



Standard Guide for Validation of Laboratory Information Management Systems¹

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1. Scope

1.1 This guide describes an approach to the validation process for a Laboratory Information Management System (LIMS).

1.2 This guide is for validation of a commercial LIMS purchased from a vendor. The procedures may apply to other types of systems, but this guide makes no claim to address all issues for other types of systems. Further, in-house developed LIMS, that is, those developed by internal or external programmers specifically for an organization, can utilize this guide. It should be noted that there are a number of related software development issues that this guide does not address. Users who embark on developing a LIMS either internally or with external programmers also should consult the appropriate ASTM, ISO, and IEEE software development standards.

1.3 This guide is intended to educate individuals on LIMS validation, to provide standard terminology useful in discussions with independent validation consultants, and to provide guidance for development of validation plans, test plans, required standard operating procedures, and the final validation report.

2. Referenced Documents

2.1 ASTM Standards:

- E 622 Guide for Developing Computerized Systems²
- E 623 Guide for Developing Functional Requirements for Computerized Systems³
- E 624 Guide for Developing Implementation Designs for Computerized Systems³
- E 627 Guide for Documenting Computerized Systems²
- E 919 Specification for Software Documentation for a Computerized System²
- E 1013 Terminology Relating to Computerized Systems²
- E 1384 Guide for Content and Structure of the Electronic Health Record (EHR)²
- E 1578 Guide for Laboratory Information Management Systems (LIMS)²
- E 1639 Guide for Functional Requirements of Clinical

Laboratory Information Management Systems (CLIMS)

2.2 IEEE Standards:⁴

- 100 Standard Dictionary of Electric and Electronic Terms
- 610 Standard Glossaries of Computer-Related Terminology
- 729 Glossary of Software Engineering Terminology
- 730.1 Standard for Software Quality Assurance Plans
- 730.2 Guide for Software Quality Assurance Plans
- 828 Standard for Software Configuration Management Plans
- 829 Standard for Software Testing Documentation
- 830 Guide for Software Test Documentation
- 1008 Standard for Software Unit Testing
- 1012 Standard for Software Verification and Validation Plans
- 1016 Recommended Practice for Software Design Descriptions
- 1028 Standard for Software Reviews and Audits
- 1042 Guide to Software Configuration Management
- 1058-1 Standard for Software Project Management Plans
- 1063 Standard for Software User Documentation
- 1074 Standard for Developing Software Life Cycle Processes

1228 Standard for Software Safety Plans

2.3 ISO Standards:⁵

- 9000 Quality Management and Quality Assurance Standards - Guidelines for Selection and Use
- 9000-3 Guidelines for Application of ISO 9001 to Development, Supply, and Maintenance of Software
- 9001 Quality Systems—Model for Quality Assurance in Design, Production, Installation, and Servicing
- 9002 Quality Systems—Model for Quality Assurance in Production and Installation
- 9003 Quality Systems—Model for Quality Assurance in Final Inspection and Test
- 9004 Quality Management and Quality System Elements—Guidelines
- 9004-2 Quality Management and Quality System Elements, Part 2 Guidelines for Services
- 9004-4 Guidelines for Quality Improvements
- 10005 Guidelines for Quality Plans
- 10007 Guidelines for Configuration Management

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² Annual Book of ASTM Standards, Vol 14.01.

³ Discontinued 1994; see 1993 Annual Book of ASTM Standards, Vol 14.01.

⁴ Available from Institute of Electrical and Electronic Engineers, Inc., 445 Hoes Lane, P. O. Box 1331, Piscataway, NJ 08855-1331.

⁵ Available from International Organization for Standardization, 1 rue de Varembe, Case postale 56, CH-1211 Genève 20, Switzerland.

- 10011-1 Guidelines for Auditing Quality Systems, Part 1 Auditing
- 10011-2 Guidelines for Auditing Quality Systems, Part 2 Qualification Criteria for Auditors
- 10011-3 Guidelines for Auditing Quality Systems, Part 3 Managing Audit Programs
- 8402 Quality Vocabulary
- 2382 Data Processing Vocabulary

3. Terminology

3.1 *Definitions*—This guide defines terminology used in the validation of computerized systems. The standards listed in Section 2 provide additional definitions that the reader may want to review before beginning their validation process.

