework use of evidence does not conflict with the OSBI CSD's responsibility to preserve and protect the integrity of evidence, the following sources of evidence will be used in the order listed as appropriate. With the exception of evidence returned for destruction, **at least half of all samples must be retained or returned to the submitting agency.**

1. Evidence which has been resubmitted to the OSBI CSD for destruction or which is eligible for destruction based on state statute (e.g. toxicology samples).
2. Evidence from a no-analysis case, where there is no possibility for a later request for analysis. For example, evidence from a no-analysis case where the suspect has pled guilty.
3. Evidence from adjudicated cases or no-analysis cases with a potential that a request for analysis will be received later.
4. Evidence from active, non-adjudicated cases.

C. Notifying Customers of Non-Casework Use of Evidence

Notification will be provided to customers regarding non-casework use of evidence and whenever practical, an attempt will be made to obtain written authorization for non-casework use of evidence.

1. Notifications may be handled in the following manner:
 - a) All customers will be notified of the OSBI's policy regarding non-casework use of evidence by the posting of this policy and the attachment ([OSBI CSD QPA 6.2.1](#)) on the OSBI website.
 - b) For evidence samples retrieved from destruction evidence, no further notification is required.
 - c) For evidence from adjudicated cases, or no-analysis cases with no possibility of future requests, a second notification will be sent to the investigating agency or the prosecuting agency, at a minimum. This notification may be through a letter, memo, or e-mail. Alternately, this notification may be done by adding a statement to the Criminalistics Examination Report which states that a portion of the evidence is being retained in accordance with OSBI CSD QP 6.2 and the notification posted on the OSBI website.
 - d) For evidence from non-adjudicated cases, the second notification must be sent to the prosecuting agency and defense counsel. Including a statement in the report like that described in II.C.1.c above will be considered notification of both prosecution and defense.
2. As indicated in [OSBI CSD QP 6.2.1](#), authorization will be documented through the submittal of evidence for analysis or destruction.

D. Documenting Non-Casework Use of Evidence

When evidence is used for non-casework purposes it will be documented as indicated below.

1. Re-analysis of casework samples will be documented according to [QP 30](#) and all applicable discipline protocols.
2. Use of destruction evidence will be documented on the destruction form.
3. Use of no-analysis, adjudicated, or non-adjudicated casework will be documented in the case record.

- a. The information may be recorded in a case narrative or as part of the examination documentation, whichever is most appropriate.
- b. The amount and item/sub-item numbers of samples taken and the purpose of sampling will be documented. The amount of sample taken may be recorded by a specific size, weight or other measurement. Alternately, if additional samples or aliquots are taken which are the same size as that used in casework analysis, the documentation could simply reflect “1 additional test sample taken for training purposes” or a similar notation.

III. Attachments

[OSBI CSD QPA 6.2.1, Rev. 0](#)

I. Scope

The security and integrity of all evidence in the possession of the OSBI CSD will be preserved. Transfer, return and/or destruction of evidence will be conducted according to the specifications of this procedure.

II. Procedure**A. Documenting Chain of Custody for Evidence Transfers**

The following process will be used to document evidence transfers at the time of the transfer.

Scan the appropriate barcodes as indicated below.

- a) Location (the person receiving the evidence or the vault/storage area where the evidence will be placed)
- b) Item (the barcodes associated with all items being moved)
- c) Process chain of custody (or hit enter/click save)

B. Evidence Transfers

The following method will be used to transfer evidence between individuals or locations. For the purposes of this section, an evidence transfer refers to evidence that has been analyzed in one unit or laboratory and is being routed to another unit or laboratory for additional analysis. Evidence which has been received (but not analyzed) by one laboratory which must be transported to another laboratory for analysis does not constitute an evidence transfer as described in this section. Evidence transports must be documented according to section II.A above.

1. Routing Evidence

- a) The individual initiating an evidence transfer will verify that an assignment for the target unit exists in the BEAST. If there is no assignment for the target unit, the individual initiating the transfer will create the assignment.
- b) Evidence transfers can be done directly from analyst to analyst. This should be done whenever necessary to ensure the efficient and timely analysis of evidence.

2. Prepare Evidence Packaging

- a) Create a new container, if necessary, for the item(s) to be transferred.
 - i. From the items tab, click on the barcode button located next to the “Cont#”

- field.
 - ii. In the window that opens, click on “New.”
 - iii. Select the appropriate package type.
 - iv. Select the item(s) that are being placed in the new container.
 - v. Click on “OK” and enter password when prompted.
 - vi. Click the barcode button again, ensure that the newly created container is selected, and then click on print label.
 - vii. Attach the barcode label to the container.
- b) Verify that the package(s) is/are properly sealed and labeled. At a minimum, packages should be labeled with the following information:
- i. case number
 - ii. item number(s) included
 - iii. analyst initials
 - iv. barcode label
 - v. county and agency (this information should be on the barcode label)
- c) For evidence that will be physically transferred by another individual (such as an evidence technician) the following steps will be taken to identify evidence pending transfer:
- i. Complete the evidence transfer form ([OSBI CSD QPA 7.3](#)). Attach the form to the evidence with a staple or paper clip. Do not use tape.
 - ii. Move the evidence to a location specified for evidence transfers.
- d) Following transport of the evidence (if applicable) to the target lab, the following actions will take place:
- i. The individual receiving the evidence at the target lab will remove the transfer form from the evidence and check to see if a case file exists for the target unit. If a case file exists, the transfer form will be routed to the target unit. If no case file has been created yet for the target unit, one will be created and the transfer form will be attached to the outside of the file and then routed to the target unit.
 - ii. The evidence being transferred will be placed in the target unit’s property room or vault in the location designated for pending evidence.
 - iii. The supervisor or designee of the target unit will review the information on the transfer form and ensure that the case is assigned to an analyst or prioritized as necessary.
 - iv. The transfer form may be retained in the case file while the assignment is in progress, but will be shredded prior to the assignment being closed.

C. Evidence Returns

With the exception of evidence samples which require or warrant retention, evidence will be returned to the appropriate submitting or requesting agency.

1. Evidence will generally be returned in person but may be returned by certified mail (with return receipt requested) or private courier (UPS, FedEx).
2. For evidence returned at an OSBI CSD facility, perform the following steps:
 - a) Scan the barcode for “return to agency.”
 - b) Scan the barcode(s) for the item(s)/container(s) being returned.
 - c) Type any comments necessary in the “Comments” field. For example, if evidence is returned to an agency other than the requesting agency, an explanation or comment should be entered.
 - d) Scan the barcode for “process chain of custody” or click on “Save.”
 - e) Enter your password and click “OK.”
 - f) Type in the name of the individual receiving the evidence, have them sign on the signature pad, and click “Save Signature.”
 - g) Give a printed copy of the evidence receipt to the individual.
3. For evidence returned at remote locations, perform the following steps:
 - a) Transfer the evidence to the custody of the individual that will be returning the evidence.
 - b) Prepare (a) hard copy evidence release form(s) ([OSBI CSD QPA 7.2](#)).
 - c) Deliver the evidence to the appropriate agency and have the individual receiving the items print and sign his/her name on each form and date the form(s).
 - d) Document the date and time of the transfer.
 - e) After returning to a CSD facility, attach an imaged copy of the signed release form to the appropriate BEAST case file.
 - f) Update the chain of custody record to reflect the return to agency. This can be done by coordinating with the FSC Evidence Unit Supervisor or the Administrative Programs Officer over the Evidence discipline.

D. Evidence Destruction

The following procedure for evidence destruction applies to items that are currently in the custody of the OSBI CSD or evidence and other items, with the exception of pharmaceutical items, brought to the OSBI CSD specifically for destruction.

1. Destruction of toxicology evidence will be handled in a manner that complies with O.S. Title 47, Section 751-759.

2. Drug and other evidence will be destroyed after receiving written authorization from the OSBI Case Agent, the submitting agency, the district attorney having jurisdiction in the case, or by applicable statutes. [OSBI CSD QPA 7.1](#) must be completely filled out by a law enforcement representative for the proper destruction.
3. Completed destruction forms should be attached to the electronic case file in the BEAST using the same procedure listed under section II.B.3.d above. The file name for the electronic copy of the destruction form should include the case number and an indication that the file is a destruction form.
4. Destruction forms do not need to be imaged for any evidence that has been returned to the agency and is then brought back to the OSBI solely for destruction.
5. The Administrative Programs Officer over the FSC physical evidence unit or the appropriate regional supervisor or physical evidence technician will coordinate the destruction of evidence.
6. Any laboratory facility may accept drug items for destruction. Items received for destruction should be maintained under seal, if possible. Destruction items are not considered evidence.
7. To update the chain of custody for destruction of evidence still in the custody of the OSBI CSD, perform the following steps.
 - a) Scan the barcode for the evidence disposition “destroyed.” If the destruction form is received by mail, scan the barcode for “destroyed, no signature.”
 - b) Scan the barcodes for the item(s)/container(s) being destroyed.
 - c) Enter the destruction number in the comments field, if applicable.
 - d) Scan the barcode for “process chain of custody.”
 - e) Enter your password and then click on “OK.”

III. Attachments

[OSBI CSD QPA 7.1, Rev. 1 Evidence Destruction Form](#)
[OSBI CSD QPA 7.2, Rev. 1 Evidence Release Form](#)
[OSBI CSD QPA 7.3, Rev. 1 Evidence Transfer Form](#)

I. Scope

The OSBI CSD recognizes that the quality of reagents, supplies, consumables, and services used are an integral part of providing quality and reliable test results. This procedure will be used to guide the purchase, receipt, and verification of reagents, supplies, consumables, and services required for testing procedures.

II. Procedure**A. Identification of Necessary Quality of Products/Services**

1. All reagents, supplies, or consumable materials used in analysis will be identified in analytical protocols proportional to the degree to which they impact the quality of the test. For example, a reagent or supply which does not impact quality may not be specifically listed or may be listed with no clarifying information (e.g. “methanol”). A reagent which does impact the quality of the testing will be described in fashion that ensures the proper quality of reagent or supply is ordered and used for the test (e.g. “methanol – 95% purity” or “methanol – reagent grade”).
2. Technical Managers (TM’s) may choose to consolidate this information into a single list to make it easier to identify reagents and supplies affecting quality of analysis as they are received.
3. Critical reagents or supplies, which must be tested to confirm the purity or quality prior to use must be identified by the discipline TM in analytical protocols or another suitable location.

B. Purchasing Quality Reagents, Supplies, and Consumables

1. An Internal Purchase Request (IPR) will be filled out with sufficient detail to identify the quality of reagent or supply needed.
2. If a substitution is necessary, the discipline TM should be consulted to determine if the substitution is acceptable.
3. IPR’s for items that affect the quality of the tests shall be reviewed and approved for technical content. This review may be done by the individual(s) with approval responsibility according to OSBI Policy 208, provided he/she has sufficient technical knowledge to ensure the appropriate quality product has been requested. If the approving individual does not have sufficient technical knowledge to conduct an appropriate review, he or she should consult with the TM.
4. When consultation with the TM is necessary for a review of technical content or for a substitution, it may be documented in one of the following ways, or an equivalent

method:

- i. by summarizing the consultation with TM in the comments field in the IPR system during the approval process;
- ii. by adding a memo or e-mail as an attachment to the electronic IPR.

C. Receiving Reagents, Supplies, and Consumables

1. Upon receipt, all reagents will be marked with the date the item is received. In addition, upon opening a container of a chemical or reagent for the first time, the analyst opening the container will initial and date the container. Date of receipt and date opened should also be marked on supplies or consumables if the age of the supply will impact the quality of the product.
2. All containers will indicate a lot number and expiration date, if applicable.

D. Storage of Reagents, Supplies, and Consumables

1. Once reagents and supplies are received, they should be stored according to the manufacturer's recommendations.
2. Reagents and supplies affecting the quality of analysis that have not been inspected or verified must be stored either in a separate location from those that have been inspected and approved, or in another manner which prevents uninspected materials from being used in casework.

E. Inspection of Quality Reagents and Supplies

1. At a minimum, the reagent or supply will be inspected to ensure that it meets the quality criteria established in the protocol. This may be as simple as comparing the part number, described quality (e.g. % purity, or grade), or other relevant information on the reagent or supply received to that included in the IPR and/or the appropriate technical protocol.
2. In addition to inspecting reagents or supplies upon receipt to ensure the proper quality of product was received, the quality of reagents and supplies will also be continually monitored through the evaluation of standards and controls established in discipline protocols. Any quality concern identified through the use of standards and controls will be handled according to QP 13, 14, or 15 as appropriate.
3. All reagents prepared in-house must demonstrate proper function. Function verification should include testing the reagent or item in the same manner it will be used in testing. Function verification should be completed prior to using the item for testing samples which could not be retested.

4. Reagents used for DNA analysis will be evaluated in a method which complies with the current Quality Assurance Standards for DNA Testing Laboratories.

F. Documenting Preparation and Inspection of Reagents/Supplies

1. If the reagent or product is inspected to verify that the proper purity or quality of reagent was received (in lieu of conducting a function verification test), a copy of the packing slip, invoice or other document should be marked to indicate the quality was verified, the date, and the initials of the person verifying the item. This documentation must be maintained if it is the only record that the quality of the reagent or product was verified.
2. Any product inserts (or Certificates of Analysis) received which indicate the quality or purity of a reagent, should be retained. Unit Supervisors are responsible for ensuring these are properly retained and communicating to staff where the inserts and/or certificates will be retained. The lot number(s) and date received should be noted on the document.
3. All laboratory prepared reagents will be entered onto an OSBI chemical formulation worksheet ([OSBI CSD QPA 8.1](#)) and will include a short narrative or description of the preparation of the reagent. Alternately, each discipline may create their own chemical formulation worksheet (hard copy or electronic), provided that it contains, at a minimum, all of the information included on [OSBI CSD QPA 8.1](#).
4. Function verifications will be recorded on the chemical formulation worksheet along with the name or initials of the analyst testing the item and the date approved. Function verification of reagents which are a component of a reagent mixture can be documented on the chemical formulation sheet for the mixture.
5. Small quantities of reagents made, used, and discarded on the same day (e.g., Takayama) will have a chemical formulation sheet on file listing the component, quantities, and making of the reagent. The chemical formulation sheet for that reagent should also contain information as to function verification. The function verification information for the reagent will be included in the case notes. A new chemical formulation sheet does not have to be made for each daily use.
6. The OSBI chemical formulation worksheets will be retained. These records must be available for inspection. The chemical formulation worksheet will be maintained in a reagent logbook for each discipline or in the Chemical Inventory system in the BEAST. The Supervisor of that unit is responsible for maintaining archived chemical formulation sheets.

7. The reagent container must, at a minimum, bear the name of the reagent, the identity of the individual preparing the reagent (for reagents prepared in-house), and the lot number. The preparation date will be used for the lot number for in-house reagents. For example, a reagent prepared on July 20, 2003 would have the lot number 72003. Expiration dates should be placed on the front of the container if applicable.
8. Documentation of reagent preparation and evaluation of reagents used for DNA analysis will be maintained according to the Forensic Biology Quality Manual.

III. Attachments

[OSBI CSD QPA 8.1, Rev. 1 Chemical Formulation Worksheet](#)

I. Scope

Critical reagents, supplies, and services which affect the quality of testing will be obtained from reliable suppliers. The following process will be used to evaluate suppliers to determine the reliability of services and/or supplies they provide.

II. Procedure**A. Evaluation**

One of the following methods will be used to evaluate suppliers of critical reagents, supplies, and services.

1. The first time a new supplier is used, an in-house verification process will be performed to verify the quality of reagent supplied. If the reagent/supply ordered does not already have a verification process established, the Technical Manager (TM) will be responsible for determining the method appropriate for verification and determining what documentation must be maintained as a record of the verification. This verification should be performed using non-probative or non-casework samples before the reagent/supply is used for testing purposes and should verify that the reagent/supply is free from contamination and yields the expected results.
2. For suppliers used prior to the original issue of this document, historical data can be reviewed to confirm the supplier's reliability.

B. Documentation

The evaluation process and approved suppliers will be documented in the following manner.

1. The TM for each discipline will maintain a list of all approved suppliers for critical reagents, supplies, and services. The list should include the name of the supplier and the reagents, supplies, or services obtained from the supplier.
2. The evaluation process will be documented in the following manner.
 - a) Examination documentation and results of an in-house verification process will be maintained in a location designated by the TM.
 - b) Evaluation of suppliers which is based on historical data will be summarized in a memo by the TM and retained.

C. Handling Quality Problems with Reagents, Supplies, and Services

When quality concerns are identified with reagents, supplies, or services, the following response will be taken.

1. If the quality concern potentially impacts or did impact testing, the non-conforming work will be reported and documented according to [QP 13](#) and a corrective action identified and implemented according to [QP 14](#).
2. The supplier will be removed from the list of approved suppliers if, based on the severity of the quality concern, the supplier's response to the concern, and the record of past transactions with the supplier, such action is appropriate.
3. Any significant quality concern will be documented and a copy of the documentation will be forwarded to the OSBI purchasing office.
4. If a supplier is removed from the list of approved suppliers, the purchasing office will be notified and steps taken to file a vendor complaint with the Department of Central Services according to OSBI Policy 208.

III. Attachments

None

I. Scope

The OSBI CSD will cooperate with customers to clarify service requests and allow monitoring of CSD progress and/or testing as necessary. This procedure outlines methods to ensure this service while protecting the confidentiality of other customers.

II. Procedure**A. Consultations with Customers**

1. In order to ensure that both OSBI CSD employees and the customer understand the services that will be provided and the customer is informed of the progress, CSD employees shall be available for consultation.
 - a) Analysts and/or Supervisors will be available to consult with customers at the time evidence is submitted. Individuals receiving evidence will contact an analyst/Supervisor when necessary.
 - b) Analysts and/or Supervisors will contact customers when significant changes (e.g. projected completion dates, additional analysis possible/recommended, etc.) occur.
 - c) Analysts will be available to meet in person with officers or attorneys to review evidence or explain testing results.
 - d) CSD employees will provide updates on the progress or status of work as requested. For open assignments which have been given to an analyst, status inquiries will be forwarded to the appropriate analyst.
2. Locations for Consultations
 - a) Consultation rooms are available at the FSC to allow for private discussion and examination of evidence.
 - b) Consultation may also be conducted in an alternate lab space or location, but CSD employees will ensure that customers are not given access to confidential information from other cases. This includes case files and evidence packaging with suspect/victim information on it.

B. Requests to Monitor/Observe Analysis

In order to ensure confidentiality of case information, limit potential for contamination, ensure security of evidence and case records, and to provide the best service possible to all customers, outside observers will not be allowed in laboratory spaces under normal

circumstances, unless expressly ordered by the court. However, under certain circumstances, such as the analysis of consumption samples, monitoring of analysis may be necessary. Requests will be evaluated in the following manner.

1. The analyst receiving a request to observe testing will forward the request through his/her Supervisor to the Criminalistics Administrator (CA).
2. The CA will evaluate the request to determine how critical the requested analysis is, whether the observation would be better accommodated through an alternate method, and what the impact of allowing the observation will be on other pending cases (e.g. loss in productivity, etc.). The CA will consult with the customer, analyst, CSD Director, and OSBI Legal Counsel as needed to determine the appropriate course of action.
3. Once the course of action is decided, the CA will inform the analyst, the customer, and any other personnel as appropriate.

C. Performing Analysis with an Observer

When observation of analysis is ordered by the court or agreed upon by the OSBI CSD, the following steps will be taken.

1. If practical, the analysis will be conducted in the training laboratory at FSC. Prior to analysis, the analyst will ensure that all instruments are up to date on maintenance and quality control checks. The analyst will also obtain a key to secure the appropriate room(s) as necessary during the analysis process.
2. If the analysis cannot be conducted in the FSC training lab, the analysis may be conducted in other OSBI CSD facilities. Under these circumstances, if necessary to ensure confidentiality of other cases, the analysis will be scheduled outside normal working hours and at the convenience of the OSBI analyst.
3. The observer(s) will wear the appropriate personal protective equipment (PPE) as instructed by the analyst.
4. The observer(s) will be asked to refrain from speaking during certain portions of analysis, such as sample transfers, to reduce the potential for distractions.
5. If the observer(s) will be in areas where DNA evidence is handled, the observer(s) will be expected to provide a DNA sample.
6. The observer(s) may be asked to provide identifying information so that a background check can be completed.

D. Alternate Methods for Monitoring/Observation of Testing

The following alternate methods of observation may also be proposed.

1. The OSBI CSD may elect to send the samples to an independent laboratory for analysis.
2. The analysis may be videotaped and a DVD documenting the analysis provided to the customer. Under these circumstances, the customer will be expected to reimburse the OSBI CSD for the cost of videotaping and providing the DVD's.
3. If possible, a closed circuit feed may be set up to enable the observer(s) to monitor analysis from a separate room or location.

E. Requests for Assistance with Verifications or Technical Reviews

In some circumstances, some customers (such as local police laboratories) may request assistance conducting verifications or technical reviews of work they've performed. These requests may be met by performing work in-house or by conducting work as if a contract employee of the customer's lab. All requests for assistance with verifications or technical reviews must be forwarded through the chain to the CSD Director, who will coordinate necessary discussion with the customer.

1. In-House Analysis

A request for verification may be handled by analyzing the case in the same fashion as any other case, following all applicable discipline and CSD policies and procedures.

2. Conducting "Contract" Work

Alternately, verifications can be conducted by OSBI employees working as "contract" employees of the customer's laboratory. In order to do this, the OSBI employee must receive training from the customer regarding the applicable policies and procedures. The OSBI employee will then perform and document the requested verification(s) according to the customer's policies. Any request for assistance performing technical reviews for another laboratory must be handled in this same fashion.

Prior to conducting this type of work, OSBI employees must prepare a Memorandum of Understanding (MOU) detailing the work to be done. The MOU should be signed by the CSD Director and an authorized employee at the customer laboratory before any work is conducted.

III. Attachments

None

I. Scope

The OSBI CSD will routinely solicit customers for positive and negative feedback to identify any potential areas for improvement of the CSD management system and testing services.

II. Procedure**A. Feedback Regarding Testimony**

In order to solicit feedback regarding courtroom testimony provided by CSD personnel, the witness critique form will be distributed as indicated in [QP 32](#).

B. General Feedback

The following methods will be used to solicit feedback regarding services provided by the OSBI CSD.

1. The OSBI Forensic Laboratory Survey ([OSBI CSD QPA 11.1](#)) is available on the OSBI website.
2. In addition, hard copies of the survey should be available at each CSD facility.
3. CSD employees may also complete applicable portions of the survey (such as additional comments – ideas for improvement or complaints) based on feedback received during a consultation or conversation with a customer. The employee receiving the feedback should ask the customer for his/her permission to document the comments and for permission to include the customer's contact information.
4. The CSD Administrative Staff or a designee may contact customers for additional input as needed.

C. Analysis of Feedback

Feedback received from customers will be used and analyzed in the following manner.

1. Suggestions for improvement will be forwarded for consideration as preventive measures according to [QP 15](#).
2. Critique forms and survey results will further be analyzed during the management system review to identify trends which indicate further opportunity for improvement. See [QP 18](#).

III. Attachments

[OSBI CSD QPA 11.1, Rev. 4](#)

I. Scope

This policy is applicable to internal and customer complaints of a technical or administrative nature. It does not apply to personnel or human resources issues which should be forwarded to the appropriate Supervisor. Complaints may identify opportunities for improvement within the quality system. Valid complaints will be dealt with in a responsible and appropriate manner.

II. Procedure**A. Filing a Complaint**

1. CSD employees wishing to file a complaint will route a memo that thoroughly describes the issue to the designated Criminalistics Administrator (refer to [QP 1](#)).
2. If a customer indicates dissatisfaction with the OSBI CSD, the person who is the recipient of the complaint should provide the name and phone number of the designated Criminalistics Administrator (CA) to the customer. The employee receiving the complaint should also offer to document and route the complaint according to this policy.

B. Verifying and Acknowledging Complaints

1. Upon receiving a written complaint, the CA, or designee, will begin a Complaint Tracking Form ([OSBI CSD QPA 12.1](#)) and assign a tracking number.
2. The CA or designee is responsible for investigating the condition(s) stated in the complaint. If the condition(s) can be verified, the complaint will be reviewed to determine its validity. Validity will be determined based on the significance and impact of the condition. The purpose of validity screening is to eliminate complaints that do not deal with substantive or appropriate issues. The verification and validity status will be identified on the Complaint Tracking Form.
3. Following the verification and validity screening, the CA will notify the complainant of the status of the complaint. This notification may be oral, written, or by e-mail. The notification will be documented on the Complaint Tracking Form.

C. Investigating and Resolving Complaints

1. The CA or designee may forward the complaint package to the appropriate Technical Manager (TM) or Supervisor for investigation and determination of appropriate action(s). When forwarding complaint packages, the CA or designee should include an appropriate timeline. The selected manager's name and date will be entered on the Complaint Tracking Form.

2. The TM or Supervisor selected in II.C.1 is responsible for investigating the situation, condition, or action that caused the complaint and recommending a course of action, if necessary, to remedy, as appropriate.
3. The selected TM or Supervisor will report the cause and recommended actions to the CA. The CA will enter the date the report was received on the Complaint Tracking Form and forward the report to the appropriate manager for approval of proposed actions. When approval is obtained, the selected manager will implement the approved actions.
4. The CA will track the progress of the complaint process to ensure timeliness and will periodically analyze the complaint instances to determine if there are systemic or underlying problems that require attention.

D. Final Notification

Upon completion of actions dealing with a complaint, the CA will notify the complainant that the complaint has been resolved. The completion date and notification date will be documented on the Complaint Tracking Form.

III. Attachments

[OSBI CSD QPA 12.1 Rev. 1](#)

I. Scope

This procedure will be used as an evaluation tool to determine the proper method for documenting and addressing work that does not comply with OSBI CSD policies and procedures or meet the agreed needs of the customer. This procedure should be applied to work that impacts the accuracy of results, the accuracy of the audit trail, and the integrity of evidence.

II. Procedure**A. Responsibilities and Authority**

1. It is the responsibility of every CSD employee to report all observations of non-conforming work.
2. Every Supervisor and Technical Manager (TM), in addition to all CSD administrative staff, is responsible for and has the authority to immediately suspend any observed non-conforming work activity that could result in erroneous reports or unreliable testing data.
3. Authority to resume work that has been suspended lies with the appropriate TM or Criminalistics Administrator (CA). The authority to resume suspended DNA work belongs to the Forensic Biology TM.

B. Levels of Non-Conformance

There are three key levels of non-conforming work.

1. Simple Correction – The nature of the non-conforming work is limited in scope and significance. The problem identified is easily corrected and does not cast doubt on the overall reliability of results.
2. Corrective Action Necessary– The nature of the non-conforming work is such that it will continue to occur without a proper root cause analysis and appropriate corrective action. While corrective action is necessary, there is still no doubt regarding the overall reliability of test results.
3. Non-conforming Results – The nature of non-conforming work is such that the reliability of test results is questioned. There is potential that erroneous or invalid results have been reported. Corrective action will be required, but it is imperative to first address suspension of work and recall of reports.

C. Simple Correction

Simple corrections will be addressed in the following manner.

1. Instances of protocol drift or other errors which are limited in scope and significance will be corrected and the correction documented in an appropriate location in the case record. The supervisor, or designee, will approve the correction before the report is issued. He or she will document his or her approval in an appropriate manner, such as initialing the correction or placing a narrative in the BEAST file.
2. Supervisors will be responsible for maintaining a log of simple corrections required and monitoring that log for patterns or repetition of the same types of errors. When simple corrections are routinely required under the same circumstances or for the same individual, the supervisor will be responsible for initiating a corrective action request according to [QP 14](#).
3. Technical managers of disciplines with more than one unit will be responsible for establishing the method for logging simple corrections that impact their discipline. TM's may defer this responsibility to supervisors, but must ensure that a periodic evaluation of simple correction logs is conducted to monitor for patterns or repetition that is spread among different units.

D. Corrective Action

Errors that occur and that will continue to occur without corrective action but which do not cast doubt on the reliability of test results will be addressed according to [QP 14](#).

E. Non-Conforming Results

When non-conforming results are identified, the following actions will be taken. It is imperative that the actions described happen as quickly as possible and that the documentation be forwarded as soon as is practical.

1. The individual that identified the non-conforming work will complete sections I, II, and III of the Non-Conforming Results Report ([OSBI CSD QPA 13.1](#)). He or she will then provide a copy of the report to the Supervisor and Technical Manager (TM). If the Supervisor and TM are the same individual, the reporting individual will also forward a copy of the report to the appropriate Criminalistics Administrator (CA).

2. Upon receiving the report, the Supervisor of the impacted area will consult with the TM as necessary and immediately conduct an initial assessment as indicated below. The supervisor's assessment will be recorded in Section IV of the form.
 - a) Evaluate the **scope** of the non-conforming work.
 - i. Determine and document whether the non-conforming work is limited to the case/event reported or if the non-conforming work may extend to other cases/work.
 - ii. If the full scope of the non-conforming work is not immediately apparent, document what steps must be taken to identify all work potentially impacted, including who will be responsible for all steps and when the review will be completed.
 - b) Evaluate the **significance** of the non-conforming work.
 - i. Document whether the non-conforming work impacted results. For example, were incorrect results reported or were results invalid due to the non-conforming work?
 - ii. If the significance of the non-conforming work is not readily apparent, determine what steps must be taken to further investigate the matter and document the plan. Include in the plan who will be responsible and when the investigation should be completed.
 - c) Based on the scope and significance of the non-conforming work, take appropriate action and **document the actions taken** on the non-conforming results report.
 - i. Have erroneous or invalid results been reported? If so, recall work or issue amended reports and contact the customer to explain the non-conforming work, as appropriate.
 - ii. Is there a need to suspend work activities? Suspension of work may pertain to an individual analyst, a particular method, etc. Any suspension of work must be clearly communicated to the employees affected and should be limited to the work activities impacted.
 - iii. Is corrective action required? Generally, corrective action will be required for non-conforming work which meets one or more of the following criteria:
 - a. The scope of the non-conforming work is broad and impacts work conducted by multiple analysts and/or over a range of time.

- b. The significance of the non-conforming work is serious. Incorrect or invalid results were reported.
 - c. The problem could recur without corrective action.
 - d. There is doubt regarding whether OSBI CSD operations comply with agency and/or CSD policies and procedures.
 - e. The non-conforming work has been previously reported, regardless of scope or significance.
- d) Determine the most appropriate method for remediation of the non-conforming work.
- i. If corrective action is not required, document the method for remediation in the appropriate location on the non-conforming work report.
 - ii. If corrective action is required, but it is most appropriate to remediate the reported non-conforming work prior to the completion of the corrective action plan, document the proposed method of remediation on the non-conforming work report.
 - iii. If corrective action is required and the remediation of the reported non-conforming work would best be addressed in conjunction with the corrective action plan, indicate this on the non-conforming work report.
- e) If corrective action is required, conduct a root cause analysis and develop a corrective action plan according to [QP 14](#).

F. Review of Non-Conforming Work Reports

Once the Supervisor has completed section IV, he or she will route the report to the TM or the appropriate CA as indicated below. The reviewer will take the steps indicated to review and approve the report.

1. Section V will be filled out by the following individual(s):
 - a) Section V, the initial review and approval will be completed by the TM, unless the TM is the same individual that completed Section IV.
 - b) If the Supervisor that completed Section IV is also the TM, then Section V will be completed by the appropriate CA.
2. The individual responsible for the review detailed in Section V will evaluate and document for each section, whether the steps taken were:
 - a) Adequate and approved, as is,
 - b) Require modification and approved with the modification specified, or
 - c) Are not adequate and require additional action.

3. If further action is needed, the reviewer will provide instruction for what additional steps must be taken and return the report back to the appropriate Supervisor.
4. Once the non-conforming work report has been approved, the TM or appropriate CA will route the completed form back to the Supervisor for implementation of the remediation, if applicable. The Supervisor will be responsible for documenting the completion of the remediation, and maintaining record of the non-conforming work report.

G. Resuming Work

When work has been suspended due to non-conforming work, the following actions will be taken to obtain and document a release to resume work:

1. If work was suspended during the evaluation of the non-conforming work, the decision to resume work will be made by the TM or the appropriate CA. Authorization to resume work will be documented at the bottom of the non-conforming work report.
2. If work was suspended because corrective action was required, the authorization to resume work will be conducted and documented according to [QP 14](#).

H. Notification of Administration

1. Supervisors will provide a copy of the non-conforming results report to the appropriate CA, the Quality Manager (QM), and CSD Director at the following steps in the process:
 - a) Once sections I through V have been completed by both the Supervisor and TM or CA.
 - b) Once sections VI and VII, if applicable have been completed.
2. The Quality Manager, CA's, and CSD Director retain the authority to direct Supervisors and TM's to take additional action and/or document additional information, if necessary.

III. Attachments

[OSBI CSD QPA 13.1 Rev. 1](#)

I. Scope

This procedure will be followed when non-conforming work or a departure from management system or technical procedures or policies has occurred and there is potential for recurrence if no corrective action is taken.

II. Procedure**A. Tracking Corrective Actions**

The following procedure will be used to track corrective actions, once they are initiated.

1. The individual who identified the circumstances requiring corrective action will complete sections I and II of the Corrective Action Request (CAR) form ([CSD QPA 14.1](#)) and will forward the CAR to the Quality Manager (QM) and to the supervisor of the impacted area.
2. The QM, or designee, will review the CAR to ensure that the issue does require corrective action according to this procedure. The QM may determine upon review, that the issue should be addressed as a simple correction or as non-conforming results. If the issue needs to be addressed as a simple correction or non-conforming results, the QM will advise the appropriate individual(s) how to proceed.
3. After completing the review the QM, or designee, will assign each CAR a tracking number and log the CAR onto the CAR tracking spreadsheet located at [\\Pm-fsc13000s\qa\Lab_System_Records](#).
4. The QM, or designee, will route the CAR to the appropriate supervisor(s) or designee(s) for cause analysis.

B. Cause Analysis

The process for developing a corrective action will start with a root cause analysis.

1. The Supervisor of the impacted area, or the appropriate designee(s), will conduct an investigation to determine the root cause(s) of the problem.
2. If the root cause is not obvious, a systematic analysis of all potential causes will be conducted. The Supervisor, or designee, will list all potential causes that are evaluated.
3. If necessary, the Supervisor, or designee, may create a committee or conduct a unit meeting to gather additional input regarding potential causes.

C. Developing Corrective Actions

Once a root cause is identified or all potential causes are listed, potential corrective actions will be listed and evaluated to determine the corrective action(s) most likely to prevent a future occurrence of the same type of problem.

1. The Supervisor, or designee, will list possible corrective actions and how each action would correct the root cause(s) or potential cause listed.
2. After compiling a list of possible corrective actions, the Supervisor, or designee, will review the list and select the corrective action(s) most likely to eliminate the problem and prevent a recurrence. The corrective action(s) must also be appropriate to the magnitude of the problem.
3. The Supervisor, or designee, will also list or describe the mechanism(s) that will be used to monitor the implementation of the corrective action(s) and determine whether the corrective action(s) has/have been effective.

D. Selecting and Implementing Corrective Actions

1. Once the Supervisor, or designee, has determined the most appropriate corrective action(s) and a method for monitoring, the supervisor will include his/her recommendation on or with the Corrective Action Request ([OSBI CSD QPA 14.1](#)).
2. The proposed corrective action plan and monitoring mechanism will be reviewed and approved by the appropriate Technical Manager (TM), or the appropriate Criminalistics Administrator (CA) if the TM is the same individual as the Supervisor.
3. Upon approval, the TM or CA will route the corrective action request back to the Supervisor.
4. The Supervisor, or designee, shall document and implement the changes required as part of the corrective action.

E. Monitoring Corrective Actions

The Supervisor will monitor activities according to the method approved as part of the corrective action plan.

1. If the monitoring of the corrective action indicates that it is not/was not effective, the matter will be re-evaluated for a subsequent cause-analysis and/or selection of an alternate corrective action. Supervisors will document the additional cause analysis and/or selection of an alternate corrective action on an attachment to the original corrective action request form.

2. If the nature of the non-conforming work indicates a failure to comply with laboratory policies/procedures or applicable accreditation standards, the appropriate areas of activity will be audited as soon as possible. The audit may be conducted following the implementation of corrective action to further assess the effectiveness of the corrective action. The supervisor of the impacted area will be responsible for coordinating and documenting this audit. He or she may request assistance from other audit trained supervisors and analysts, TM's, CA's, or the Quality Manager.

F. Authorization to Resume Work

If work was suspended in conjunction with the corrective action request, the Supervisor will be responsible for requesting authorization for work to resume. He or she will document the completion of the corrective action plan and return the form with any necessary documentation to the TM or the appropriate CA. The TM or CA will verify the corrective action plan has been completed and will indicate in Section VII of the form whether or not the resumption of work is approved.

G. Notification of Administration

1. Supervisors will provide a copy of the corrective action request form to the appropriate CA, the Quality Manager (QM), and CSD Director at the following steps in the process:
 - a) Once sections I through V have been completed by both the Supervisor and TM or CA.
 - b) Once sections VI and VII, if applicable, have been completed.
2. The Quality Manager, CA's, and CSD Director retain the authority to direct Supervisors and TM's to conduct and document additional cause analysis, monitoring, and corrective action, if necessary.

III. Attachments

[OSBI CSD QPA 14.1, Rev. 1](#)

I. Scope

All CSD employees are responsible for monitoring work flow, technical procedures, and management system practices for potential improvements and/or potential sources of nonconformities. CSD employees will follow this procedure for documenting, routing, implementing, and monitoring preventive actions which are not already covered under another procedure.

II. Procedure**A. Recommending Preventive Actions**

1. Any CSD employee who identifies a potential source for non-conforming work or improvement to the CSD technical operations or management system must submit a suggestion in writing through his/her supervisory chain. The written suggestion (which may be in memo or e-mail format) must include the following elements:
 - a. A description of the problem or opportunity for improvement,
 - b. An explanation of any potential for nonconforming work,
 - c. A proposed action plan or description of the steps necessary to implement the suggestion, and
 - d. A proposed control mechanism for monitoring the effectiveness of the suggested change.
2. Any CSD employee who receives a suggestion for improvement or preventive action from a customer shall forward the information according to section II.A.1 above.

B. Review and Approval of Preventive Actions

1. Preventive actions will be reviewed and approved at the lowest management level appropriate to the suggested change. For example:
 - a. Proposed changes to work flow processes impacting a single unit should be reviewed and approved or disapproved by the unit supervisor.
 - b. Proposed changes to technical procedures should be reviewed by any impacted supervisors and the appropriate technical manager, but approved or disapproved by the technical manager.
 - c. Proposed changes to case acceptance policies or other changes which may impact customer service or satisfaction should be routed to and approved or disapproved by the CSD Director.

2. Individuals reviewing and/or approving suggested preventive actions should research further or make modifications to the suggestion as necessary to ensure that it complies with section II.A.1 above.

C. Implementation and Documentation of Preventive Actions

1. The individual approving a preventive action will be responsible for directing the implementation of the plan and monitoring the implementation and effectiveness of the plan. Alternately, the approving individual can designate one or more individuals to implement and monitor the plan.
2. Supervisors will be responsible for maintaining documentation of preventive actions that are proposed. Supervisors will also be responsible for reporting on the status of preventive actions in accordance with [QP 18](#).
3. Individuals who review and/or approve preventive actions should ensure that the status of the review, approval, and implementation is communicated to affected employees in a timely fashion.

III. Attachments

None

I. Scope

This procedure will be used for quality and technical records to ensure that they are readily identifiable and retrievable, protected from damage, and kept confidential.

II. Procedure**A. Identification of Records**

Quality and technical records, whether hard copy or electronic, will be identified in the following manner.

1. Case files are identified by the laboratory case number. The case file may be further identified by the unit or discipline, when necessary.
2. Technical records which are not stored in the case file, such as quality control records associated with batched cases, will be identified in a manner that facilitates associating the data with the proper case(s). For example, quality control results could be identified by an instrument name or number and date/time of the run.
3. Quality records should be identified with sufficient detail to facilitate proper filing and storage.

B. Indexing/Filing Records

Technical and quality records will be indexed and filed according to the record identification.

1. Case files will be stored numerically according to the case number.
2. Quality records will be indexed according to subject, location, and/or date.

C. Collection and Storage of Records

Hard copy technical and quality records will be stored in designated areas with appropriately controlled access.

1. Unassigned case files pending analysis will be stored in a secure location designated by the supervisor.
2. Case files and technical records for cases in the process of examination will be stored in the analyst's work area or other appropriate and designated location.
3. Completed case files will be stored in a file room or other designated secure area of the appropriate CSD facility until they are archived.

4. Technical records such as quality control results, reagent logs, etc., will be stored in an orderly fashion in (a) location(s) designated by the supervisor.
5. Quality records will be stored as specified by the Quality Manager (QM).

D. Access of Records

1. Access to quality and technical records will be limited to those CSD employees that require access to conduct analysis and assist customers. This includes management, analysts, and physical evidence and analytical technicians.
2. Other CSD employees, practicum students, contractors, and visitors will be restricted from accessing technical and quality records according to [QP 20](#).
3. Access will be limited by restricting access to the physical storage location (e.g. file room).
4. Access to electronic records will be further restricted by issuing user names and passwords and setting appropriate permissions.

E. Maintenance of Records

1. Technical and quality records may be maintained in hard copy or electronic format.
2. When case files are maintained in an electronic format and no hard copy file is created, the BEAST barcode which is automatically generated to track the hard copy file will be scanned to the location code “electronic case file.”
3. Technical records will be maintained in their entirety for a minimum of ten years from the initial receipt of evidence.
4. Quality records will be maintained for a minimum of one accreditation cycle or five years, whichever is longer.
5. Management system documents will be maintained indefinitely.
6. Original records will not be removed from OSBI CSD facilities, with the following exceptions.
 - a) Case files may be removed for the purpose of referencing during courtroom testimony or meetings with attorneys or officers.
 - b) Case files or other technical records will only be removed from OSBI laboratory facilities for regular business purposes such as transfer of cases, court, conferences with court officials or investigators, or with permission of the unit supervisor or Criminalistics Administrator over the unit.

- c) Quality records may be removed from OSBI CSD facilities only at the permission of the QM.
- 7. Removal of completed hard copy case files will be documented by scanning the case file barcode or by inserting a piece of card or paper, labeled with the case number, examiner's initials and date of removal, in the specific location from which the file was removed.

F. Disposal of Records

- 1. Hard copy records may be disposed of once converted to an electronic format for archiving.
- 2. With the exception of drug and toxicology records, technical records will be retained in either hard copy or electronic format, indefinitely. Drug and toxicology technical records will be maintained in hard copy or electronic format for a minimum of six years. After six years, the following documents may be disposed of and not retained in either hard copy or electronic format.
 - a) any subpoenas that have been placed in the file
 - b) officer reports and information, with the exception of the officer affidavit and consent to test blood for toxicology cases
 - c) instrument scans
 - d) any duplicate documents
- 3. When disposing of quality and/or technical records, the documents will be shredded or otherwise disposed of in a manner that ensures the confidentiality of the information within the documents is protected.

G. Electronic Storage of Records

Electronic records will be stored utilizing the BEAST Laboratory Information Management System (LIMS), the LaserFische system, or on a network server.

- 1. Documents stored in the BEAST will be protected in the following manner:
 - a) Access to the documents will be limited through the use of a user name and password with appropriate permissions specified.
 - b) Alterations to completed documents will be tracked through the system's audit log.

- c) Information in the system will be replicated between servers at each CSD facility and backed up on a regular basis.
2. Documents stored on a network server will be protected in the following manner:
- a) With the assistance of the Information Technology Services (ITS) Division, access to the folder(s) will be limited to the appropriate individuals.
 - b) Back up of the files will be coordinated through the IT Division.
 - c) Records stored in this fashion will be saved as pdf files or another file format which prevents unintended and/or unauthorized alteration.

III. Attachments

None

I. Scope

This procedure details the administrative and technical documentation which must be maintained for all forensic analyses performed.

II. Procedure**A. Documentation Required for Case Records**

The following documentation is considered a technical record of analysis performed and must be maintained.

1. Administrative Documentation

- a) submission information
- b) evidence inventory
- c) conversations and communications
- d) copies of reports
- e) documentation of administrative and technical review

2. Examination Documentation

- a) Examination documentation includes notes concerning the analysis of evidence, scans, chromatographs, and all other documents produced and used to derive conclusions in the analysis of the case. The following details must be included in the examination documentation or another appropriately specified location in the case record:
 - i. The method used for analysis.
 - ii. The condition of the item, as it was received.
 - iii. If applicable, a reference to the sampling plan used for analysis and date and location of sampling.
 - iv. The date(s) that analysis was conducted or the start and end dates of analysis, at a minimum.
- b) Examination documentation must be sufficient to establish an audit trail and identify relevant documentation (calibration records, staff records, etc.).
- c) Examination documentation will include information identifying factors affecting uncertainty of measurement, where possible. This includes the identity of instruments used, personnel conducting each step of analysis, software used, etc.

d) Observations, data, and calculations will be recorded and appropriately identified at the time they are made.

3. Supporting Technical Documentation

a) Quality control results, including standards, ladders, calibrators and positive and negative controls are considered technical documentation/records.

b) Supporting technical documentation may be stored in the case file or in an alternate designated location.

B. Guidelines for BEAST Case Files

1. All administrative documentation should be maintained in the BEAST case file.

2. Any examination documentation which can be readily documented electronically should be maintained within the BEAST.

3. Alterations made to records in the BEAST will be tracked through the program's audit log.

C. Guidelines for Hard Copy Case Files

1. Any administrative or technical documentation which is required to be retained but is not readily incorporated into the BEAST case file should be maintained in the hard copy case file. For example, documentation received in hard copy format (faxes, instrument printouts) must be maintained in the hard copy file if they are not imaged into the BEAST case file.

2. No evidence items should be stored in the Criminalistics case file, with the exception of latent lifts, photos, and/or negatives. Latent lifts will be placed into a sealed manila envelope that is clearly marked with case number, examiner's initials, barcode, and date. The items may then be placed into the case file, providing the case files are located in a secure evidence storage location.

3. All notes, forms and documents generated by OSBI personnel (with the exception of latent evidence) shall be utilized on one side only. No two-sided forms or yellow sticky notes will be used.

D. Maintenance of Hard Copy Case File Documentation

1. All paperwork in the case file will be clearly identified with the Criminalistics' case number and handwritten initials.

2. Additionally, all Criminalist generated paperwork will be clearly identified with the examiner's initials, date, and item numbers if applicable. In some circumstances it may be acceptable for the date not to be printed on each page, provided the date the work was generated can be determined through other documentation in the file.
3. Notes generated by analysts shall be sequentially numbered for each assignment completed. When analysts conduct analysis at a later time on a separate assignment, the subsequent notes may be sequentially numbered (starting with page 1) or added to the sequence of notes already in the file. The first page of notes will be labeled to indicate the total number of pages in the set. (e.g. 1 of 5)
4. The case file will be orderly, complete and concise, thus facilitating administrative and technical review. Notes will be neat, readable, and written in ink. Notes may be typed.
5. Any corrections to notes will be made by an initialed single strikeout. Nothing in the handwritten information should be obliterated or erased and additions to notes (interlineations) must be initialed by the person making the entry.
6. It will be the responsibility of the examining Criminalist to ensure the contents of the case file are in compliance with the above sections.
7. It will be the responsibility of the unit supervisor to oversee case work and case files completed by their unit to ensure they are in compliance with all existing policies and analytical protocols.

E. Documenting Limited Samples

All evidence received of insufficient quantity to allow a representative sample after testing should be documented in the following manner.

1. The District Attorney involved in the case will be advised by the examining Criminalist prior to limited quantity samples being analyzed and consumed. A letter from the appropriate prosecuting attorney authorizing the consumption of those samples will be placed into the case record.
2. The evidence will be photographed. A ruler or size standard will be included in the photograph, if possible.
3. Photographic documentation will be made as necessary according to the appropriate discipline protocol.

4. Every reasonable attempt will be made to comply with any special request regarding the analysis of limited quantity evidence. These contacts and any special requests should be documented in an appropriate fashion (memo, e-mail, narrative, etc.) in the case record. The Division Director should be notified of any special requests.

III. Attachments

None

I. Scope

Internal audits of the OSBI CSD facilities and functions will be conducted according to this procedure.

II. Procedure**A. Scheduling Audits**

1. The OSBI CSD will conduct internal audits annually for each of the following facilities:
 - a) Forensic Science Center (FSC)
 - b) Northeast Regional Laboratory
 - c) Northwest Regional Laboratory
 - d) Eastern Regional Laboratory
 - e) Southwest Regional Laboratory
2. If necessary, audits of facilities may be further sub-divided into specific units/functions, provided all units of each facility are audited annually.
3. A schedule of audits including the Unit/Program and audit dates for the following calendar year will be issued by the Quality Manager (QM) during the fourth quarter of each calendar year.
4. Audit schedules may be adjusted depending on conflicts with auditors or lab staff. All changes must be requested through the appropriate Criminalistics Administrator(CA) and approved by the QM.

B. Conducting Audits

1. Prior to the audit, the QM, or designee, will assemble an audit team.
 - a) The audit team at a minimum will consist of a lead auditor (normally the Quality Manager or designee) and other auditors responsible for specific areas or disciplines as assigned by the lead auditor.
 - b) Each auditor shall have training in the audit process. Training may be provided by an approved external source or conducted by the QM or designee.
 - c) Additional interagency or outside personnel may be requested to help in the audit process. These individuals will be included on the audit team at the approval of the CSD Director.

2. Once the audit team has been identified, the QM, or designee, will prepare a checklist of assignments indicating which criteria each individual will assess.
3. At the scheduled time of the audit, the audit team will assemble at the designated location. An opening meeting may be conducted if appropriate. Each auditor will review the appropriate documentation and/or conduct interviews in order to determine whether the work/operations conform(s) to the standard and applicable policies and procedures.
4. After completing the review and/or interviews, each auditor will report to the lead auditor and provide a summary of what was reviewed. In addition, each auditor will list or describe the objective evidence observed for any findings or non-conformances.
5. An exit meeting will be conducted to inform the supervisor(s) and/or laboratory director of the results of the audit.
6. The QM, or designee, will compile the information provided by auditors into an audit report. The audit report will be in the appropriate format as indicated below.
 - a) All OSBI CSD audits will be reported referencing the most current accreditation standards. Findings will be reported on a corrective action request (CAR) form ([CSD QPA 14.1](#)). Other minor issues that require correction and any suggestions or observations will be summarized in memo format.
 - b) For OSBI CSD facilities conducting DNA analysis, an audit report will also be completed using the most current version of the Quality Assurance Standards audit document.
7. The audit report(s) should be completed and provided to the appropriate supervisor(s) and/or lab director within 2 weeks of the audit.
8. Within 30 days of receipt of the audit report, the appropriate supervisor(s) and/or lab director will send a response to the CSD Director. Responses to any CAR's will be documented according to [QP 14](#). A response will also be required to address any minor issues or recommendations provided in the memo. This response should address what corrections and/or preventive measures have been taken and why.
9. If the root cause and corrective action cannot be completed within 30 days, the response will include a plan for completing these steps and (a) projected completion date(s).
10. The QM, or designee, will monitor the progress of corrective action plans submitted until all corrections and corrective actions are completed.
11. When necessary, the customer will be informed of non-conforming work and work will

be recalled. The QM, or designee, will monitor non-conformances arising from audits to ensure that customer notifications and work recalls are conducted in an appropriate and timely fashion.

C. Completing/Closing Audits

Once all corrections and corrective actions/corrective action plans have been completed and documentation provided to the CSD Director, the CSD Director will evaluate with the QM and/or the appropriate CA the completed audit response.

1. If necessary, the CSD Director or designee will request and/or obtain verification of the correction(s) and/or implementation of corrective action(s).
2. Once the response has been reviewed and determined to be complete, the QM or designee will send a memo to the supervisor and/or lab director indicating the audit is closed.

D. Submitting Annual Audit Reports

Each OSBI CSD laboratory must submit an Annual Accreditation Audit Report to ASCLD/LAB within 30 days of the laboratory's accreditation anniversary date.

E. Notifying NDIS

Each year, the CODIS Supervisor will prepare appropriate documentation to notify the NDIS Custodian of internal and external audits as required by the current NDIS Procedures. Biology and CODIS Supervisors will prepare responses to external DNA audit findings and forward them to the Biology Technical Manager. The Technical Manager, with assistance from the CODIS Supervisor if needed will forward a copy of external audit reports to the NDIS Custodian in the manner and timeframe required by NDIS Procedures.

F. Conducting Line Inspections

1. Unit Supervisors and regional laboratory Supervisors will conduct a line inspection annually.
2. Line inspections will be documented on the Line Inspection Form ([CSD QPA 17.1](#)).
3. Line inspections will be conducted approximately six months after the annual internal quality assurance internal audit. Line inspections can be conducted as many times as the Supervisor deems necessary outside of the annual requirement.
4. Supervisors conducting line inspections are to note any deficiencies and take corrective action as appropriate.

5. The Supervisor will provide copies of all line inspections to the CSD Director, the appropriate Criminalistics Administrator, and the QM.

III. Attachments

[OSBI CSD QPA 17.1, Rev. 1 Line Inspection Form](#)

I. Scope

The following procedure will be used to conduct management reviews. A list of topics covered by management reviews is included in section [4.15.1](#) of the Quality Manual.

II. Procedure**A. Committee Structure**

The following committees are established for the purpose of conducting management reviews.

1. The primary committee for conducting management review is the Quality Improvement Committee (QIC).
 - a) All CSD supervisors, technical managers, and the administrative staff will serve as members of QIC.
 - b) The Quality Manager (QM), or designee, will chair QIC.
2. The following subcommittees are established for the purpose of conducting appropriate portions of the management review. The QM may assign tasks to subcommittees to facilitate the management review process.
 - a) Chemistry Subcommittee
 - i. CSD Supervisors that have been or will be trained in drug analysis and toxicology will serve as members of the Chemistry subcommittee.
 - ii. The Controlled Substances Technical Manager (TM) will chair the Chemistry Subcommittee.
 - b) Biology Subcommittee
 - i. Any Forensic Biology TM, CODIS Administrator, and any CSD Supervisor that has been or will be trained in forensic biology casework or database analysis will serve as members of the Biology subcommittee.
 - ii. The Forensic Biology TM will chair the Biology Subcommittee.
 - c) Identification Subcommittee

- i. CSD Supervisors that have been or will be trained in the disciplines of Firearms, Latent Evidence, and Trace Evidence will serve as members of the Identification Subcommittee.
 - ii. The supervisor of the FSC Latent Evidence Unit will chair the subcommittee.
3. Subcommittees may solicit assistance from other qualified analysts as necessary to complete tasks assigned to them. Subcommittee chairs are responsible for avoiding conflicts of interest when completing tasks assigned to the subcommittee. For example, chairs should ensure that subcommittee members do not review their own proficiency tests.

B. Meeting Schedules and Agendas

1. QIC will meet at least quarterly.
2. QIC schedules and agendas will be coordinated by the QM.
3. Subcommittees will meet as needed.
4. Subcommittee meeting times and agendas will be coordinated by the subcommittee chairperson.

C. Documenting Management Reviews

The findings from management reviews and summaries of actions taken will be documented as follows.

1. During the first quarter of each calendar year, each supervisor will review the ASCLD/LAB Guiding Principles of Professional Responsibility for Crime Laboratories and Forensic Scientists with each employee in his/her unit/lab. Upon completion of the review, supervisors will include a statement documenting the review in the appropriate management system review report.
2. At least annually, management reviews will be summarized into a written annual report by each Unit and Lab Supervisor. This management review report will be prepared in memo format and must include, at a minimum, the topics listed below. Upon completion of the report, supervisors shall forward the review through the appropriate Technical Manager and Criminalistics Administrator to the CSD Director.
 - a. Documentation of review of ASCLD/LAB Guiding Principles (if applicable);
 - b. An evaluation of laboratory objectives, indicating whether they are still

- appropriate for the unit and/or lab;
- c. An indication of the status of annual policy/procedure review, including whether or not the policies and procedures are deemed suitable;
 - d. A summary of any reports made by the supervisor or managerial staff during the year, which might include legislative changes that have impacted the unit/lab, staffing levels, etc;
 - e. A summary of the outcome of internal audits, line inspections, and safety audits, including the status of any corrective action that was required;
 - f. A summary of corrective and preventive actions submitted or performed by the unit/lab during the reporting period and the current status of those actions;
 - g. A summary of any external assessments conducted during the reporting period, including the status of any necessary corrective actions;
 - h. A summary of the results of any inter-laboratory comparisons or proficiency tests completed during the reporting period;
 - i. An evaluation of the volume and type of work submitted and performed by the unit/lab that highlights any changes during the reporting period;
 - j. A summary of any customer feedback received during the reporting period;
 - k. A summary of any complaints received during the reporting period including the status of any improvements implemented as a result of the complaint(s);
 - l. Any recommendations for improvement;
 - m. Any other relevant factors (which might include quality control activities, resources, and staff training).
3. If management reviews generate findings which require action, the CSD Director will appoint an appropriate designee to monitor the actions taken to ensure that they are completed in a timely fashion.

III. Attachments

None

I. Scope

OSBI CSD employees will be properly trained and qualified prior to performing independent or unsupervised testing.

II. Procedure**A. Training Program Structure**

Each discipline will have a written training program as outlined below.

1. Each discipline's training manual/program must include the following sections:
 - a. Application of ethical practices in forensic sciences
 - b. General knowledge of forensic science
 - c. Applicable criminal and civil law and procedures
 - d. Overview of quality system
 - e. Applicable Safety Topics (blood borne pathogens, chemical hygiene, etc.)
 - f. Discipline specific topics
 - g. Presentation of evidence in court (when applicable)
2. Each section of the training manual/program will include a list of goals which must be met for the trainee to have the skills necessary to complete the duties listed in the trainee's job description.
3. Each section of the training program will also establish the tasks or activities that will be completed by the trainee in order to meet the stated goal.
4. Successful completion of the training program will be assessed through the use of competency tests as indicated below.
5. The training program should be reviewed annually. The discipline Technical Manager (TM) and the CSD Director must approve any changes.

B. Conducting Training

1. The TM will be responsible for the assignment of training for any new employee assigned to that discipline.
2. Prior to training any individual, an assessment should be done to identify his/her specific training needs. This assessment may include a review of his/her education, experience, and/or a quiz or other competency evaluation to assess his/her knowledge/skill level.

3. Based on the results of the assessment, the training program can be modified according to the knowledge, skills, and abilities of the trainee.
4. OSBI Criminalists competent in the assigned discipline may act as trainers, at the request of the TM.
5. New training goals will not be assigned until both the trainer and trainee are satisfied that current goals are understood.

C. Competency Tests

1. Regardless of education, qualifications, or past experience, each analyst must demonstrate competence in each **category of testing** through the successful completion of a competency test(s) prior to release for casework or crime scene duties.
2. In order to successfully complete a competency test, the analyst must achieve the intended results. Any discrepancies must be reviewed and re-training conducted as necessary to achieve the expected results, prior to the test being accepted as satisfactory.
3. The minimum components and objectives of a competency test are as follows:

- a. Analysis of an adequate number of unknown samples:

Unknown samples should be prepared and assigned under the direction of the TM. The samples should encompass the range of samples which the employee will be expected to test upon successful completion of the test. The number of samples should be sufficient to evaluate an employee's ability to select and perform proper testing methods in accordance with laboratory policy.

- b. Written report:

Results of competency tests should be reported in the same fashion as casework or database analysis. The report should then be reviewed to evaluate the employee's ability to accurately and clearly convey testing results and the significance of the results.

- c. Written and/or oral examination:

Each employee must also demonstrate an adequate knowledge of the area being tested through the completion of a written and/or oral examination. This portion of the competency test should include a mock trial, when appropriate. Prior to the assignment of a written and/or oral examination, the TM or designee should

identify the knowledge necessary to perform testing (e.g. specific technical knowledge necessary to perform testing, conduct trouble-shooting, etc. and/or specific knowledge and ability to convey the knowledge clearly to lay-people), develop questions to assess the knowledge level of the individual, prepare a scoring mechanism and/or key, and set the minimum score that will be accepted as passing.

4. The trainee must successfully complete all phases of the competency test before being released to perform independent testing.
5. Upon the successful completion of a/the competency, a memo indicating the successful completion and approval for the trainee to begin analysis will be submitted by the trainee's TM to the appropriate Criminalistics Administrator (CA) and the Quality Manager (QM). The memo should clearly specify the area of analysis the individual is approved to perform.

D. Documenting Training

1. Training Records

The FSC administration will maintain training records for each CSD employee. This includes information concerning the job description, education, training and continuing education of the employee. Individual training records will be stored in the appropriate folder located at [\\Pm-fsc13000s\qa\Individual Records](#). Each CSD employee is responsible for placing a copy of the following documents in his/her folder:

- a) all transcripts indicating any degrees conferred
- b) certificates or agendas for continuing education/professional development classes attended
- c) updated transcripts when additional courses are completed
- d) all memos approving individuals to perform analysis

2. Supervisory Training File

The supervisor and/or appropriate TM will be responsible for maintaining a file for each trainee which includes:

- a) A copy of a completed checklist of the knowledge and skills for each trainee.
- b) Copies of all evaluation tools used (written tests, sample lists, etc).
- c) The expected and reported results for any/all competency test(s).

3. Monthly Training Updates/Memos

The progress of each trainee will be documented in the form of a monthly update stating what training was completed during the previous month. The trainer will provide the monthly update.

- a) A copy of the monthly training update will be provided to the trainee, Supervisor, Technical Manager, and appropriate CA.
- b) The monthly update is in addition to any documentation or evaluation required by OSBI Human Resources.
- c) The update should be submitted to the appropriate individuals by the 5th working day of the month.

E. Re-training

When re-training employees the following steps should be taken.

1. Re-training should begin with an assessment of the training needs of the individual. This may include completing a cause analysis related to non-conforming work or evaluating contributing factors for incorrect responses on competency tests.
2. Re-training may be conducted using specific portions of the discipline training manual, or may be developed and tailored to the needs of the individual being re-trained. The training program used must be written and approved by the TM before use.
3. Any re-training conducted should be documented in the same fashion as initial training.

F. Continuing Education

Employee development is critical to the quality program of the laboratory. Laboratory employees must keep current on the latest techniques and technologies. The OSBI CSD supports the continuing development of its employees through various methods, including the following.

1. Attendance at professional meetings is encouraged. Employees should refer to OSBI Agency Policy 202.1 for more information on regulations concerning attendance at professional meetings. Required materials concerning the course must also be turned in to the training office.
2. OSBI CSD employees are also encouraged to recommend training classes which can be hosted in the FSC training rooms. Recommendations for training classes should be forwarded through the supervisory chain to the CSD Director.
3. The OSBI encourages attendance at local colleges and universities in areas related

to the job description of the employee by offering tuition assistance when funding is available as outlined in OSBI Agency Policy 202.2. A copy of a completed transcript for any course related to professional development should be forwarded to the QM.

4. An employee development plan is a part of each annual evaluation as required by Oklahoma law and OSBI Agency Policy 214.
5. Other sources of training and development include:
 - a) Courses offered by the Council on Law Enforcement Education and Training (CLEET)
 - b) Courses offered by the Office of Personnel Management
 - c) In-house seminars, employee conferences, training and technical meetings
 - d) FBI, DEA, or other outside training
6. An evaluation of any grant funded training or meeting will be documented on form OSBI CSD QPA 19.1 and submitted to the CA responsible for overseeing grants.
7. Current literature review is important to the development of employees. The TM (or his/her designee) of each discipline will assign/and or circulate articles of interest for their discipline. The TM will make sure each analyst has had an opportunity to read each article.

III. Attachments

[OSBI CSD QPA 19.1, Rev. 0](#)

I. Scope

The OSBI CSD maintains the integrity and prevents contamination of evidence and ensures the confidentiality of records by limiting access to restricted areas to authorized personnel.

II. Procedure**A. Facilities**

1. Access to laboratories and evidence rooms will be limited to employees assigned to the laboratory unit, evidence technicians assigned to the physical facility, Criminalistics Administrative personnel, and other personnel as authorized by the CSD Director on a limited or permanent basis. All other personnel including service/maintenance personnel will only have access to the laboratory and evidence room areas when accompanied by employees authorized to have access.
2. Laboratories and evidence rooms will remain locked. Any exception must be authorized by the laboratory administration. At the end of each workday, the last person leaving is to ensure the unit and building are secured and alarmed.
3. No exterior or lab access door shall be propped open under normal circumstances. If an exterior or lab access door is propped open (e.g. while moving equipment), the employee propping the door open will ensure that appropriate OSBI personnel are present to monitor who enters and exits, until the door is closed. It will be the responsibility of any employee opening an outside door during normal work hours to ensure that door is locked at the end of that activity.
4. When arriving at work and leaving work, Forensic Science Center (FSC) employees should use the employee break room entry.

B. Lock Security

Keys, proximity devices, and access cards which provide access to laboratory spaces or evidence storage areas will be tracked as indicated below. Each key, proximity device, and access card, should be engraved or stamped with a unique identification if not already provided. Excess keys which are not issued for use may not require a unique identification. However, these keys should be maintained in a secure location and be labeled in a manner that they can be associated with the coordinating lock.

1. Tracking Keys

- a) Key control logs ([OSBI CSD QPA 20.2](#)) will account for the number of assigned keys to a laboratory unit and a sign out log will document who has keys to each door, locking refrigerator, freezer, or work area drawer. A master key log will be maintained ([OSBI CSD QPA 20.4](#) or [OSBI CSD QPA 20.4b](#)) for all keys within a unit or regional laboratory. The total number of keys will be verified annually and recorded on the Master Key Log.
- b) Laboratory units having proximity access devices (key fobs, etc.) will use [OSBI CSD QPA 20.2](#) to document the issue and receipt of proximity access devices, excluding identification badge proximity access devices which are handled according to section II.B.2.a, below. Cipher lock codes will be maintained on [OSBI CSD QPA 20.3](#).
- c) Assigned keys will not be copied or loaned. Proximity access devices will not be loaned. Cipher lock combinations, access codes, and alarm codes will not be further disseminated by the individuals receiving the combination or code.
 - i. Loss of assigned keys and proximity access devices will be handled as per OSBI Directive 211. The employee shall immediately report the loss to their Supervisor.
 - ii. Previous key control log forms must be archived by the responsible person(s) designated in II.B.2. When key control log forms are revised, the older revision may be used until a change in key assignment occurs.
- d) Occasionally, a common key is issued to a Supervisor or a key is needed for only a short period of time. In those instances, the person checking out the key must sign and date when the key is removed and when it is returned using [OSBI CSD QPA 20.1](#).
- e) The drying stall key control log ([OSBI CSD QPA 20.5](#)) may be used when temporarily issuing drying stall keys to individuals outside the OSBI CSD.

2. Issuing Keys, Proximity Access Devices, and Lock Codes

The following individuals are responsible for issuing keys, proximity access devices, and lock codes and maintaining their corresponding logs:

- a) Forensic Science Center (FSC)–
 - i. Identification Badge Proximity Access Devices: Human resources personnel or a designee will issue an identification badge/proximity access device to each CSD employee and assign the employee a personal identification number (PIN). New or modified FSC access privileges should be requested through the supervisory chain to the CSD Director for approval. The CSD Director or designee will then forward approved requests to the FSC Building Manager, who will be responsible for programming the identification badge proximity access device with the authorized level of access to FSC.
 - ii. The FSC Building Manager will issue FSC building entry keys and maintain records documenting the issuance of those keys.
 - iii. Unit Supervisors are responsible for unit keys used for temporary evidence storage areas within their assigned sections.
- b) Regional Laboratories – The Laboratory Supervisor is responsible for tracking keys and access codes.

3. Changing Access Codes/Privileges

At a minimum, cipher lock combinations, access codes or privileges and alarm system codes will be changed or deleted under the following conditions:

- a) When necessary to prevent unauthorized access to a laboratory or unit by a former employee.
- b) When a situation or circumstance involving a potential security breach occurs as determined by the Unit Supervisor or Criminalistics Administrator over the unit.
- c) Employee PIN's should also be changed when necessary due to technical difficulties, such as repeatedly and inadvertently entering duress code.

4. Responsibilities for Updating Codes

Updating access codes for cipher locks and alarm systems will be the responsibility of the following individuals:

- a) Forensic Science Center – The FSC Building Manager will be responsible for controlling the levels of access granted to employees for FSC. Modifications to access privileges should be requested, approved, and routed as indicated in section II.B.2.a above.
- b) Regional Laboratories - Laboratory Supervisor will be responsible for updating access codes.

C. Evidence Storage and Security

1. Each laboratory site shall establish a secure and organized evidence storage room(s) and/or building which must be approved by the CSD Director prior to use. This approval will be documented by a memorandum from the CSD Director to the respective laboratory facility and archived at the FSC.
2. Evidence rooms will be kept neat and clean and will have limited and controlled access. Evidence rooms will have fire and security alarm systems.
3. Each laboratory evidence room or building shall at a minimum meet the following standards:
 - a) An inside room or building with no windows, or, if windowed, the windows must be covered with secure steel bars or grate; or the room must be on an upper floor not easily accessible; or monitored by suitable motion detectors or other devices.
 - b) Doors must have secure locks.
 - c) Sufficient shelving or floor space must be available so that all evidence can be stored in a safe and orderly manner.
4. Prior to being logged in and labeled, evidence may be held in a secure, approved designated temporary evidence holding area or in an Evidence Room. The evidence or the holding area should be labeled to identify it as pending log in. All other evidence placed in an evidence room shall be marked for identification with a Criminalistics case number, the submitting agency, and barcode.
5. No CSD personnel may store evidence at home, in their vehicle, in their office/work area, or at any other such place. This does not apply to latent lifts, impressions, or images of latent prints/impressions which may be maintained by Latent Evidence analysts in their desk areas. CSD personnel may also temporarily maintain custody of evidence while working with the evidence, transporting the evidence, traveling to and from court, or for any other short-term investigative or prosecutorial purpose.

D. Alarm Systems

1. All buildings in which laboratories and evidence rooms are located will have security alarm systems approved by the CSD Director. The security alarm will monitor the facility when it is not occupied. This approval will be documented by a memorandum from the CSD Director to the respective laboratory facility and will be archived at the FSC. Changing and redistribution of access codes/levels will be done as described in sections II.B.3 and II.B.4.
2. Each laboratory facility will maintain a call list for the alarm system. The list is maintained by the person responsible for maintaining keys, magnetic cards, and keypad lock codes in II.B.2 and updated as outlined in II.B.4. Copies of the call list shall be forwarded to the alarm monitoring company in the event the alarm company does not provide the list. The call list will be kept on file with the Regional Laboratory Supervisor in regional laboratories or Administrative Office at the FSC. These same individuals are responsible for archived lists when changes are made.

E. Enforcement

It will be the responsibility of the immediate unit Supervisor to ensure and monitor compliance with this policy.

III. Attachments

[OSBI CSD QPA 20.1, Rev. 1 Temporary Key Control Log](#)

[OSBI CSD QPA 20.2, Rev. 1 Key Control Log](#)

[OSBI CSD QPA 20.3, Rev. 1 Keypad Control Log](#)

[OSBI CSD QPA 20.4, Rev. 1 Master Key Log](#)

[OSBI CSD QPA 20.4b, Rev. 1 Excel Master Key Log](#)

[OSBI CSD QPA 20.5, Rev. 1 Drying Stall Key Control Log](#)

I. Scope

This procedure will outline the process for recommending and evaluating research of new methods or instruments.

II. Procedure**A. Research**

Research may be conducted to develop new methods or evaluate the suitability of non-standard methods. When conducting research, the following steps should be taken.

1. Obtain approval and guidance for purchasing, storing, and using any new reagents. Refer to OSBI Policy 121.1 concerning proper procedures for procuring new reagents.
2. Prepare a written research plan including the following information:
 - a) goal(s) and objective(s) of the research
 - b) description of research project(s) to be performed
 - c) budget
3. Route the research plan through the Research Committee to the CSD Director.
 - . The CSD Director may present the research plan during an administrative staff meeting for discussion if necessary.
5. The CSD Director will notify the Research Committee chairperson if the plan was approved or denied.
6. Following the completion of the research, a summary should be prepared and routed in the same fashion as the research proposal. The summary should indicate whether the research was successful and what further action is recommended.
7. When enough data from research has been obtained to determine that it is desirable to introduce a method for use in casework, the research should be summarized with a recommendation to proceed to validation. A more thorough and detailed validation plan should then be submitted according to section [QP 21.2](#).

III. Attachments

None

I. Scope

All analytical methods utilized by the OSBI CSD will be validated prior to use to ensure each method is fit for its intended use. This procedure explains the process for validating and approving methods used by the CSD.

II. Procedure**A. Proposing and Reporting Validation Studies**

Any new or modified method, or method to be used in a fashion outside the original intended scope, will be validated prior to use in testing.

1. Obtain approval and guidance for purchasing, storing, and using any new reagents. Refer to OSBI Policy 121.1 concerning proper procedures for procuring new reagents.
2. Prepare a validation plan and route the plan in the same fashion as that required for research plans.
3. The plan should include the following sections and/or attachments:
 - a) goal(s) and objective(s)
 - b) financial impact, including:
 - i. a list of additional equipment, reagents, and supplies necessary to complete the validation and projected cost
 - ii. a projected cost/sample, including a comparison to any existing method used
 - c) study descriptions, including:
 - i. purpose/type of study (precision, sensitivity, etc.)
 - ii. validation parameters for testing (e.g. temperature, time, volume, etc.)
 - iii. types of samples (pristine, simulated forensic, non-probative) and approximate numbers of sample
 - iv. controls and any standard reference materials
 - v. method for evaluating data
 - d) references, including:
 - i. any applicable recommendations/requirements from a Scientific Working Group
 - ii. any applicable peer reviewed literature
 - iii. draft protocol

4. Following the completion of the validation study, the results will be summarized for review. Validation reports should include the following sections/attachments:
 - a) range and accuracy of values obtainable (this may include uncertainty of measurement, detection limit/sensitivity, selectivity/specificity, reproducibility, etc.)
 - b) quality control measures which must be incorporated with the method
 - c) final draft of the technical protocol and any quality control/calibration protocols required
 - d) final cost/benefit analysis (this should summarize both the technical and financial benefits and liabilities)

B. Review of Validation Reports

1. The initial review of validation reports will be conducted by the appropriate QIC subcommittee.
2. The subcommittee should ensure that the report is complete and then, based on their review, recommend the method be approved or recommend further validation.
3. Completed validation reports which are recommended for approval should be forwarded to the appropriate CA, the Quality Manager (QM), and the CSD Director. A summary of the subcommittee recommendation, including any dissenting opinion(s), should be attached to the validation report.
4. If the CA, QM, and CSD Director unanimously agree with the subcommittee then the validation will be approved.
5. If the CA, QM, and CSD Director do not unanimously agree, they may return the validation for additional action. Alternately, the Quality Manager may forward the report to all QIC members including all comments and recommendations.
6. The Quality Improvement Committee can then approve the technique or remand the validation for additional study.
7. QIC review and approval may be conducted and documented via e-mail if necessary.
8. The validation plan and all related documentation are to be archived with the Technical Manager of the discipline.

9. Approval of validation summaries may be documented by writing on a copy of the validation summary. It may also be further documented through the issuance of an approved protocol.

C. Performance Checks

A performance check will be conducted to verify that a validated method works as expected on additional or new instruments of the same kind. Performance checks may also be appropriate when other minor modifications are made. Performance checks will be conducted under the instruction of the appropriate Technical Manager. The Technical Manager and/or the appropriate Supervisor will be responsible for maintaining the appropriate documentation.

III. Attachments

None

I. Scope

This procedure will be used for the estimation and documentation of uncertainty of measurement for applicable testing methods.

II. Procedure**A. Tests Which Require Estimation of Uncertainty of Measurement**

The following criteria will be used to determine whether or not an estimation of uncertainty of measurement must be calculated.

1. Estimation of uncertainty of measurement will be calculated for quantitative measurements where numerical values are reported and there is a reasonable expectation that a customer will use the result to determine, prosecute, or defend the type or level of criminal charge.
2. Estimation of uncertainty of measurement is not required for qualitative tests which do not result in numerical values or for quantitative tests where the numerical value obtained is not reported.

B. Steps for Estimating Uncertainty of Measurement

1. Specify what is being measured.
2. Indicate the method for measurement, specifically the equipment or instrument used to take the measurement.
3. List all potential sources of uncertainty in a measurement of uncertainty budget.
4. Review the uncertainty budget and eliminate, if necessary, any potential sources which based on the previous experience of discipline experts, do not significantly impact the uncertainty of measurement.
5. Gather measurement data through an appropriate process. The discipline Technical Manager (TM) will determine the process for gathering sufficient data or will identify previously collected data, such as data from validation studies, proficiency tests, or quality control steps, that will be used for the calculation.
6. The TM will then select the appropriate formula for calculating the uncertainty of measurement. If necessary, the TM will consult a qualified statistician or metrology expert to determine the most appropriate formula.
7. The TM will be responsible for reviewing and updating uncertainty of measurement

calculations. Reviews should be conducted annually at a minimum. Updates should be completed whenever there is a significant change to the uncertainty budget.

C. Documenting Uncertainty Calculations

1. The TM will be responsible for maintaining documentation of any uncertainty of measurement calculations conducted. Documentation must include the data elements listed in section II.B above.
2. Analysts must document in the case record those factors which impact the uncertainty of measurement. This may include the identification of the instrument or measuring device used or an indication of the size or capacity of the measuring device.

D. Reporting Uncertainty of Measurement

1. Reporting uncertainty of measurement will be conducted as follows:
 - a. The uncertainty of measurement will be reported upon request by the customer.
 - b. The uncertainty of measurement should also be reported when the measurement and applicable uncertainty of measurement overlap an established and applicable legal threshold. For example, if the legal threshold for trafficking methamphetamine is 20 grams, the reported measurement is 20.02 grams and the uncertainty of measurement is +/- 0.03 grams, then the uncertainty of measurement should be reported.
2. When the uncertainty of measurement is reported, it must be reported using the same units of measurement and the same number of significant figures as the reported measurement. The associated confidence level of the estimated uncertainty of measurement must also be reported.
3. If the uncertainty of measurement is reported without a customer requesting it, the reporting analyst should ensure that the customer understands what the uncertainty of measurement is. This can be accomplished by calling the customer and explaining the new reporting language prior to or shortly after releasing the report. Alternately, the TM may prepare and have approved by CSD Administration a written explanation of uncertainty of measurement that can be attached as an appendix to reports or e-mailed to customers as necessary.

III. Attachments

None

I. Scope

The OSBI CSD will use the following procedure to ensure adequate traceability of measurements as applicable and required by quality procedures for calibration of equipment, reference standards and reference materials.

II. Procedure

A. Internal Calibrations

In the event OSBI CSD personnel perform internal calibrations or comparisons to establish traceability, the appropriate TM shall develop and issue a written procedure which complies with the current Measurement Traceability Policy issued by ASCLD/LAB. A list of current ASCLD/LAB documents and policies is located at http://www.ascl-d-lab.org/programs/intl_testing_overview.html.

B. External Calibrations

In order to ensure that external calibrations of measuring equipment/instruments, reference standards or reference materials provide adequate traceability, the following steps will be used to select an appropriate vendor and maintain adequate documentation of the calibration.

1. The vendor conducting the calibration must demonstrate and provide documentation of **competence** and adequate **measurement capability** and **traceability**.
 - a. **Competence** will be demonstrated by selecting a calibration laboratory which is ISO/IEC 17025:2005 accredited. The accreditation must be from an IAAC or ILAC MRA signatory or another verified through ASCLD/LAB as an adequate accrediting body. Listings of current IAAC MRA and ILAC MRA signatories may be found on the internet at <http://www.iaac.org.mx/English/Members.php> and http://www.ilac.org/documents/mra_signatories.pdf respectively.
 - b. **Measurement capability** should be verified by determining that the calibration laboratory's scope of accreditation includes a measurement range applicable to the instrument or measuring equipment being calibrated.
 - c. **Measurement traceability** should be verified by obtaining documentation from the calibration laboratory which documents traceability to NIST standards.
2. The calibration certificates received from outside calibration laboratories should include:

- a. the measurement results and the associated uncertainty of measurement, and/or
 - b. a statement of compliance with a metrological specification.
3. Instruments, equipment, reference standards, and reference materials which have been externally calibrated to obtain appropriate traceability will not be adjusted by OSBI CSD personnel. If intermediate checks are performed and results indicate that the item is no longer within the calibrated range, the item will be taken out of service until an appropriate external calibration can be performed.

C. Calibrations to Alternate Reference Standards or Materials

If calibrations cannot be made in SI units, the OSBI CSD will provide confidence in reported measurements by:

1. establishing traceability to a certified reference material (see [QP 26](#)), or
2. using a specified method or consensus standard agreed on by OSBI CSD management and the customer.

III. Attachments

None

I. Scope

The OSBI CSD will furnish CSD facilities with equipment and instrumentation which will provide the correct performance for the analysis conducted. This procedure will be used to track, maintain, and verify calibration of equipment and instruments. This procedure will also be used to ensure safe handling, transport, storage, use, and maintenance of measuring equipment.

II. Procedure**A. Inventory**

1. The Supervisor of each unit and of each regional laboratory must have access to an inventory of their instruments/equipment and analysis software.
2. The inventory for instruments/equipment should include the following information:
 - a) Name
 - b) Manufacturer, model number, and serial number
 - c) OSBI asset number
3. As each unit or laboratory receives new instruments/equipment, the Supervisor will send information concerning the new instruments/equipment to the FSC Administrative Office.
4. A file containing original paperwork (manufacturer's information, etc.) for instruments/equipment should be readily accessible by the Supervisor.

B. Calibration and Maintenance Procedures

1. Equipment procedures (protocols) will be established as necessary to ensure the following criteria are met:
 - a. Calibration programs or procedures will be established for key values of instruments where those properties have a significant impact on the accuracy or validity of test results.
 - b. Equipment will be checked (e.g. function verification) or calibrated before being placed into service to ensure it meets the necessary specifications established in OSBI CSD protocols.

- c. Equipment used for tests which has a significant impact on accuracy or validity of tests results shall be calibrated before being put into use.
 - d. For measuring equipment (or test equipment with measuring functions) that has a significant impact on the accuracy or validity of a reported test result, the calibration procedure must ensure that any calibrations and measurements made are traceable according to [QP 23](#). This requirement does not apply if it is documented that any error contribution from the calibration has a negligible impact on the total uncertainty of measurement.
 - e. Calibration or calibration checks will be conducted as needed for measuring equipment (or test equipment with measuring functions) for which calibration does not significantly contribute to the total uncertainty. The calibration or calibration check should be adequate to ensure that the equipment or instrumentation can provide the necessary or appropriate level of uncertainty of measurement.
2. Equipment procedures will require and specify the maintenance of the following records for equipment and software which has a significant effect on the accuracy or validity of test results:
 - a. **user’s manuals** or manufacturer’s instructions or a reference to the location of the manuals/instructions;
 - b. **calibration records** which include the dates, results, and copies of calibration reports/certificates, as well as records of any adjustments, acceptance criteria, and the due date for the next calibration;
 - c. **results of function verifications** (at a minimum the record should include the date of the verification, initials of the person conducting the verification, a description of the verification activity, and identification of any reference standards or materials used);
 - d. **maintenance plan and records** of maintenance (at a minimum maintenance records should include the date of the maintenance, initials of the person conducting the maintenance, and a description of the maintenance activity);
 - e. **records of any damage, malfunction, modification, or repair.**
 3. Maintenance procedures should include a maintenance plan, maintenance contract information, and routine preventive maintenance.
 4. Whenever an applicable laboratory instrument/equipment is taken out of service, an entry will be made in the logbook including the “out of service” date and again the “in service” date. Additionally, the instrument/equipment must have an “out of service” sign placed on it or be otherwise isolated and/or identified to prevent use until it is placed back into service.

C. Measuring Equipment

1. At a minimum, measuring equipment, such as pipettors, balances, pH meters, etc., will be handled, transported, stored, used and maintained according to manufacturer's recommendations in order to prevent deterioration and/or contamination. If additional procedures are necessary to adequately prevent deterioration and/or contamination, the discipline Technical Manager will issue a protocol detailing the appropriate additional steps.
2. The manufacturer's user's manual should be maintained for all measuring equipment. In the event that the manufacturer's user's manual is not available, the discipline Technical Manager will be responsible for issuing a protocol detailing proper handling, transportation, storage, use, and maintenance of the equipment.

III. Attachments

None

I. Scope

This procedure will be followed to ensure traceability of reported measurements, when applicable, and to ensure proper calibration and handling of reference standards.

II. Procedure**A. Calibration of Reference Standards**

1. Reference standards will be calibrated in a fashion that provides traceability as described in [QP 23](#), unless the contribution to uncertainty of measurement associated with the measuring or test equipment is negligible.
2. Calibration Requirements
 - a) Reference standards used to calibrate instrumentation or to provide traceability for measurements (weight, length, temperature, etc.) that are reported must be properly calibrated.
 - b) Reference standards shall be calibrated in a manner that ensures compliance with [QP 23](#).
 - c) Calibration certificates will be maintained at the CSD unit using the reference standard. The original calibration certificates for reference standards used by multiple units will be maintained with the standards. Units using the standards should maintain a copy of the calibration certificate for easy reference. Expired certificates will be maintained by the appropriate unit Supervisor or Technical Manager.

B. Transport and Storage

1. Calibrated reference standards will be stored separate from evidence in a secure location that does not invalidate their performance as reference standards.
2. The current location of calibrated reference standards used by multiple units will be tracked through the use of a sign out sheet or equivalent tracking mechanism.
3. Calibrated reference standards will be used for their intended purpose only, unless it can be demonstrated that alternate uses do not invalidate their performance as reference standards.
4. Manufacturers' recommendations will be followed for the handling, storage, transport, and use of reference standards to ensure that they are protected from damage.

III. Attachments

None

I. Scope

This procedure outlines the legal, safety, transportation, storage, handling, and use requirements for reference materials, such as drug standards and certified reference materials (such as NIST Standard Reference Materials or SRM's). These requirements will be followed to prevent contamination or deterioration of reference materials and in order to protect their integrity.

II. Procedure**A. General Requirements**

When it is not possible or appropriate to trace reported results to SI units, the OSBI CSD will ensure the reliability of reported results, wherever practicable, through the use of certified reference materials.

1. Certificates of analysis from manufacturer's will be maintained in a location designated by the Supervisor and/or Technical Manager.
2. Reference materials will be labeled with the date of receipt and receiving analyst's initials, date opened, initials of the individual opening the reference material, and expiration date, if applicable.
3. Reference materials should not be stored with evidence samples.
4. The safe handling, transport, storage and use of reference materials will be conducted according to the manufacturer's instructions or an approved discipline protocol to prevent contamination, deterioration and to protect integrity.
5. Whenever possible, only one lot number of standards will be open and in use at a time.

B. Drug Standards

1. Drug standards are considered reference materials.
2. Drug standards will be received, handled, and logged according to applicable discipline Quality Manuals.

III. Attachments

None

I. Scope:

This procedure provides for assistance by criminalists at crime scenes. This procedure will be followed to ensure proper notification of crime scene requests and responses and documentation of activities conducted at a crime scene.

II. PROCEDURE:**A. Requests for Crime Scene Assistance:**

The OSBI CSD provides crime scene assistance through two groups, the Latent Evidence Unit (LEU) and the Crime Scene Response Team. Requests for assistance by a Criminalist in one of these groups will be routed as indicated below.

1. Latent Evidence Unit

- a. The supervisor or designee will be notified of the request and determine if the scene warrants the use of a criminalist.
- b. In the event that the supervisor or designee is unavailable, the Criminalistics Administrator (CA) responsible for LEU will be contacted and make assignment as needed.
- c. The Division Director should be notified (via phone, blackberry, e-mail, etc.) when criminalists attend a crime scene.

2. Crime Scene Response Team

Participation in the Crime Scene Response Team is voluntary. Individuals interested in participating may express their interest to the CSD Director who will make the final determination whether an individual is eligible to participate.

- a. Individuals seeking assistance from the Crime Scene Response Team will contact the CSD Director who will notify the appropriate individual(s) from the Team.
- b. If the CSD Director is not available, the CA assigned to FSC Biology, Firearms, Trace, et al, will be contacted.

B. Crime Scene Memorandum:

1. All crime scene responses require a Crime Scene Memorandum ([OSBI CSD QPA 27.1](#)).
2. Copies of the crime scene memorandum will be distributed as follows:

- a. One copy each will be forwarded to the CSD Director and the Evidence Administrative Programs Officer. At the latest, this copy should be turned in by 10:00 A.M. on the following Monday.
 - b. The original will be placed in the criminalistics case record, if applicable.
 - c. A copy may be retained by the responding criminalist(s) and/or supervisor.
3. If the responding OSBI Criminalistics personnel did not leave together, travel together in the same unit and return together, separate crime scene memos must be completed.
 4. Separate crime scene memos will be filled out when a Criminalist responds to different locations (scenes) on the same case.

C. Crime Scene Narrative Report:

1. LEU criminalists collecting, preserving, diagramming, photographing, or analyzing evidence at a crime scene will complete a Crime Scene Narrative Report . Morgue calls do not require a crime scene narrative.
2. Crime scene response team members are provided for support only. While they may assist with photographing or diagramming the scene, the report regarding crime scene activities will remain the responsibility of the investigative personnel on scene.
3. The narrative report will include a general description of the Criminalist's activities at the crime scene, an itemized list of what was inventoried or collected, and if testing was performed at the scene. The narrative report can be included with the Criminalistics Examination Report and issued as one document. A copy of a Crime Scene Narrative Report can be distributed to district attorneys, investigating officers and other appropriate parties. The original Crime Scene Narrative will be retained in the case file.

III. Attachments

[OSBI CSD QPA 27.1, Rev. 0](#)

I. Scope

This procedure establishes consistency in the format, content, style, and distribution of analytical reports.

II. Procedure

A. Generating Analytical Reports

1. OSBI CSD analytical reports will be generated in the BEAST Laboratory Information Management System (LIMS). Although the procedure may vary some from discipline to discipline, this is typically done in the following manner:
 - a. The analyst will click on the “Analysis” button from the “Assignments” tab in the BEAST.
 - b. This should open exam log and/or matrix panels that the analyst can use to document the analysis.
 - c. Once the necessary data has been entered, clicking on the “Send to Word” button should open a draft report. Analysts should verify that the information in the report is complete and properly formatted.
2. When a report has been approved, **but not distributed (hard copy or pdf via website)**, it may be corrected with assistance from a LIMS administrator using the following **reset** process:
 - a. Reset the report – this creates a new assignment and the matrix data is still in place.
 - b. Delete the reset report – this deletes the current report number along with the report.
 - c. From the matrix, regenerate the report. The analyst can then proceed to sign, route for review, and approve the report.
3. When a report has been distributed and needs to be corrected, the following process will be used to issue an **amended report**:
 - a. Create an assignment with the same items selected for the assignment that were included in the report being corrected.
 - b. In the lower right corner of the “Assignments” tab, indicate the report number that is being corrected in the “Amended From” field.

- c. Click on the “Analysis” button, navigate to the proper “Send to Word” button, and create a report. All of the exam log and matrix panels should be blank, so the report generated should not include any results.
 - d. Copy and paste the necessary content from the report being amended into the new report document. Do not copy the signature block.
 - e. Make the necessary corrections to the content of the report, identify the report as an amended report according to section II.B below, and then conduct review and approval of the report according to applicable procedures.
4. Occasionally, case circumstances will change and requested analysis will no longer be necessary. When this happens and notification is received before the analysis begins, the following procedure will be used to issue a **“no-analysis” report**.
- a. First, enter a narrative in the BEAST case file.
 - i. From the “Case Info” tab, click on the “Narrative” button.
 - ii. Click on the “Add” button in the lower left corner. Then enter details of the conversation, including the first and last name of person, the agency, and what analysis is no longer needed.
 - iii. Select “NA” for no analysis communication in the “Type” field.
 - iv. When all information has been entered, click “save.”
 - b. Next, generate the no analysis report.
 - i. From the “Assignments” tab, highlight the appropriate assignment, and click the “Edit” button at the bottom of the screen.
 - ii. In the “Lab/Format” field in the lower left corner of the screen, select “NOAN” and, if necessary, select the appropriate analyst in the “Analyst” field. Click “Save.”
 - iii. Click on the “Notes” button and enter the amount of time spent on the “no analysis.”
 - iv. Click on the “Analysis” button and navigate through the exam log/matrix panels and click on the appropriate “Send to Word” button. Verify the information is correct, and then save the report. Print a sufficient number of reports for mailing, if necessary.
 - c. Finally, close the assignment.
 - i. Back on the “Assignments” tab, click on the “No Analysis” button.

- ii. Click on “Yes” in the next window.
 - d. The analyst closing the assignment is responsible for ensuring that the evidence is moved or routed appropriately so that it can be returned to the proper agency in a timely fashion.
5. In the event that a request for analysis is cancelled after analysis has started but before it has been completed, a **partial analytical report** will be issued. At a minimum the report should address the partial results in one of the following two ways:
 - a. The report can be issued indicating that analysis was started but then cancelled at the request of the customer or due to case circumstances (suspect pled guilty, DA declined to file charges, etc). The report should include the name (first and last) of the customer who cancelled the service request or confirmed that analysis was no longer necessary. The report may then indicate that results of testing are not reported since the analysis was not completed. This type of report must be administratively reviewed, but no technical review is required.
 - b. Alternately, the report may include results from any testing/items which were completed and a statement similar to that described above to address any testing which was not completed.
6. When evidence is sent to a contract laboratory for analysis, an OSBI report is not required. However, the requesting agency should be notified in writing (e.g. report, letter, or memo) that the evidence was sent to a private lab for analysis. At a minimum, this written notification should include the requesting agency case number, OSBI Lab case number, a description of the evidence and/or packages sent, and the name of the laboratory the evidence was submitted to.

B. Numbering Reports

1. Reports and notifications generated in the BEAST will be assigned a sequential number by the BEAST. This number will not be modified by analysts.
2. On amended reports, the analyst will indicate below the report number, “Amended From Report X” where X is the number of the report that was amended.
3. When issuing an amended report for a report that was generated prior to the implementation of the BEAST, the analyst will indicate in the report what report was amended. If the report does not have a number, the issuing analyst and date of issue will be referenced for identification purposes.

4. Similarly, if a case has been worked both before and after the implementation of the BEAST and a duplicate report number is assigned by the BEAST, it is recommended that the analyst place a statement of explanation in the report.

C. Report Content

The format of analytical reports will include the following information. The following requirements do not apply to reports or notifications which do not contain results of analysis. This includes no-analysis reports and CODIS Database Entry Notifications.

1. All OSBI CSD analytical reports will have the title, “Criminalistics Examination Report.”
2. All OSBI CSD analytical reports will include the date the report is issued.
3. The name and address of the OSBI Laboratory issuing the report will be reflected in the header of the report. If part or all of the analysis is conducted at another location, this shall be documented in the case record.
4. All OSBI CSD analytical reports will contain and be uniquely identified by the case number and a report number.
5. The name and agency and/or address of the requesting officer will be located in the report header.
6. All OSBI CSD reports will reflect the date evidence was first received by the CSD. For evidence that has left the custody of the OSBI CSD and then been resubmitted, the first date that the evidence was received back into CSD custody will be reflected in the report.
7. All OSBI CSD reports will include a description of the evidence submitted and an unambiguous identification of the items tested.
8. All current OSBI CSD reports, with the exception of no-analysis reports, shall include a signature block showing the type-written name and title of the analyst issuing the report and an electronic signature.
9. Requesting agency case numbers should be listed on the first page of the report immediately below the header. Be sure to indicate the agency number by using the initials or title of the agency with the agency case number.
10. Analysis of evidence or results sections will include identification of the item tested and test results.

11. Where relevant, OSBI CSD reports should include a statement indicating that the reported results apply only to the items tested.
12. All DNA reports will include qualitative or quantitative interpretive statements; the amplification system or loci used for analysis; and a disposition of the evidence, as applicable.
13. When associations are made, the significance of association shall be communicated clearly and properly qualified in the report.
14. When comparisons are made and result in an elimination, the elimination shall be clearly stated in the report.
15. When results are inconclusive, the report shall clearly communicate why no definitive result can be made.

D. Analytical Report Format

1. All OSBI CSD reports will be numbered to indicate the page number and total number of pages in the report.
2. OSBI CSD reports will follow proper rules of grammar and correct spelling in the Criminalistics Examination Report.
3. For DNA reports, the method of analysis used will be included in the report.
4. All Criminalistics Examination Reports will be in Times New Roman font. The preferable font is size 10 or 12. Only the certification block and any report footer should have a font size of less than 10 (with a minimum font size of 7).
5. All OSBI CSD reports will reflect the results of analysis, including units of measurement, where appropriate.
6. The signature block will be located on the right side at the bottom of the report.
7. The report certification block will be placed next to or above the signature block.
8. Opinions and interpretations reported will be appropriately identified in the report.
9. Multiple submittals should be listed in a paragraph format immediately preceding the description of those items of evidence.
10. With the exception of the Drug and Toxicology reports, the "Analysis of

Evidence" should be written in semi-narrative style. Item numbers and analytical results are included under this section.

III. Attachments

None

I. Scope

Criminalists will accurately document the number of items submitted, the number of examinations performed, and the time expended analyzing items of physical evidence in each sub-case worked .

II. Procedure

- A. Criminalistics statistics will be recorded by clicking the “Notes” button on the “Assignments” tab in the BEAST. Criminalists may, at their discretion, maintain a personal log book.
- B. Each Criminalist who inventories and/or analyzes items in a sub-case will record the appropriate numbers in the BEAST.
- C. It is the responsibility of each Criminalist to accurately record their statistics at the conclusion of the analysis.
- D. Worksheets can be utilized to document the kinds and numbers of examinations made on all items of a sub-case analyzed. Worksheets will be retained in accordance with [QP 16.1](#) and [QP 16.2](#).
- E. It is acknowledged that some analytical procedures utilize reference standards, controls, or both and these shall be counted as examinations as well.
- F. It is not the intent of this policy to mandate standard analytical procedures for Criminalists to follow for the analysis of specific kinds of items. However, each Criminalist shall perform sufficient analytical procedures to support the opinion rendered by the report. The following procedures are not all-inclusive. Any additional procedures to be used for statistical purposes must be approved by the appropriate Criminalistics Administrator.
- G. ANALYTICAL PROCEDURES: Any of the following listed analytical procedures performed on an item will be counted as ONE examination . Items inventoried for report purposes only (no analysis cases) will not be counted as items analyzed or counted as examinations.
 - 1. Drug analysis
 - a. Any measurement to determine quantity of an item (i.e., weight, volume, count).
 - b. Macroscopic or microscopic examination for a specific purpose.
 - c. Reference/literature search.
 - d. Spot/color test (includes Duquenois-Levine test).
 - e. pH determination.
 - f. Thin layer chromatography examination.

- g. Wet chemistry extraction.
 - h. Individual instrument analysis.
 - i. Determination of a quantitation curve/quantitative analysis.
2. Toxicology
- a. Alcohol (Volatile) Analysis.
 - b. Immunoassays – Immunoassays for each individual drug assay will each be considered one examination.
 - c. Blood Drug Screen – Each screen for different classifications of drugs (bases, acid – neutral) will be considered one examination.
 - d. Quantitations for Blood Drug Concentrations – Quantitations which require separate extractions due to drug classifications will each be considered one examination.
 - e. Reference/literature search.
3. Forensic Biology
- a. Serology
 - 1) Macroscopic examination (including use of alternate light source) or microscopic examination of a single item for a specific purpose.
 - 2) Collection, preservation, and labeling of a single sample removed from an item.
 - 3) Any screening examination or presumptive test performed on a single sample.
 - b. Hair Analysis
 - 1) Slide preparation for individual unknown or known sample.
 - 2) Any measurement of individual unknown hair.
 - 3) Determination of racial origin of a single unknown hair.
 - 4) Determination of body area of origin for a single unknown hair.
 - 5) Comparison of single unknown hair to a known reference sample.
 - 6) Preparation of a scale case.
 - 7) Assessment of a single hair root for suitability for nuclear DNA profiling.
 - 8) Assessment of a single hair shaft for suitability for mtDNA sequencing.
 - 9) Reference/literature comparison.
 - c. DNA STR Analysis
 - 1) Visual evaluation of the biological material.
 - 2) Isolation/extraction of DNA from biological material.
 - 3) Real Time PCR analysis for quantitation of human DNA in a sample.

- 4) Amplification setup and run in the thermal cycler.
- 5) ABI 310 or 3130 setup and run per injection sample.
- 6) Application of GeneMapper ID (GMID) software and evaluation of raw data for each injection of each sample.
- 7) Application of GMID software and assessment of quality of data for each injection of each sample.
- 8) Profile assessment/comparison for each genetic locus compared for each sample .
- 9) Composite population calculation entire “OSBI STATS” calculation per sample.
- 10) Evaluation of a previously collected sample for feasibility of testing.
- 11) Division, packaging and labeling of a previously collected sample for independent testing and/or retention in the laboratory.
- 12) Reference/Literature Search.

4. Firearms & Toolmark

- a. Visual examination.
- b. Determination of class characteristics.
- c. Determination of individual characteristics.
- d. Comparison of individual known samples to individual unknown sample.
- e. Firearm function examination.
- f. Reference/literature search.
- g. Any distance determination procedure.
- h. Chemical procedure to identify obliterated serial number.
- i. Shot cup or wad examination.
- j. Comparison of cutting or pry tool to individual cut or pry mark.
- k. Comparison of any two individual fractured items.

5. Latent Print

- a. Macroscopic or microscopic examination.
- b. Standard dusting powder examination.
- c. Forensic (fluorescence) dusting powder examination.
- d. Light source examination.
- e. Superglue fuming examination development.
- f. Ninhydrin examination development.
- g. Dye staining examination development.
- h. Image enhancement.
- i. Sudan black B examination.
- j. Zinc chloride examination.
- k. DFO examination.
- l. Side/back lighting examination.
- m. UV light examination.

- n. Microscopic enhancement.
 - o. Electrostatic dust lifter.
 - p. Small Particle Reagent (SPR)
 - q. Attempt to obtain viable impression from morgue sample.
 - r. Comparison of unknown latent impression to any known impression.
 - s. Reference/Literature Search.
 - t. Casts
 - 1) Visual examination for comparison suitability.
 - 2) Comparison to physical items for class characteristics.
 - 3) Comparison to physical items for individual characteristics.
 - 4) Reference/Literature Search
6. Trace Evidence
- a. Sample preparation.
 - b. Macroscopic or microscopic examinations for a specific purpose.
 - c. Specialized light examination (either macroscopic or microscopic).
 - d. Spot/color test.
 - e. Microcrystal examination.
 - f. Thin layer chromatography examination.
 - g. Melting point determination.
 - h. Solubility determination.
 - i. Density determination or comparison.
 - j. Refractive index determination or comparison.
 - k. Individual instrument analysis.
 - l. Reference/literature search or comparison.
 - m. Any physical measurement or comparison.
 - n. Arson
 - 1) Macroscopic or microscopic examination for a specific purpose.
 - 2) Sample preparation or isolation.
 - 3) Burn tests.
 - 4) Individual instrument analysis.
 - 5) Reference/Literature Search.

III. Attachments

None

I. Scope

Proficiency tests will be used to monitor the quality of results provided by OSBI CSD analysts. This procedure will be used to conduct external and internal proficiency testing, including re-analysis of casework samples.

II. Procedure**A. Scheduling Proficiency Tests**

Each year, the Quality Manager (QM) or designee will coordinate with Technical Managers to prepare a draft proficiency test schedule that meets the following guidelines:

1. If possible, each analyst working in a discipline containing multiple categories of testing, will be administered a proficiency test, annually, for each category of testing in which he/she performs analysis. At a minimum, each analyst and technician performing analysis shall complete a proficiency test for each category of testing in which he/she performs analysis once per five year accreditation cycle.
2. Each year, every OSBI laboratory (FSC, NERL, NWRL, SWRL, ERL) shall complete at least one external proficiency test per discipline in which they provide service.
3. When practical, a minimum of one external proficiency test should be completed per category of testing performed per laboratory. Some categories of testing, such as GSR or general Trace analysis, may be more effectively addressed through internal proficiency tests, such as re-analysis casework.
4. Each analyst and technician performing analysis shall complete one proficiency test (internal or external) per discipline in which he/she performs casework, annually.
5. Each Forensic Biologist conducting DNA analysis must complete two external proficiency tests per year. One proficiency test must be completed in the first six months of the calendar year and the second in the last six months of the calendar year. The time between tests must be at least four months, but not longer than 8 months. For calculating time periods between tests, the vendor due date will be used.
6. Forensic Biologists who are qualified for multiple DNA methods or technologies must complete proficiency tests as follows:

- a) Forensic Biologists must complete a test on each method at least once per year. Examples of different methods are Identifiler Plus for casework, Identifiler for database samples, manual extraction or setup, automated extraction or setup, etc.
- b) Forensic Biologists must complete at least two tests approximately 6 months apart for each technology they are qualified to use for casework or database samples. Examples of different technologies are STR analysis, Y-STR analysis, and mitochondrial DNA analysis.
- c) Results from different technologies, but not different methods, may be reported on the same test. For example, Identifiler Plus casework results and Y-STR analysis may be reported on the same test.

B. Ordering External Proficiency Tests

1. The QM or designee will then prepare an internal purchase request to order a sufficient number of external tests as indicated by the schedule.
2. External proficiency tests will be obtained from ASCLD/LAB approved test providers, where available.

C. Assigning External Proficiency Tests

1. Upon receipt of test samples from an external provider, the QM or designee will determine which employee each test will be assigned to and what analysis the analyst will need to conduct. The QM will also retain the “Release of Data to Accreditation Bodies” form for proficiency tests from Collaborative Testing Services.
2. The QM or designee will then log the proficiency tests into the BEAST as follows:
 - a) Open the BEAST receive.exe program.
 - b) From the main screen click on the “New Lab Case” button.
 - c) Click on the manual submission button on the screen that comes up. This icon looks like a hand on top of a piece of paper.
 - d) Click OK on the screen that opens next.
 - e) Enter PT as the case type, 00 as the county, and 3 as the priority.
 - f) Enter the appropriate test provider in the “Department” field.
 - g) Enter “Quality Manager” in the case officer field.
 - h) Enter “UPS” or the most appropriate submission type in the “Submission Type” field.
 - i) Enter the UPS or other tracking number from the box or package that the

- proficiency tests were received in. This can also be done by scanning the barcode on the shipping label.
- j) Enter the proficiency test number, including an abbreviation for the provider and the letter designator for the participant code, in the “Department Case” field.
 - k) Enter a name in the “Submitted By” field. For example, C.T. Services can be used as the “Submitted By” name for a CTS proficiency test.
 - l) Enter the date that the proficiency tests were received as the “offense date.”
 - m) On the “Names” tab in the Quick Create screen, enter names based on the proficiency test scenario.
 - n) On the “Containers” tab, enter the appropriate container designator, package type, item number(s) and service request(s). Refer to QP 5 if necessary.
 - o) On the “Items” tab, enter a description of what was submitted in the “Description” column.
 - p) Complete the case creation process by clicking the “Quick Create” button at the bottom of the screen.
3. After the case has been logged in, the QM or designee will attach the evidence barcode generated to the sample packet or container and the file folder barcode will be attached to the proficiency test data sheets.
 4. Next, the QM or designees will assign the proficiency test “case” to the appropriate analyst.
 - a) From the main BEAST screen, click on the “Assignments” button at the top of the screen.
 - b) Use the search tab to bring up the list of recently created proficiency test cases. This can be done by searching on a Case Type of “PT” or a Priority of “3.”
 - c) From the list of cases that comes up, highlight the appropriate case and click on the “Assign” button in the lower left corner.
 - d) Select the appropriate analyst, priority, and status.
 - e) Enter in the “Comments” field when the test must be returned to the QM and, if necessary, any specific analysis instructions and click “OK.”
 - f) Repeat this process until all proficiency tests are assigned.
 5. Once all proficiency tests have been logged in and labeled with barcodes, the QM or designee will scan the barcodes for the case files to the FSC File Room location. The data sheets will be provided to the appropriate analyst and no further custody transactions will be tracked for the data sheets.
 6. The sample packs will be transferred to the custody of the appropriate analyst or transferred to/routed to the appropriate property room location.

D. Preparing Internal Proficiency Tests

Internal proficiency tests may consist of re-usable, expired external proficiency tests, external proficiency tests which can be analyzed multiple times, internally prepared proficiency tests, or re-analysis casework.

1. Internal proficiency tests may be prepared from re-used external proficiency tests by following these steps:
 - a) The samples will first be packaged in a fashion which precludes associating the samples with the original manufacturer test number.
 - b) The samples will be assigned a proficiency test number in the following format IPT-QM-YY-NNN, where YY is a two digit representation of the year the proficiency test will be assigned, and NNN is a chronological number assigned to the test.
 - c) The test number and the expected results associated with the test must be recorded.
2. An external proficiency test may be assigned to one or more analyst(s) as an internal proficiency test when the test samples lend themselves to multiple examinations (e.g. comparison of latent print images ordered on a DVD) . In these situations, the test will be assigned in the same fashion as an external proficiency test and must be completed prior to the release of the manufacturer’s results. The department case number for an external test that will not be submitted to the manufacturer will include a suffix with “IPT” and a letter to designate the participant. (e.g. CTS11-518-IPT-A)
3. Internal proficiency tests prepared “from scratch” in-house will be prepared by the appropriate Technical Manager using the following guidelines:
 - a) The composition of the proficiency test will be determined based on the type of procedure and samples tested.
 - b) All proficiency tests should be prepared using samples, materials, and methods that ensure the uniformity, integrity, and identity of the proficiency testing samples.
 - c) Duplicate testing samples should be prepared and retained when possible. Alternately, test samples should be prepared in a fashion that will facilitate re-testing, in the event of a potential discrepancy.
 - d) Proficiency test samples that use comparisons or produce qualitative results should be prepared in such a way that they contain sufficient class/individual characteristics for meaningful analysis and comparison.
 - e) For those proficiency tests that evaluate procedures which produce quantitative results, samples should contain an amount of testing material sufficient to enable a conclusion to be drawn from the results of the analysis.

- f) Proficiency tests should include appropriate controls among the samples submitted, where appropriate or necessary. Standard reference materials may be used as part of the control system if available for a particular examination.
 - g) Each set of proficiency samples must be labeled with a test set identifier using the format IPT-TMD-YY-NNN, where TMD is a designator for the discipline or Technical Manager (e.g. CDS, TM's initials, etc.), YY is a two digit abbreviation of the year the test will be assigned/created, and NNN is a chronological number assigned to the test.
 - h) After analysis, any remaining portion of each proficiency test sample will be returned to the QM or designee for possible reassignment and/or re-analysis and comparison if circumstances dictate.
4. Cases may be selected for re-analysis at the discretion of the Technical Manager, following the guidelines set forth in section II.J, below.

E. Assigning Internal Proficiency Tests

1. Re-used external proficiency tests will be assigned by the QM or designee, using the procedure outlined in section II.C, with the following exceptions:
 - a) II.C.2.f - In the department field, enter IPT to indicate the case is an internal proficiency test.
 - b) II.C.2.h and i – For the submission method, choose hand delivered no signature.
 - c) II.C.2.j – Enter the internal proficiency test case number.
 - d) II.C.3 – The barcode for the case file may be placed on a hard copy of the RFLE and routed to the appropriate analyst.
2. Internal proficiency tests created in-house will be assigned by the appropriate TM or designee, using the procedure outlined in section II.C, with the same exceptions listed in II.E.1. TM's or a designee must provide the QM a record of the test number and the expected results at the time the test is assigned to an analyst.

F. Analysis of Proficiency Tests

1. Proficiency tests will be analyzed using the same analytical protocols as casework. This includes all verifications and administrative and technical reviews required by current OSBI policy.
2. All proficiency tests will be handled in compliance with the OSBI CSD QM and QP's. This includes policies regarding itemization and evidence handling.
3. Proficiency tests will be analyzed to the full capability of the analyst assigned. Each analyst will conduct all appropriate examinations for which he/she is qualified based on the case information provided with the test.

4. The same examination documentation generated during testing will be generated during the analysis of proficiency tests.
5. Results of proficiency tests will be reported to the QM in an OSBI report or according to the normal protocol documentation for any discipline which does not issue a standard OSBI report (e.g. database units).
6. External proficiency test data sheets will be completed by the analyst in their entirety. This includes any section which requires the analyst to list a narrative statement of how their conclusions would be reported. It is not acceptable to put “See Attached Report,” since only the data sheets are submitted to the vendor.
7. Upon completion of the proficiency test, the analyst will turn in all hard copy documentation to the QM.
8. If all proficiency test documentation is electronic, the analyst will e-mail the QM to notify him/her that the test is complete.
9. Remaining samples will also be returned to the QM, or they may be archived according to the unit’s standard procedures for database proficiency samples. When multiple analysts in the same lab or unit have participated in the same proficiency test, all sample packets from the unit/lab should be returned at one time to the QM.

G. Review of Proficiency Tests

1. Upon receipt of a completed test, the QM will review the following documentation to determine if the expected results were obtained.
 - a. External tests – manufacturer’s results and consensus results (once received)
 - b. Re-used external tests – manufacturer’s results and consensus results
 - c. Internal tests created in-house – expected results provided by TM
 - d. Re-analysis cases – memo from TM indicating whether second results were consistent with original analysis.
2. Provided that results are consistent, the QM will mark the results page/memo as satisfactory, initial and date the page and forward the page to the analyst, any other participants, and the TM (if necessary) for notification. Each individual will initial or sign and date the page to document his/her notification and return the page to the QM for archival.
3. Any test which does not yield consistent results will be forwarded to the appropriate QIC sub-committee for a detailed review and cause analysis according to [QP 13](#).

4. When necessary or practical, notifications of analysts or TM's may be done via e-mail or an alternate method, provided that adequate documentation is maintained to demonstrate that the notification has been received.

H. Review of DNA Proficiency Tests

1. For DNA proficiency tests, the QM will route the results page(s) to the Biology TM and the following information will be noted by the TM:
 - a. Are results satisfactory or unsatisfactory?
 - b. Are all reported inclusions correct (if applicable)?
 - c. Are all reported exclusions correct (if applicable)?
 - d. Are all reported genotypes and/or phenotypes correct or incorrect according to consensus results or within the laboratory's interpretation guidelines?
 - e. Are results reported as inconclusive or not interpretable consistent with written laboratory guidelines?
2. For any DNA proficiency test identified with non-administrative discrepancies that affect the typing results and/or conclusions, the TM will inform the appropriate CODIS Administrator at the time of discovery. The CODIS Administrator will sign/initial and date the pages to indicate he/she has been notified.
3. The TM will sign/initial and date the page(s) and then forward them to the appropriate analyst and any other participant for him/her to sign/initial and date to indicate he/she received notification of test results.
4. Any test which does not yield consistent results will be forwarded to the Biology sub-committee for a detailed review and cause analysis according to [QP 13](#).
5. When necessary or practical, notifications of analysts, other participants, TM's, or the CODIS Administrator may be done via e-mail or an alternate method, provided that adequate documentation is maintained to demonstrate that the notification has been received.

I. Documentation of Proficiency Tests

The following documentation should be maintained for each test completed:

1. Test set identifier
2. How the samples were obtained (test provider for external) or created
3. Identity of analyst and other participant(s), if applicable
4. Date of analysis
5. Date of completion

6. All data sheets, notes, and work products (scans, electropherograms, etc.)
7. Report of results
8. Provider's proficiency test evaluation, for external tests
9. Quality Improvement Committee (QIC) sub-committee review of any discrepancies noted, if applicable
10. Notice to the analyst/participant of the proficiency test results and review
11. Notification to TM of results and review, for DNA proficiency tests
12. Notice of any deficiencies and corrective action (if needed)

J. Re-examination

Re-examination casework may be assigned to meet an internal proficiency test requirement or as an additional quality control measure, at the discretion of the TM. The guide for re-examination cases is as follows:

1. Re-examination cases will be selected and issued by the TM of each discipline. Re-examination cases may be selected from evidence already authorized for destruction, if appropriate.
2. If possible and practical, the original analytical results will be unknown to the second analyst.
3. Re-examination results will be reported in the same fashion as original analysis with the following exceptions:
 - a) Provided that the re-examination results are consistent with the original results, the report header will be edited to reflect that it is reported to "case file" instead of a requesting agency or district attorney's office.
 - b) A statement or footer will be placed on the report indicating that the results reported were generated from a re-examination of the evidence conducted as part of a routine OSBI quality control process.
4. Upon completion of the re-examination, the re-examining analyst will provide his/her results to the TM for review **prior to approving his/her report of analysis**.
5. Once completed, all documentation of the re-analysis will be placed in the original case file.
6. The Technical Manager will issue a memo stating the case number, analysts involved in retesting, whether the test was assigned as an internal proficiency test or additional QC, and a summary of results indicating whether results were concordant or discrepant to the QM.
7. Inconsistent results will be evaluated as non-conforming work according to [QP 13](#).
8. The QM will retain re-examination documentation memos.

III. Attachments

None

I. Scope

All case record documentation is subject to review. Review assists in ensuring the quality of the work product. Two types of review will be used: administrative review and technical review.

II. Procedure

A. General

1. The review process does not shift responsibility for an analyst's findings to the reviewer. Each Criminalist has the ultimate responsibility for their casework.
2. Prior to submitting case files for administrative or technical review, each criminalist will thoroughly review his/her own reports and case records. Upon completion of this review the analyst will click on the "Sign" button on the right side of the assignments window. This will lock the report and documentation in the BEAST and prevent unintentional changes.
3. Administrative and technical reviews may be done consecutively in either order (AR-TR or TR-AR) or concurrently (at the same time).
4. Additional administrative or technical reviews (above the minimum required by policy) may be done as necessary or desired by CSD Management.
5. Cases requiring correction will be routed to the appropriate analyst for correction. When routing cases for correction, the reviewer must include an explanation in the comments field describing what must be corrected. **Changes made during or after the review process must be recorded.**

B. Technical Review

Technical reviews will be performed according to the procedure below to ensure that reported conclusions are correct and reasonable, in accordance with validated scientific knowledge, and supported by the examination documentation in the case record.

1. Qualifications of Technical Reviewers:

An individual performing technical reviews must meet the following requirements:

- a. He/she must be a qualified individual who is not the author or co-author of the report **or examination records** being reviewed.
- b. He/she must have received authorization from the discipline Technical Manager (TM) to conduct the technical review based on experience gained through training

and casework experience in the category of testing he/she is reviewing. Analysts may not be authorized to perform technical reviews until they have performed some casework in the category of testing being reviewed.

- c. He/she must have knowledge of the discipline protocols/quality manual applicable to the case he/she is reviewing.

2. Scope of Technical Reviews:

A technical review is a thorough review of the entire case file/record including the Criminalistics Examination Report. Verifications of latent evidence identifications and firearms/toolmarks identifications do not constitute a technical review.

Technical reviews are not required for reports which do not contain the components described in section II.B.3 below (e.g. analytical results, conclusions, associations, etc). Some examples of reports which may not contain these elements include CODIS Database Entry notifications and amended reports, if the correction made does not impact the results, conclusions, or associations of the original report.

For technical reviews, a legible photocopy or fax of the entire file, including the signed report, are to be sent for review only when the original case file cannot be examined on site. **ORIGINAL CASE FILES WILL NOT BE MAILED.**

3. Parameters for Technical Review Process:

Discipline protocols and/or quality manuals may describe more specific portions of examination records which must be checked to complete a technical review. However, any discipline specific review procedures must include a verification of the following, at a minimum:

- a. compliance with all applicable sections of the discipline quality manual and protocols and CSD policies and procedures;
- b. accuracy of the report and that all reported results and/or conclusions are supported by the data in the case record;
- c. any associations are properly qualified in the report;
- d. report contains all information required by discipline quality manual and protocols.

4. Documenting Technical Reviews:

Technical reviews will be documented using one of the following methods:

- a. Using the BEAST to complete the following steps:
 - i. click on the “Tech Rev” button from the assignments tab

- ii. complete the checklist in the window that opens (if applicable)
- iii. click the “Tech Rev” button in the lower right corner
- iv. enter your password and click “yes”

- b. Completing a hard copy version of the checklist (if applicable) and signing and dating the hard copy. The hard copy must then be placed in the case file or scanned and saved to the image vault in the BEAST. This method should only be used for documenting additional technical reviews or if technical difficulties with the BEAST necessitate an alternate method.

5. Handling Technical Review Discrepancies

If an analyst and technical reviewer disagree whether a report and case file meets the criteria listed in section II.B.3, the following steps will be taken to resolve the discrepancy or disagreement.

- a. The analyst and the technical reviewer will discuss the issue and attempt to resolve the issue together.
- b. If the criminalists cannot resolve the issue together, they will seek input from the appropriate Supervisor(s) and/or the Technical Manager.
- c. Any disagreements which cannot be resolved by the Supervisor and/or Technical Manager will be brought to the attention of the Criminalist Administrator assigned to that discipline for further evaluation.

Any non-conforming work identified during review will be handled according to [QP 13](#).

6. Frequency of Technical Reviews:

- a. All case files of Criminalist I’s will be technically reviewed until the Technical Manager of the discipline feels that the Criminalist is fully competent and no longer requires 100 percent technical review. The Technical Manager of the discipline determines the level (within policy) of technical review for employees within their discipline.
- b. The minimum level of technical review for all Criminalists will be at least six cases per month or 20 percent of the cases worked per month, whichever is less.
- c. The Technical Manager of each discipline will determine the number or percentage of cases and the types of cases which must be technically reviewed.
- d. The Technical Manager of the discipline is responsible for designing a system to monitor the number of cases technically reviewed ensuring that the minimum

number specified are reviewed.

C. Administrative Review

1. All OSBI case files, with the exception of BEAST generated no analysis reports, will be administratively reviewed by an individual other than the reporting analyst in accordance with the following procedure, prior to being released to an outside agency.
 - a. Administrative reviews may be performed by any casework qualified analyst or supervisor. A casework qualified analyst or supervisor may conduct administrative reviews for any discipline, not just the discipline for which he/she is qualified to perform casework.
 - b. Administrative reviews may also be performed by a properly trained laboratory technician. Administrative review training for a laboratory technician must include a review of QP 31 and the discipline specific case documentation which he/she will be responsible for reviewing. The appropriate supervisor and/or technical manager will be responsible for documenting the training completed and the date that the laboratory technician has been approved to conduct administrative reviews.
2. Administrative reviews will include the following:
 - a. a review of the report(s) for spelling and grammatical accuracy;
 - b. a review of all administrative and examination records to ensure they are labeled with the case number and initials of the appropriate individual(s) and/or any alternate or additional identification required by discipline or laboratory policy/procedure;
 - c. a review of the report to ensure that all key information required by [QP 28](#) is included;
 - d. a review of the report (for DNA reports) to ensure that it includes a description of the technology and loci or amplification system used and a disposition of the evidence.
3. Documentation of administrative review will be done by one of the following methods:
 - a. Completion of the technical review documentation process for an AR/TR conducted concurrently. It must be documented in the discipline procedure or checklist that the review includes both administrative and technical review.

- b. Completion of a similar BEAST method developed for a specific discipline with the LIMS Administrator's assistance/guidance which has been written into the discipline quality manual or protocol.
- c. Completion of a hard copy or electronic version of [OSBI CSD QPA 31.1](#). A hard copy should be signed and dated by the reviewer and placed in the case file or scanned and saved to the image vault in the BEAST. Alternately, if the form will be placed in the image vault in the BEAST by the reviewer, it may be completed electronically with the analyst's typed name and date.

III. Attachments

[OSBI CSD QPA 31.1, Rev. 1](#)

I. Scope

Each OSBI CSD employee will use this procedure to actively solicit feedback regarding testimony he/she provides.

II. Procedure**A. Procedures for Using the Witness Critique Form**

1. Whenever possible, testifying employees shall give a Witness Critique Form ([OSBI CSD QPA 32.1](#)) to judges, prosecutors, defense attorneys, and other persons in a position to evaluate their testimony. Employees will actively encourage them to objectively complete the form and mail it to the OSBI CSD Quality Manager (QM). Employees may directly accept a completed form to return to the QM.
2. Supervisors and peers will use the Witness Critique Form when observing and evaluating the testimony of laboratory employees.
3. The QM will document all Witness Critique Forms received and forward them to the immediate supervisor of the evaluated employee.
4. The supervisor shall review each evaluation with the employee. Strengths and deficiencies will be noted and discussed, and recommendations for improvements may be made. Both the supervisor and employee will sign the form after the review and discussion. The supervisor shall then return an imaged copy of the signed review form to the QM.
5. The QM will maintain a file of all completed Witness Critique Forms and document who has been evaluated and the numerical rating. Summaries of the results of the evaluations may be given to appropriate supervisors and administrators.
6. The completed Witness Critique Forms and summaries may be used for annual performance evaluations. Supervisors are responsible for maintaining a copy of or other documentation of witness critique forms reviewed with employees throughout the year.
7. Supervisors shall ensure that all their testifying employees are evaluated with a Witness Critique Form at least once each calendar year.
8. Supervisors are strongly encouraged to personally observe the testimony of each of their testifying employees. Once a year is recommended. More frequent intervals are encouraged for less experienced personnel. It is desirable that employees testifying for the first time be observed by their supervisor or a senior peer.

B. Other Methods of Monitoring and Evaluating Employee Testimony

1. It is recommended that supervisors periodically read the transcripts of testimony given by their employees when such transcripts are readily available, and discuss the testimony with the employee.
2. It is recommended that supervisors periodically telephone or personally contact one or more officers of the court to solicit feedback on the testimony of their employees. The information obtained should be recorded on the Witness Critique Form, discussed with the employee, and forwarded to the QM.

C. Corrective Action

When the rating received on a witness critique form is less than satisfactory (average score less than 2 and/or any individual rating of 0), the Supervisor will coordinate with the Technical Manager to review the circumstances of the testimony to determine whether the analyst needs additional training. The supervisor will document the review in the appropriate section of [QPA 32.1](#).

In the event that it is determined that erroneous or misleading testimony was provided, the Supervisor of the testifying employee will follow [QP 13](#) and [QP 14](#).

III. Attachments

[OSBI CSD QPA 32.1, Rev. 1 Witness Critique Form](#)

I. Scope

Laboratory analysis reports and other information concerning OSBI CSD work product are made confidential by State Statutes. Case information may only be provided as allowed by law.

II. Procedure**A. Persons or Agencies Authorized to Receive OSBI Analytical Reports**

1. Title 74 O.S., § 150.2, paragraph 1 states the OSBI shall “maintain a nationally accredited scientific laboratory to assist all law enforcement agencies in the discovery and detection of criminal activity.”
2. Title 74, O.S., § 150.5(D) states: “all records relating to any investigation being conducted by the Bureau, including any records of laboratory services provided to law enforcement agencies pursuant to paragraph 1 Section 150.2 of this title, shall be confidential and shall not be open to the public....” and “the person or entity authorized to initiate investigations (laboratory services) in this section shall receive a report of the results of the requested investigation.” Therefore, a copy of an OSBI laboratory analysis report can be provided to the following requestors of forensic laboratory services:
 - a) Governor
 - b) Attorney General
 - c) Council on Judicial Complaints
 - d) A Legislative Committee with Subpoena Powers
 - e) Chief Medical Examiner
 - f) Director of the Oklahoma State Bureau of Narcotics and Dangerous Drugs Control
 - g) Law Enforcement Agency or Officer
 - h) District Attorney
3. Title 74, O.S., § 150.5(D) further states: “The person or entity requesting the investigation (forensic laboratory services) may give that information only to the appropriate prosecutorial officer or agency having statutory authority in the matter...” therefore, the requestor of forensic laboratory services may provide, or authorize the OSBI Laboratory to provide, a copy of a laboratory analysis report to a prosecutor or prosecutorial agency with jurisdiction in the case.
4. Title 74, O.S., § 150.5(D) also states “officers and agents of the Bureau may disclose, at the discretion of the Director, such investigative information (laboratory analysis reports) to officers and agents of federal, state, county, or municipal law enforcement agencies and to district attorneys, in the furtherance of criminal investigations within their respective jurisdictions.” The OSBI Director has given

authority over laboratory report dissemination to the CSD Director. Unless specifically requested not to by the submitting agency, the CSD Director has authorized that prosecuting attorney offices with jurisdiction may always receive copies of laboratory analysis reports.

5. Regarding the provision of laboratory reports and information to OSBI Investigative Division personnel, authority is provided in two areas. First, intra-agency sharing of forensic laboratory and criminal investigative information in furtherance of criminal investigations or prosecutions is not prohibited by the confidentiality statute, and is desirable and beneficial. Second, the state statute provides for, at the discretion of the Director, the disclosure of records of laboratory services to officers and agents of state law enforcement agencies in the furtherance of criminal investigations within their respective jurisdictions. The person granting the information will document all requests and disseminations in the respective laboratory case file as to the information provided.

B. Criminalistics Analysis Report Dissemination

1. The appropriate number of reports will be made and distributed as follows:
 - a) A minimum of one signed report should be issued to the prosecuting authority having jurisdiction and/or the requesting agency. In some circumstances it may be appropriate to send a report to the requesting agency only. For example, some case types such as property crimes where no suspect has been identified do not need to be distributed to the prosecuting authority.
 - b) No hard copy report needs to be distributed if the receiving agencies have already been issued access to the reports on-line. The list of agencies which don't require hard copy reports will be issued at the discretion of the LIMS administrator.
 - c) An electronic copy of each report prepared using the BEAST Laboratory Information Management System (LIMS) will be maintained in the BEAST.
2. One copy of the report can be retained with the evidence.
3. The only exception to the above is in a situation noted on the RFLE limiting report distribution at the request of the submitting officer, or with the permission of the appropriate Criminalistics Administrator or CSD Director.

C. Inquiries Concerning Case Related Information

1. Occasionally, calls are received requesting information on laboratory cases from the news media, family members, and others.
 - a) All calls from the news media will be routed through the Oklahoma State Bureau

of Investigation Public Information Office.

- b) Other calls from families, friends or others interested in information concerning a CSD case should be directed to the original submitting agency in the case.
- c) The CSD Director must approve any exceptions to the release of case information.

2. Discovery orders will be handled as per OSBI Policy 226.

III. Attachments

None

Rev. #	Effective Date	History
0	10-4-11	<p>Original issue. Combined past QM and QP's into one document with single history and approval. Replaces QM Rev 3, QP 1 Rev 3, QP 2 Rev 1, QP 3 Rev 2, QP 4 Rev 1, QP 5 Rev 4, QP 6 Rev 4, QP 7 Rev 4, QP 8 Rev 3, QP 9 Rev 1, QP 10 Rev 1, QP 11 Rev 3, QP 12 Rev 3, QP 14 Rev 4, QP 15 Rev 2, QP 16.1 Rev 2, QP 16.2 Rev 1, QP 17 Rev 1, QP 18 Rev 1, QP 19 Rev 1, QP 20 Rev 2, QP 21 Rev 1, QP 22 Rev 1, QP 23 Rev 0, QP 24 Rev 2, QP 25 Rev 1, QP 26 Rev 1, QP 28 Rev 2, QP 30 Rev 3, QP 31 Rev 2, QP 32 Rev 1, and QP 33 Rev 1.</p> <p><u>QM</u> updates- Added organizational chart back as OSBI CSD QMA 1, Rev 1. Updated 4.1.5.e,f,i, and 4.1.7 to reference QMA 1. Added 2nd sentence to 4.5.</p> <p><u>QP 1</u> – Revised section II.A.2.a,b,c to match administrative re-organization.</p> <p><u>QP 5</u> – Revised section II.A and II.F to require RFLE. Revised section II.C.1 re: documentation of unloaded firearms. Revised section II.D.1 & II.D.4 for clarification. Added RFLE as OSBI CSD QPA 5.1. Added “or other identifying mark” to section II.B.2.</p> <p><u>QP 6</u> – Renamed as QP 6.1. Added section II.B.5.c to clarify when evidence has been subdivided. Added language to section II.F.4 and 5 regarding proper storage of evidence (toxicology samples and DNA extracts).</p> <p><u>QP 6.2</u> – Inserted new policy re: using evidence for non casework purposes (e.g. training, research, validations, etc). Added notice to customers as QPA 6.2.1.</p> <p><u>QP 7</u> – Incorporated deviation into section II.B.</p> <p><u>QP 8</u> – Added section II.E. 2 to clarify that reagents/supplies are continually evaluated through controls/standards.</p> <p><u>QP 10</u> – Incorporated deviation (deletion of 2nd portion of II.E.1).</p> <p><u>QP 11</u> – Updated QPA 11.1 (Rev. 4) to new contact person for complaints.</p> <p><u>QP 16.2</u> – Revised section II.C.1 to clarify police reports are not required to be retained.</p> <p><u>QP 18</u> – Minor grammatical changes to clarify subcommittee assignments.</p> <p><u>QP 19</u> – Revised section II.D.1 to reflect storage of training records on QA server. Added section II.F.6 and training evaluation form (OSBI CSD QPA 19.1).</p> <p><u>QP 21</u> – Split into QP 21.1 (Research) and QP 21.1 (Validations).</p>

QP 27 – Inserted Operations Policy 400, Rev 3. Revised to match current practices. Added crime scene memo as QPA 27.1.

QP 28 – Added II.A.4.d to prompt return of evidence for no-analysis cases. Revised section II.B.1 and II.C to clarify different requirements for analytical and no-analysis reports and CODIS notifications. Corrected typo in II.C.14. Added “Analytical to header of II.D. Added “and any report footer” to II.D.4.

QP 29 – Inserted Operations Policy 600 Rev 1. Major revision to incorporate use of BEAST and match current testing methods.

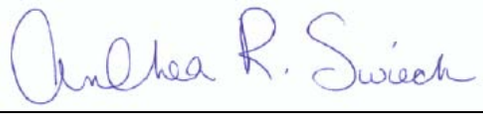
QP 30 – Added language to II.D. and II.D.2 to clarify that external proficiency tests may be assigned to multiple analysts with subsequent analysts’ tests considered as internal. Inserted sections II.F.2 and II.F.4. Major revision to section II.J.

QP 31 – Added 2nd paragraph in II.B.2. Added language to II.C.1 re: who can conduct AR’s and appropriate training.

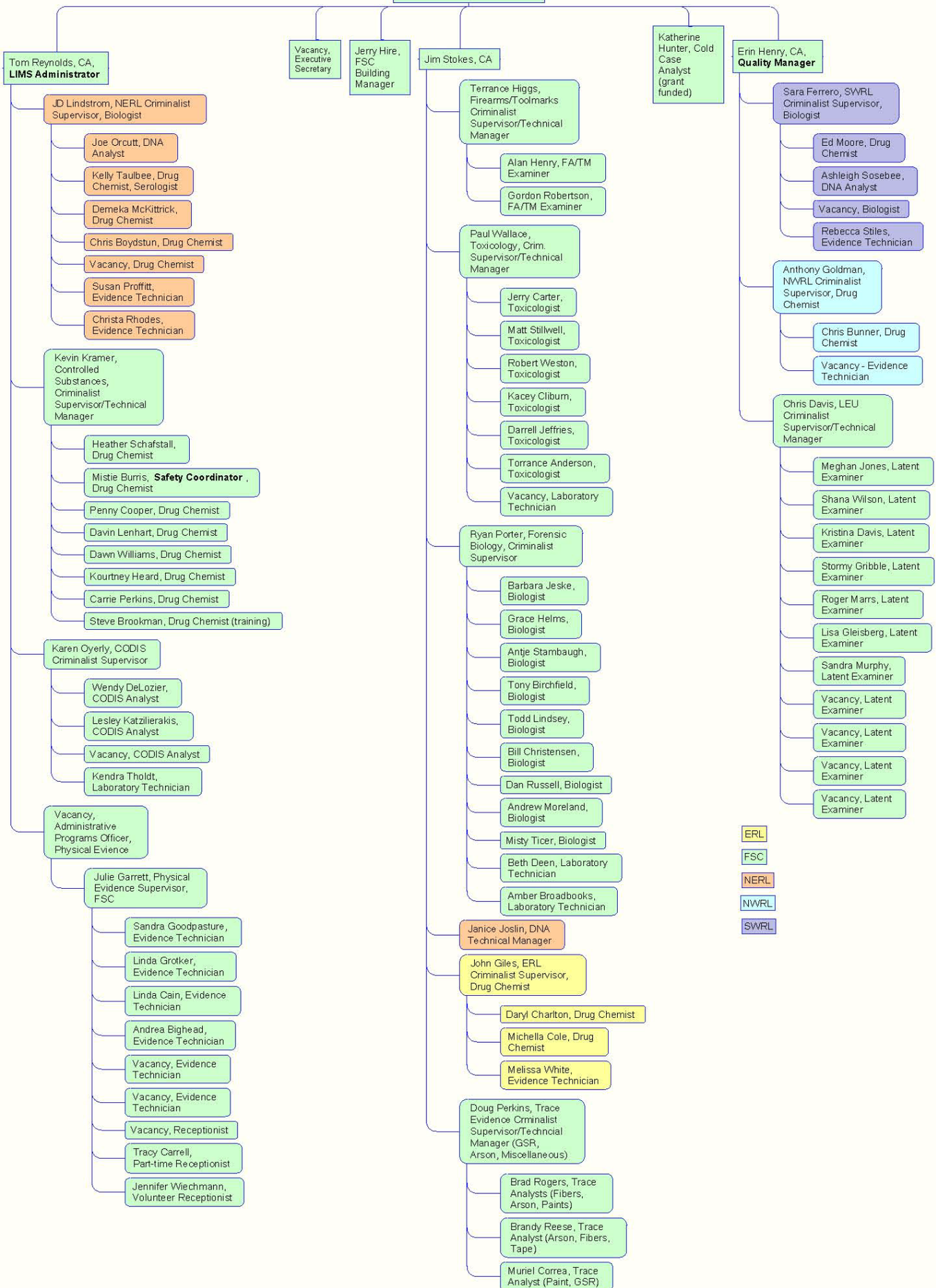
QP 32 – Deleted “the original or” from last sentence in II.A.4. Replaced “will” with “may” in 1st sentence of II.A.6.

QP 33 – Revised section II.B to better match current report distribution practices.

Quality Manager  Date 10-12-11
Erin N. Henry

Division Director  Date 10-12-11
Andrea R. Swiech

Andrea Swiech CSD Director



- ERL
- FSC
- NERL
- NWRL
- SWRL



Notice to Customers

ASCLD/LAB-*International* accreditation requirements state that laboratories must notify customers (investigators, prosecuting attorneys, etc.) in certain circumstances listed below. ASCLD/LAB has established that labs may conduct these notifications on a case by case basis or through a general notification made available to all customers. This notice will serve as a general notification to customers for these areas. Submitting evidence to the OSBI Criminalistics Services Division (OSBI CSD) indicates consent with these terms.

Review of Requests for Analysis (ISO/IEC 17025:2005 Std. 4.4.1):

Each request for forensic analysis is reviewed by OSBI CSD personnel. The OSBI CSD will use the review process to ensure:

1. that the customer's needs are understood (ex: which items need to be processed by the Latent Evidence Unit, etc.), and
2. that the OSBI CSD can meet those needs.

However, the OSBI CSD will determine the most appropriate method(s) of analysis (ex: which chemical processing would best develop latent prints on an item submitted for latent processing) based on the information provided by the customer. Once the OSBI CSD accepts a request for analysis, the accepted request is considered a contract between the requestor and the OSBI.

Changes to Contracts (ISO/IEC 17025:2005 Std. 4.4.4):

In addition, the OSBI CSD may select the item(s) most appropriate for analysis and/or elect to not analyze all items submitted based on the needs and circumstances of the case. The OSBI does not consider this a change to the "contract," and this may be done without additional notice to the customer.

The OSBI CSD does strive to provide the highest quality and most valuable forensic analysis possible. For that reason, if analysts conducting testing identify alternate and/or additional testing that may prove beneficial to our customers, the OSBI CSD may notify the customer on a case by case basis. This notification will always be done if the proposed analysis will require consumption of the evidence and/or limit future examinations.

Subcontracting Analysis (ISO/IEC 17025:2005 Std. 4.5.2):

The OSBI CSD will transfer evidence between OSBI CSD laboratories in order to accommodate efficient analysis.

In limited circumstances, the OSBI CSD may subcontract analysis, if the OSBI CSD cannot provide the service necessary. This includes sending out samples for mitochondrial DNA (mtDNA) analysis and for the DNA identification of human remains and/or associated reference samples.

Any unidentified human remains and/or any reference samples associated with missing persons or unidentified remains that are subcontracted will be sent to the University of North Texas Missing Persons/Unidentified Remains program. Any other evidence requiring mtDNA analysis will be sent to the Federal Bureau of Investigation (FBI) or one of their regional mtDNA labs.

Deviations from Analytical Procedures (ISO/IEC 17025:2005 Std. 5.4.1):

The OSBI CSD utilizes analytical methods that are generally accepted in the forensic science community and that have been validated by OSBI CSD personnel and documented in written protocols. In addition, the OSBI CSD maintains a policy to allow for suggesting, evaluating, approving, and documenting deviations to policy and procedure when necessary. These deviations are not communicated on a case by case basis, but are documented according to policy and can be discussed with customers upon request.

Selection of Methods (ISO/IEC 17025:2005 Std. 5.4.2):

OSBI CSD analytical methods are documented in written protocols and in some circumstances the analytical method used is also referenced in the case file and/or case record. In some circumstances, the analytical methods used may be listed in the examination report, as required by accreditation and quality standards. In any case in which a report does not list the analytical methods used, the OSBI CSD will provide this information upon request. In addition, a list of current services and/or analytical methods currently in use by the OSBI CSD is located in [OSBI CSD QMA 4](#).

**Oklahoma State Bureau of Investigation - Criminalistics Services Division
(OSBI CSD)
Evidence Acceptance Requirements**

The following requirements must be met for the OSBI CSD to accept evidence for analysis:

1. The evidence must be submitted by an individual or agency authorized to request services from the OSBI CSD. The agencies and individuals authorized to request services are listed in Title 74, Sections 150.2 and 150.5. The OSBI CSD cannot accept evidence from private citizens or other individuals/agencies not listed in statute.
2. The evidence must be relevant to an investigation which is expected to result in criminal charges being filed. Evidence relevant to civil investigations or non-criminal product cases such as food or drugs suspected of being old, faulty, etc., will not be accepted for analysis.
3. The evidence must not include any explosive devices, explosive samples, or post-blast samples.
4. The evidence must not include syringes, which under normal circumstances will not be accepted for analysis. Exceptions to this will be evaluated on a case by case basis and exceptions must be approved by a Criminalist Supervisor, Criminalistics Administrator, or designee.
5. Evidence must be submitted in person or through a delivery service such as the United States Postal Service (USPS), United Parcel Service (UPS), or Federal Express (FedEx). Evidence in digital form (images of latent prints, etc.) will not be accepted by e-mail.
6. Evidence must be collected and packaged in a manner that preserves the integrity of the evidence. Evidence which is packaged in a manner that would invalidate the results of testing will be refused. Refer to the OSBI Evidence Collection Manual for information regarding appropriate collection and packaging of evidence.
7. Evidence must be properly collected, packaged, and sealed. Refer to [QMA 3](#) for evidence sealing guidelines.
8. The evidence must have a legitimate associated service request. A listing of available services is detailed in [QMA 4](#). Evidence will not be accepted for the purpose of long-term storage or if the OSBI CSD cannot meet the needs of the customer.

9. Evidence samples submitted for the purpose of comparison (paints, fibers, projectiles, DNA, etc.) must be accompanied by the appropriate reference samples. In most cases, comparison samples will not be accepted unless both the questioned and reference samples are provided. However, this does not apply to samples submitted for comparison to a database. For example, DNA cases with no suspect identified may be submitted with the evidence sample(s) and victim reference sample.

**Oklahoma State Bureau of Investigation - Criminalistics Services Division
(OSBI CSD)
Evidence Packaging and Sealing Guidelines**

Evidence submitted to the OSBI CSD must be stored in an appropriate container under proper seal. The seal must be sufficient to prevent item(s) contained from being lost, removed, or contaminated by outside sources. A container is considered “appropriate” and “properly sealed” only if its contents cannot readily escape and only if entering the container results in obvious damage/alteration to the container or its seal.

1. All evidence must be packaged in a suitable container that protects the evidence from loss, cross-transfer, or contamination.
 - a) Some evidence, such as weapons submitted for serial number restoration or test fire, can be tagged and would not require a container.
 - b) Other evidence, such as liquids may require multiple containers. For example, containers of liquid samples must be stored in a plastic bag, bucket, or other container that will contain the liquid if the immediate container leaks.
 - c) Glass containers should be placed inside a container that will also protect the glass from breaking.
 - d) Containers used to protect the immediate evidence package from leaks or breakage may be treated as convenience packages. However, only one item should be in each “convenience package” to prevent contamination in the event of a leak.

2. All evidence must bear a proper seal. A proper seal includes the initials or other identification of the person sealing the evidence and an acceptable evidence container seal. The following should be used as a guide for acceptable evidence container seals:
 - a) Boxes: A box container seal includes the long seam at both the top and bottom of the box. Boxes should be sealed with two-inch tape, IPG N8315 or equal/better. Evidence tape may also be acceptable.
 - b) Sacks, bags: All sack-like containers should be sealed by folding down the flap of the sack across the top and placing a continuous piece of 2 inch tape, IPG N8315 or equal/better, across the fold and around the sack edges. Evidence tape may also be acceptable.

- c) Envelopes: The top (unsealed) flap of the envelope should be sealed along or across the seam using two inch tape, IPG N8315 or equal/better, or using evidence tape.
 - d) Cans: One continuous piece of evidence tape across the top and down the sides of the container (including buckets with lids) or two separate pieces of evidence tape across from each other are to be used to seal all can-like containers. The seal must be marked or initialed. No clear adhesive tape is to be used on cans if it can be peeled off without evidence of removal.
 - e) Kits: Boxed sexual assault kits, GSR kits, and other purchased kits are acceptable with the seal provided by the kit manufacturer. Envelope-style sexual assault kits shall meet the evidence sealing requirement for envelopes as listed in 2.c above.
 - f) Bulky Evidence: Some items of evidence do not lend themselves to a container. In those cases, the area of interest for analysis should be isolated, protected and marked or initialed. Examples are doors or car bumpers.
3. Evidence which is properly sealed and marked for identification may then be placed in unsealed containers such as boxes or bags for the purpose of grouping items of evidence or for the convenience of carrying the evidence without that container having to meet the requirements of identification and sealing, as long as evidence security requirements are otherwise met. These containers should be marked as a “convenience package” or “convenience container.”
 4. Heat sealing a container of an item of evidence is also acceptable. Identifying marks or initials of the person sealing the evidence must be present across the heat seal.
 5. The submitting officer will be expected to correct improperly sealed evidence prior to the evidence being accepted by the lab. If evidence received by the laboratory has an acceptable evidence seal but is not initialed or marked and that individual is not available to remedy the problem, the receiving personnel will place a piece of evidence tape across the evidence seal at an approximate 90 degree angle and initial across the tape or place the entire evidence container in a heat sealed container and initial across the heat seal.

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)
Facilities and Available Services**

FACILITIES:

The OSBI CSD provides services at the following 5 facilities:

OSBI Forensic Science Center (FSC)
800 East Second Street
Edmond, OK 73034
(405) 330-6724

OSBI Eastern Regional Laboratory
701 West Carl Albert
McAlester, OK 74501
(918) 423-6672

OSBI Northwest Regional Laboratory
1305 E. Garriott
Enid, OK 73701
(580) 242-2600

OSBI Northeast Regional Laboratory
1995 Airport Parkway
Tahlequah, OK 74464
(918) 456-0653

OSBI Southwest Regional Laboratory
5 Northeast 22nd Street
Lawton, OK 73507
(580) 355-6144

For the convenience of OSBI CSD customers, evidence may be submitted at any CSD facility. OSBI CSD personnel will transport evidence between facilities when necessary to provide the appropriate or most timely analysis.

SERVICES:

The following services/analytical methods are available. However, the OSBI reserves the right to select the most appropriate method and to select the item(s) most appropriate for analysis (see [“Notice to Customers” – OSBI CSD QMA 1.1](#)). If a particular test method or service is desired for a specific item, please contact a Criminalist from the discipline in question for assistance with the review of the request.

Biology (FSC, SWRL, and NERL):

1. Screening

Evidence can be screened for biological material including blood, semen, and hair.

2. Confirmatory Testing

Tests are available to confirm the presence of blood and semen.

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)
Facilities and Available Services**

3. Hair Evaluation

Hair samples can be evaluated to determine whether the hair is animal or human and, if human, whether adequate sample is present for nuclear or mitochondrial DNA testing.

4. DNA Analysis

The OSBI CSD can perform two types of Short Tandem Repeat (STR) DNA analysis – autosomal and/or Y-STR analysis. Y-STR analysis generates a DNA profile based on locations on the Y-chromosome only, which means in order to generate a profile, the sample must contain male DNA. Y-STR analysis is only available at the FSC and NERL facilities.

The OSBI CSD can forward evidence to an FBI Regional Mitochondrial DNA Laboratory for analysis.

5. Database Entry/Search

All eligible DNA profiles obtained during the analysis of casework can be entered into the state CODIS (Combined DNA Index System) database and national database (NDIS).

Controlled Substances (FSC, ERL, NERL, NWRL, SWRL):

1. Controlled Substance Identification

Identification of controlled and some non-controlled substances.

2. Controlled Substance Quantitation (FSC Only)

Some evidence items can be analyzed to determine the concentration of the controlled substance.

3. Clandestine Laboratory Analysis

Analysis can be conducted on clandestine laboratory samples to detect controlled substances, precursors, and chemicals related to the illegal manufacture of controlled substances.

4. Poison Identification

Some poisons such as Strychnine can be identified by the drug lab. Other compounds such as Ethylene Glycol (antifreeze) that can be used as poisons can also be identified.

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)
Facilities and Available Services**

Firearms/Toolmarks (FSC):

1. Function Test

Guns submitted for analysis can be tested to determine if the weapon is functional.

2. Trigger Pull Analysis

Analysis can be performed to determine the amount of trigger pull required to fire a gun.

3. Fired Bullet and Casing Analysis

Fired projectiles and/or fired casings can be compared to other fired evidence (bullets/casings) or to a suspect gun.

In addition, fired projectiles and fired casings can also be examined and may sometimes provide information regarding potential makes and models of guns that could have fired the evidence. This is dependent on the amount and type of characteristics present on the fired evidence.

4. Serial Number Restoration

When requested, analysis can be performed to attempt to restore the serial number of a gun.

5. Distance Determination

In some cases, evidence can be examined to determine an approximate distance between an object and the point/location from which a gun was fired.

6. Database Entry/Searching

Test fires from suspect weapons or fired evidence can be evaluated to determine suitability for entry into the Integrated Ballistic Identification System (IBIS). Items entered into IBIS will be automatically searched against the region (Oklahoma and Texas). The OSBI can request searches through other regional databases as well.

7. Toolmark Analysis

Analysis can be conducted to determine, if possible, whether or not a particular tool was used to generate impressions or striations on the item submitted (padlock, window frame, etc.). In addition, analysis can be done to determine if the toolmarks on multiple evidence items were made by the same tool.

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)
Facilities and Available Services**

Latent Evidence (FSC):

1. Footwear Analysis

Photos or casts of questioned footwear impressions can be compared to known shoe samples.

The OSBI CSD cannot examine questioned footwear impressions without known shoes for comparison purposes.

2. Tire Impression Analysis

Photos or casts of questioned tire impressions can be compared to casts or photos of known tire impressions. Tires will not be accepted for comparison purposes.

The OSBI CSD cannot examine questioned tire impressions without known tire impressions for comparison purposes.

3. Latent Print Analysis

Processing:

Items suitable for latent print development which have been properly collected and packaged can be processed to detect and lift/capture latent prints for comparison or AFIS entry.

4. Latent Print Comparison

Questioned latent prints submitted or recovered from items submitted for processing can be compared to known inked impressions submitted or to known impressions from retained records when the subject's information (name, race, sex, date of birth, and SID number) is provided.

5. Database Entry/Searching

All latent prints (including palm prints) of appropriate quality that are not identified to a known can be evaluated for entry into the Oklahoma Automated Fingerprint Identification System (AFIS).

The OSBI CSD can also request a search be conducted using the Integrated Automated Fingerprint Identification System (IAFIS), which searches records from the FBI files.

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)
Facilities and Available Services**

Toxicology (FSC):

1. DUI Cases

Blood or urine collected from individuals suspected of driving under the influence can be analyzed for the presence of alcohol or drugs.

2. Drug Facilitated Sexual Assault

Blood and/or urine collected from an individual reporting a drug facilitated sexual assault can be analyzed for the presence of impairing substances.

3. Alcoholic Content

Liquids suspected of containing alcohol can be analyzed to determine the presence and quantity of alcohol. (Ex: suspected moonshine)

4. Poisons

Samples suspected of containing poison can be tested for select poisons, such as the active ingredient in Visine.

5. Toxic Vapors

Blood may also be analyzed for other substances which cause impairment such as toxic vapors inhaled by a suspect (i.e. huffing).

Trace Evidence (FSC):

1. Ignitable Liquids Residue Analysis

Properly packaged samples of fire debris can be analyzed for the presence of ignitable liquids such as gasoline, paint thinner, or diesel, etc.

2. Primer Gunshot Residue Analysis (GSR)

Evidence submitted using an OSBI GSR Evidence Collection Kit can be analyzed for the presence of elements that are characteristic of gunshot residue (lead, antimony, and barium).

3. Manufactured Fibers:

Questioned fibers can be analyzed and compared to reference or known samples submitted to determine if the questioned and known sample may have originated from the same source. This comparison applies to man-made fibers only.

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)
Facilities and Available Services**

Analysis of questioned fibers can also be conducted to determine the composition of the fiber(s). However, this analysis is limited to the material (e.g. nylon, acetate, etc.) and color. The OSBI CSD does not have the capability to indicate what item(s) may have been a source of the questioned fiber(s).

The OSBI CSD does not perform hair comparisons.

4. Paint Evidence:

Questioned paint samples can be analyzed and compared to known samples, when available, to determine if the questioned and known samples may have originated from the same source.

If known paint samples are unavailable, then unknown samples may be submitted for possible Make and Model determination utilizing the Paint Data Query (PDQ) database.

5. Elemental/Chemical Analysis:

Evidence can be analyzed to determine its elemental composition. The most common application of this analysis is to identify the presence of poisonous materials such as lead, arsenic, and mercury. Elemental analysis can also be conducted to identify elements used in clandestine drug manufacturing, such as phosphorus and iodine.

6. Fracture Matches:

Miscellaneous types of evidence that are torn or broken can be compared to a sample suspected to be the source of the evidentiary sample. For example, duct tape removed from a victim can be compared to a roll of duct tape found in a suspect's possession.

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)**

Alternate Service Providers

In the event that the OSBI CSD cannot provide a particular service requested by a customer, OSBI CSD personnel may assist the customer in locating an appropriate agency or organization that can provide the service. The list below summarizes some of the agencies or organizations that may be able to provide service to customers. This list is intended to be used as a tool to provide assistance to customers and is not intended to serve as a guarantee of service or endorsement of the agencies and organizations listed.

Alcohol, Tobacco, Firearms:

Bureau of Alcohol, Tobacco, and Firearms (ATF):

Firearms, Explosives, Arson—1-800-283-4867

Arson Hotline—1-888-283-3473 (24 Hours)

Explosives Hotline—1-888-283-2662 (24 Hours)

Firearms Hotline—1-888-283-4867 (24 Hours)

Animal/Agricultural:

Animal Disease and Diagnostic Lab

Livestock Diseases

405-744-6623

Department of Agriculture:

Herbicides and Insecticide Poisoning, Animal Food Poisoning

405-521-3864

OSU School of Veterinary Medicine:

Animal Deaths

405-262-5291

Aviation:

Federal Aviation Administration:

Investigations and evidence involved in airplane accidents

405-954-3011

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)
Alternate Service Providers**

Bomb Squads:

Agency	Jurisdiction	Phone
OCPD	All incorporated areas of OKC except for State property	405-297-3477 405-297-1000
OCSO	All unincorporated areas of OKC except for State property	405-713-1044
MWCPD	All incorporated areas of Midwest City	405-739-1388
Edmond PD	All incorporated areas of Edmond	405-354-4420
Norman PD	All incorporated areas of Norman	405-321-1444
Tulsa PD	All incorporated areas of Tulsa	918-596-9222
OHP	All areas of the State that do not have a bomb squad in their jurisdiction.	405-425-2435 405-682-4343 405-202-3763
US Army Fort Sill	Fort Sill EOD will assist any PD or FD 24/7	580-442-8885 580-442-2313

Crime Scene Cleanup:

Heartland BioClean
405-802-6246

AEGIS Biosystems (Edmond)
405-341-4667

Environmental Management, Inc. (Guthrie)
405-282-8510

Traumatix Solutions (Glen Poole)
918-605-2556

Environmental Cleanup, Inc. (Oklahoma City)
405-677-0565

Ferguson Environmental Resources, Inc.
405-495-6336

Crime Scene Cleanup (Reimbursement and Referral):

District Attorney's Council (DAC) Victim Services
405-264-5006

Emergency Management:

Oklahoma Department of Civil Emergency Management
1-800-800-2481 or 405-521-2481

Entomology (Analysis of Insects):

Oklahoma State University
Zoology Department
Stillwater, OK
405-744-5527

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)**

Alternate Service Providers

Environmental Concerns:

Department of Environmental Quality State Lab:

Poisoning of ground water/public or private water supply/soils
405-752-1000

US EPA-ERT Environmental Response Center
2890 Woodbridge Avenue, building #18 (MS 101)
Edison, NJ 08837-3679
1-732-321-4398 <http://www.ert.org>

Food/Drug:

Food and Drug Administration:

Retail Food/Medication (for Human Consumption) Tampering
405-231-4544 (Oklahoma City)
214-253-5200 (Dallas, 24 Hours)

Health Departments:

Non-criminal tampering/contamination of food or over the counter medications
405-271-5243 Oklahoma State Department of Health
405-425-4348 Oklahoma County Health Department
918-595-4301 Tulsa County Health Department

Oklahoma State Department of Health

Biological contamination of foods (e.g. bacterial or viral contaminants)
405-271-5070

Forensics (Fee for Service):

The BODE Technology Group

Springfield, Virginia
703-646-9740

Reliagene

New Orleans, LA
800-256-4106

LabCorp

1912 Alexander Drive
Research Triangle Park, NC 27709
800-533-0567

Serological Research Institute

3053 Research Drive
Richmond, CA 94806
(510) 223-7374

Sorenson Forensics, LLC

2495 S. West Temple
Salt Lake City, UT 84115
800-824-3457 or 888-488-1122

www.sorensonforensics.com

Tarrant County M.E.'s Office

200 Feliks Gwozdz Pl.
Fort Worth, TX 76104
807-920-5700

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)**

Alternate Service Providers

Hazardous Materials:

OHP HAZMAT

405-425-2017
Level A hazmat

Poisoning Deaths:

Office of the Chief Medical Examiner
405-239-7141

Soil Analysis:

FBI Trace Analysis Unit
Attn: Bob Fram
2501 Investigation Parkway
Quantico, VA 22135
703-632-8449

Toxicology:

Bexar County Medical Examiner's Office

Bexar County Forensic Science Center
7337 Louis Pasteur
San Antonio, TX 78229-4565
210-335-4000

Northwest Toxicology (Lab One)
Salt Lake City, Utah
1-800-322-3361 or 1-801-268-2431

El Sohly Laboratories

Oxford, Mississippi
1-662-236-2609

University Toxicology Lab
University Hospital, Rm EB 400
P.O. Box 26307
Oklahoma City, OK 73126
405-271-3840

National Medical Services (Can analyze hair samples for drugs)

3701 Welsh Road
Willow Grove, PA 19090
(800) 522-6671
www.nmslab.com

**Oklahoma State Bureau of Investigation – Criminalistics Services Division
(OSBI CSD)**

Alternate Service Providers

Weapons of Mass Destruction/Terrorism:

63rd Weapons of Mass Destruction Civil Support Team

63rd WMD CST 405-228-5880

Note: normally dispatched via Oklahoma Dept of Civil Emergency Management for response.

FBI Counter Terrorism Unit

405-290-7770 or 405-290-3615

Explosives, radiation

State Department of Health (Biological Weapons (e.g. Anthrax))

Public Health Laboratory Services

P.O. Box 24106

Oklahoma City, OK 73214

Phone: 405-271-5070

Fax: 405-271-4850

405-271-4060 Epidemiologist on call 24 hours
405-271-4341 Security, alternate 24 hr number

OSBI Criminalistics Services Division
Deviation Request Form

I. Explanation of Request

Name: _____ Date: _____

Applies to (Policy/Procedure): _____

Describe Requested Deviation: _____

Specify the Instance/Circumstance for which the Deviation is Requested: _____

Reason for Deviation: _____

II. Technical Review and Authorization

Merits: _____

Risks/Impact: _____

Duration of Authorization: _____

Restrictions/Limitations: _____

Authorized/Rejected _____ (signature) Date: _____

III. Quality Assurance Manager Authorization

Acceptability Within General Quality Assurance Principles? YES/NO

Significant Negative Impact to Division-Wide Quality Standards? YES/NO

Restrictions/Limitations: _____

Authorized/Rejected _____ (signature) Date: _____

IV. Criminalistics Division Director Authorization

Authorized/Rejected _____ (signature) Date: _____

Effective Date: _____



OKLAHOMA STATE BUREAU OF INVESTIGATION
REQUEST FOR LABORATORY EXAMINATION

FOR LABORATORY USE ONLY

Requesting Officer: (TYPE / PRINT - OFFICER'S NAME)

Agency: Phone No.:

Submitting Officer: (Personnel delivering evidence to OSBI Laboratory)

(TYPE / PRINT - OFFICER'S NAME) (OFFICER'S SIGNATURE) Agency

Has there been any previous evidence submitted on this case? YES NO Approx. Date:

Related OSBI Lab #:

Requesting Agency Case #:

TYPE OF OFFENSE:

DATE OF OFFENSE:

COUNTY OF OFFENSE:

SUBJECT/SUSPECT(S):

DNA on file

(For LAB Use Only) (Name, DOB, race, sex, SSN)

Yes No 1.

Yes No 2.

Yes No 3.

VICTIM(S):

(Name, DOB, race, sex)

ITEMIZED DESCRIPTION OF EVIDENCE:

(Attach additional pages if necessary)

- 1.
2.
3.
4.
5.
6.

TYPE OF EXAMINATION(S) REQUESTED:

SEND A COPY OF REPORT TO: (include address)

Copy of report to DA's OFFICE: No Yes

LAB #

Received: (Signature) (Method)

Transp. To LAB:

On (Date) by (Name - Signature)

Received: (Name - Signature)

Transp. To LAB:

On (Date) by (Name - Signature)

Received: (Name - Signature)

DNA Database search: (Initials) on (Date)

Tracking # Container Description

Notification Regarding Non-Casework Use of Evidence

The OSBI Criminalistics Services Division (CSD) sometimes encounters a need to use evidence for purposes beyond the analysis requested in the furtherance of criminal investigations. This notification serves to explain those circumstances and what the OSBI CSD Policy is regarding non-casework use of evidence.

Non Casework Use of Evidence

Re-analysis of Casework:

Annually, the OSBI CSD randomly selects cases to re-analyze as part of our quality assurance program. Re-analysis of casework is conducted and documented in the same fashion as the original examination. Results of re-analysis are not reported to customers unless the results are not concordant with the original examination. The OSBI CSD is in the process of transitioning to web-based distribution of reports. Re-examination reports will be available on the website, but they will be marked to indicate that the results were obtained during routine re-examination.

Research, Validation, and Training:

The other areas where evidence may be needed for non-casework use, include research, validation, and training. Research and validation projects are periodically conducted to evaluate, identify, or incorporate new methods or instrumentation which allow the OSBI CSD to improve efficiency, increase services, or make other quality improvements. Training is conducted for new CSD analysts so that they will be well qualified to analyze casework. When gathering samples for research, validation, and training, the OSBI CSD prepares “mock” evidence samples or uses samples from evidence submitted for destruction whenever possible. However, some circumstances require use of evidence from adjudicated or non-adjudicated cases.

For most cases, the amount taken for research, validation, or training will be comparable to the amount taken for the casework analysis. However, there may be occasions where a more sizable amount needs to be collected. For example, when a large amount of a controlled substance not frequently seen in casework is received, it is beneficial to the OSBI CSD to collect a larger amount to set aside for future training needs.

Notification

As described in OSBI CSD Procedure QP 6.2, the OSBI CSD has notified customers through the posting of this policy. Customers will receive a subsequent notification when samples are collected from active cases. The OSBI CSD will strive to utilize evidence from active cases as little as possible and the removal of evidence for non-casework purposes will be documented in the case record. **The OSBI CSD will not consume more than half of any sample for non-casework purposes, unless the evidence has been re-submitted or otherwise authorized for destruction.**

Submittal of evidence to the OSBI CSD for analysis or destruction indicates that the customer is aware of and accepts this policy.



OSBI
EVIDENCE TRANSFER FORM
LABORATORY USE ONLY

Requesting Criminalist: _____
(Type or Print – CRIMINALIST NAME)

From Unit: _____

LAB # _____

Container Description	Item Number(s)	Evidence Description

TARGET UNIT:

TYPE OF ANALYSIS:

Additional Instructions:

This form contains information for routing convenience only. This form should be shredded upon completion of the related assignment.

OKLAHOMA STATE BUREAU OF INVESTIGATION
CHEMICAL FORMULATION WORKSHEET

Reagent: _____

Lot #: _____

Expiration: _____

Components	Quantity	Supplier	Lot #	Exp. Date

Preparation:

Analyst: _____

Date: _____

Reagent function verification procedure:

This lot **DOES** **DOES NOT** function as expected (Circle one).

Analyst: _____

Date: _____



OSBI Forensic Laboratory Survey

We are dedicated to providing the highest quality services to you. To assist us in doing so, we are asking for your input. Please, take a few minutes to complete this survey. The information you provide will only be shared with the appropriate department Supervisor.

If you have a compliment, concern or complaint, we welcome the opportunity to visit with you about it. We ask that you provide your name and telephone number or email address at the end of this survey if you would like a member of our staff to contact you.

What was the nature of your visit or contact with the OSBI Laboratory?

Laboratory being evaluated: Forensic discipline used: Name of OSBI employee who assisted you:

Please, rate the following based on your experience with our laboratory. Skip a section if it does not apply to your experience with us.

<p>OSBI Staff</p> <p>4 = Excellent 3 = Good 2 = Fair 1 = Poor</p>	<p>___ Knowledgeable</p> <p>___ Accessible</p> <p>___ Friendly</p> <p>___ Professional</p> <p>___ Appearance</p> <p>___ Follow-through</p>
<p>Communications</p> <p>4 = Excellent 3 = Good 2 = Fair 1 = Poor</p>	<p>___ Accuracy of written materials</p> <p>___ Quality of written materials</p> <p>___ Ease of understanding written materials</p> <p>___ Telephone assistance (time on hold, call transfers, wait time for returned calls)</p>
<p>Timeliness of Responses</p> <p>4 = Excellent 3 = Good 2 = Fair 1 = Poor</p>	<p>___ Waiting time in person</p> <p>___ Waiting time by phone</p> <p>___ Waiting time by e-mail/letter</p>
<p>Our Services</p> <p>4 = Excellent 3 = Good 2 = Fair 1 = Poor</p>	<p>___ Turn-around time of examinations</p> <p>___ Breadth of services offered by lab</p> <p>___ Availability of personnel for court, etc.</p> <p>___ Ability of personnel to explain test results</p> <p>___ Adequacy of training courses offered</p>
<p>Facilities</p> <p>4 = Excellent 3 = Good 2 = Fair 1 = Poor</p>	<p>___ Accessibility</p> <p>___ Location</p> <p>___ Cleanliness</p> <p>___ Hours of operation</p>
<p>Website</p> <p>4 = Excellent 3 = Good 2 = Fair 1 = Poor</p>	<p>___ Ease of use</p> <p>___ Information on website</p> <p>___ Design of website</p>

Additional Comments

Do you have an idea to improve customer service or a comment you would like to share:

Do you have a complaint you would like to discuss:

Your Contact Information

If you would like a response to your survey, provide your name and contact information below:

Name

Email Address

Telephone Number (with area code)

Thank you for taking the time to complete this survey!
You can e-mail the completed survey to jimmy.stokes@osbi.ok.gov or print it out and mail/fax to:

**OSBI, Attn: Jim Stokes
800 East 2nd Street
Edmond, OK 73034
Fax 405-330-6207**

OSBI LABORATORY DIVISION COMPLAINT TRACKING FORM

Complainant
Name:

Date:

Organization:

Phone #:

Description

Date Received

Received by:

Logged-in Date:

#:

Was the circumstance verified?

Yes No

Date:

By:

Is the complaint valid?

Yes No

Date:

By:

Complainant notification date:

By:

Assigned to:

Date:

Plan received date:

Completion date:

Final Notification Date:

By:

Notes:

**OKLAHOMA STATE BUREAU OF INVESTIGATION
CORRECTIVE ACTION REQUEST**

Section I		Tracking #:	
Name:	Laboratory:	Date:	
Section II			
Description of event which requires corrective action:			
Section III – Root Cause Analysis		Assigned To:	
Describe the steps taken to determine what the root cause(s) is/are and what is determined to be the most likely root cause.			
Section IV – Description of Corrective Action Plan			
Based on the root cause determined, list potential methods of corrective action and then determine which corrective action plan is most likely to prevent recurrence and is most appropriate to the magnitude and risk that the problem presents. Describe the steps that will be taken to prevent the problem from recurring. Include who will be responsible for various steps of the plan and deadlines for completing each step. Include steps to remediate or correct the work originally identified as non-conforming if it has not already been documented.			
Supervisor:		Date:	
Technical Manager -or- Criminalistics Administrator:		Date:	

**OKLAHOMA STATE BUREAU OF INVESTIGATION
CORRECTIVE ACTION REQUEST**

Section V – Monitoring Plan

Describe the method that will be used to monitor the completion and effectiveness of the corrective action plan. Include who is responsible and deadlines for the completion of individual steps or reviews. If necessary, include attachments.

Supervisor:

Date:

Technical Manager -or-
Criminalistics Administrator:

Date:

Section VI – Completion of Corrective Action Plan

Sign below to indicate that the corrective action plan has been completed. Include a statement indicating whether the corrective action has been effective.

Supervisor:

Date:

Technical Manager -or-
Criminalistics Administrator:

Date:

Section VII – Approval to Resume Work (if applicable)

Technical Manager -or-
Criminalistics Administrator:

Date:



**OKLAHOMA STATE BUREAU OF INVESTIGATION
CRIMINALISTIC SERVICES DIVISION
LINE INSPECTION CHECKLIST**

Laboratory Unit/Region: _____ Supervisor: _____ Date: _____	Related References: Laboratory Quality Policy QP 17 OSBI Directive 230 CALEA Standard 53.1.1 Checklist Created 11-7-01 Revised 12-31-10
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Line inspections are annual inspections and evaluations of each regional laboratory or laboratory unit conducted by the regional or unit supervisor approximately 6 months after their internal annual audit (staff inspection). It is to determine if employees are adhering to applicable Bureau and Division policies, procedures, rules, and regulations. It is also an inspection of facilities, property, evidence, equipment, activities, and personnel. Regional and unit supervisors will note all deficiencies, take corrective action as appropriate, and archive copies of the reports.

Inspection Criteria	Yes	No	N/A	Comments	Corrective Action (CA) Needed	CA Due Date	CA Completion Date
Technical protocol manuals current & available?							
Are employee's laboratory operations and quality manuals current?							
Do employees have their credentials, badge, CLEET card, , phone card, and valid driver's license?							
Were annual safety audits performed when required?							
Do all employees wear the appropriate PPE when required?							
Are laboratory bench supplies adequate?							
Are instruments properly calibrated and maintained?							
Are instrument calibration and maintenance logs complete and current?							
Were Quality Audit deficiencies corrected?							
Are administrative reviews performed as per policy?							
Are technical reviews performed as per policy?							
Are PMP evaluations up-to-date on all employees?							
Are technical procedures followed in casework?							
Is evidence maintained securely under proper seal?							



**OKLAHOMA STATE BUREAU OF INVESTIGATION
CRIMINALISTIC SERVICES DIVISION
LINE INSPECTION CHECKLIST**

Inspection Criteria	Yes	No	N/A	Comments	Corrective Action (CA) Needed	CA Due Date	CA Completion Date
Is evidence chain-of-custody complete and documented?							
Is evidence marked for identification?							
Are laboratory and office areas clean, safe, and orderly?							
Is analysis well documented in the case files?							
Is completed case evidence returned quickly?							
Is security of the laboratory and evidence rooms maintained?							
Do employees follow time and leave policies?							
Are periodic staff meetings held with employees?							
Is reference literature appropriate and available to employees?							
Are appropriate controls and standards used in casework?							
Are critical reagents function tested before use?							
Is analyst's testimony being monitored?							
Are sufficient numbers of staff trained in first aid?							
Do employees receive ongoing annual training?							
Is monthly case reconciliation performed?							
Is emphasis placed upon closing older cases?							
Are resources directed toward reducing case turnaround times?							
Are laboratory reports written as per policy?							
Do employees understand organizational values?							
Is each analyst in compliance with the proficiency testing policy?							
Are employees in compliance with the appearance policy?							

Unit/Region _____ Date _____ Supervisor _____

The supervisor maintains the inspection form until all corrective actions are completed and verified through a re-inspection. When completed, the original form is sent to the Division Director of Criminalistics. The unit or regional supervisors maintain a copy, and copies are sent to their Criminalistic Administrator and the Quality Manager. The line inspection is a valuable tool for managing a laboratory unit or region, and maintaining our laboratory quality program. Inspections are required to be done once in each calendar year, however supervisors have the discretion to perform line inspections more frequently as necessary.



OSBI Training Report Form

For training conferences attended, please use this form to capture the requested information. The information will be used by the OSBI Laboratory Administration and grant organizations.

Date of this report: _____ Grant number (if applicable): _____

Individual(s) who attended training: _____

Training attended, location, and dates: _____

List each new technique, method, procedure, idea, and/or concept learned and implemented by you that was a direct result of the training attended.

1. _____
2. _____
3. _____
4. _____
5. _____

For each new technique, method, procedure, idea and/or concept listed above, cite **specific improvement(s)** in the quality, timeliness (improved case turn around times), and/or case backlogs of the forensic science services provided by you that were learned at the training.

1. _____
2. _____
3. _____
4. _____

Please submit this completed form within 7 working days of the completion of the training to the Lab Administration.

Thank you for your cooperation in providing this information.

TO: Criminalistics Services Division Director

SUBJECT: Crime Scene Memo OR MORGUE

FROM:

Criminalistic Case #:

Type of Offense:

Date of Offense:

Victim(s):

COMPLETE THE FOLLOWING FOR LATENT PRINT CRIME SCENES

Which techniques were utilized at the crime scene?

Dusting: Yes No
If Yes, Dusting time was Less Than or More Than 5 hrs

Reagent Spray: Yes No
If Yes, which reagent? LCV Other: _____

Alternative or high intensity light: Yes No

Suspect(s):

Request for Assistance From:

Agency:

	DATE	TIME
Request for Assistance Received		
Left for Scene(s):		
Arrived at Scene(s)		
Left Scene(s) to Return		
Returned from Scene(s)		
Total Time Out (hrs/min)		

COMPLETE THE FOLLOWING FOR ALL CRIME SCENES

Were blood, bodily fluids or excretions present at the scene?
 Yes No

If Yes, were you working in the immediate vicinity of the blood, bodily fluids, or excretions? Yes No

If Yes, was the biological material wet and/or dry?

Was luminol utilized at the crime scene? Yes No Unknown

If Yes, were you working in the immediate vicinity where it was used? Yes No

Which of the following personal protective equipment was utilized by Criminalists responding with your group (i.e. those that traveled together)?

Gloves: Yes No Protective Eyewear: Yes No

Respiratory Protection: APR Dust Mask None

Disposable Booties: Yes No

SCENE INFORMATION:

Location: County

Comments/Explanation:

Address City

Atmospheric Conditions of Scene: Sunny Rain Snow Cloudy Night Windy Other:

Approximate Temperature: Outside Scene Indoor Scene

List of Individuals met at the scene and the agencies they represent:

Officer in Charge of Scene:

Summary of what Criminalistic Personnel did at scene (Processed house,car,etc. List serial numbers if applicable. Include brief description of scene.)

Name(s) of OSBI Criminalistics Personnel responding to scene(s), indicate which ones traveled together.

OSBI Unit # used for transportation: Total Mileage Out:

- DISTRIBUTION OF COPIES OF MEMO:**
- Copy to be forwarded to Criminalistics Services Division Director
 - Copy to be forwarded to Physical Evidence Technician Supervisor
 - Original to be placed in the criminalistics case record, if applicable
 - Copy for the Criminalist(s) personal file (Optional)

(NOTE: If the OSBI personnel responding did not leave together, travel together in the same unit and return together, separate crime scene memos must be completed for each group.)

CRIMINALIST COMMENTS:

OKLAHOMA STATE BUREAU OF INVESTIGATION CRIMINALISTIC SERVICES DIVISION

CASE FILE ADMINISTRATIVE REVIEW FORM

Analyst: _____ **Case #:** _____ **Report No.:** _____

By signing or completing this form, I verify that the following items have been reviewed and verified to be in compliance with applicable CSD policies and procedures:

1. The report has been reviewed for spelling and grammatical errors.
2. All administrative and examination records have been reviewed to ensure they are labeled with the case number and initials of the appropriate individual(s) and/or any alternate or additional information required by discipline or laboratory policy and procedure.
3. The report has been reviewed to ensure that all information required by QP 28 is included.

Corrections Required (if applicable):

Examiner's response (if appropriate):

Admin. Review (sign and date): _____

Second Admin Rev. (initial and date): _____

Signatures indicate any necessary corrections have been completed.



OKLAHOMA STATE BUREAU OF INVESTIGATION

800 East Second Street

Edmond, OK 73034

Fax: 405-330-6207 Phone: 405-330-6724

Attn: *Laboratory Quality Assurance Manager*

Witness Critique

In furtherance of our goal to provide accurate and reliable testimony in a professional manner, we have developed this questionnaire to collect information that will help the OSBI laboratory better evaluate our employees' courtroom testimony. Please take a minute to answer the questions below, then return the form. Your opinions, observations and suggestions are important to our organization, and will help us improve the quality of service we provide to you.

Thank you for your assistance.

Witness' Name	_____	Court #	_____
Court/County	_____	OSBI Lab #	_____
Testimony Date	_____	Defendant	_____

Please rate the examiner's testimony:

	<u>Excellent (3)</u>	<u>Good (2)</u>	<u>Fair(1)</u>	<u>Poor(0)</u>
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Please leave blank if not applicable.

- | | | | | |
|--|--------------------------|--------------------------|--------------------------|--------------------------|
| 1. Courtroom demeanor and appearance | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2. Ability to convey information in an understandable manner | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3. Poise and professionalism during direct examination | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4. Poise and professionalism during cross examination | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5. Use of court exhibits/visual aids (if applicable) | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6. Testimony based upon scientific principles
(to be rated by lab reviewers only) | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |

Remarks/Comments/Suggestions (please explain poor ratings): _____

Your Name (optional) _____	Judge <input type="checkbox"/>	Prosecutor <input type="checkbox"/>	Defense <input type="checkbox"/>
	OSBI Supervisor <input type="checkbox"/>	Peer <input type="checkbox"/>	Other <input type="checkbox"/>

(FOR INTERNAL USE BY OSBI ONLY)

REVIEWED WITH EMPLOYEE:

_____ Employee

_____ Date

_____ Supervisor

Additional Training Required (circle one and initial): YES NO N/A