3.1.1 *acceptance criteria, n*—the specifications used to accept or reject a computer system, application, function, or test action.

3.1.2 *change control, n*—the process, authorities for, and procedures to be used to manage changes made to a computerized system or a system's data, or both. Change control is a vital activity of the Quality Assurance (QA) program within an establishment and should be described clearly in the establishment's SOPs.

3.1.3 *configuration management, n*—a discipline applying technical and administrative direction and surveillance to identify and document the functional and physical characteristics of a configured item, to control changes to those characteristics, to record and report change implementation status, and to verify compliance with specified requirements.

IEEE

3.1.4 *customization, n*—the process of adding new software to or altering a LIMS so that it may perform functions not planned by the original system designers. This entails creating new software, compiling software modules, and linking modules to produce new executable programs. If done by the vendor, it may be considered and validated as part of the vendor system. See related definition for "customized system" in Terminology E 1013.

3.1.5 *delivered system, n*—the LIMS, as initially supplied by the vendor before any static configuration data have been added. In some cases, the vendor may contract with the laboratory to enter some configuration data on behalf of the laboratory, in which case the delivered system is still considered to be the default system before such customer-specific information has been added. When the vendor performs this task, they are an agent of the laboratory, and the customer shall meet the on-site validation requirements in Section 7.

3.1.6 *dynamic testing, n*—the actual testing of various functions and procedures using the LIMS software while in operation.

3.1.7 *installation qualification (IQ), n*—documented verification that all key aspects of the installation adhere to approved design intentions as defined in system specifications and that manufacturers' recommendations are suitably considered.

3.1.8 *LIMS, n*—acronym for Laboratory Information Management System that refers to computer software and hardware that can acquire, analyze, report, store, manage data, and process information in the laboratory.

3.1.9 *LIMS data loading (configuration), n*—the process of

entering static data into appropriate data structures, such as tables or database records, to make a LIMS suitable for operation in a particular laboratory. This information may include items like names and addresses of laboratory customers, names of laboratory personnel, descriptions of tests performed by the laboratory, specifications, calculations, templates, or descriptions of LIMS reports, etc. In this process, no new functionality is added to the LIMS that was not originally planned by the system designers. Addition of configuration data may affect the behavior of the system.

3.1.10 *LIMS tailoring, n*—see *LIMS data loading (configuration)*.

3.1.11 *operational qualification (OQ), n*—documented verification that each unit or the entire system operates as intended throughout its full operating range.

3.1.12 *quality assurance unit (QAU), n*—the body of individuals responsible for design and interpretation of quality standards, such as validation procedures and processes (not product testing).

3.1.13 *source code, n*—a computer program expressed in human-readable form (programming language) that shall be translated into machine-readable form (object code) before it can be executed by the computer.

3.1.14 *static testing, n*—a structured review of the source code.

3.1.15 *stress testing, n*—the running of test protocols designed to test the limits of LIMS functions.

3.1.16 *test plan, n*—see *test protocol*.

3.1.17 *test protocol, n*—a written procedure describing a set of actions and their expected outcomes that when executed provides documentary evidence that specific functional requirements for the LIMS work as specified.

3.1.18 *validation, n*—the process of establishing documented evidence that provides a high degree of assurance that a specific process, system, or item consistently meets its predetermined specifications or quality attributes.

3.1.19 *validation plan, n*—the document that identifies all systems and subsystems involved in a specific validation effort and the approach by which they will be qualified and validated, including identification of responsibilities and expectations.

3.1.20 *validation team, n*—the group of individuals responsible for the validation process. This team may consist of representatives of the laboratory, QAU, Management Information System (MIS) organizations, or outside consultants.

3.1.21 *vendor audit, n*—an independent review and examination of system records and activities in order to test the adequacy and effectiveness of data security and data integrity procedures, to ensure compliance with established policy and operational procedures, and to recommend any necessary changes.

ANSI

3.1.22 *vendor audit team, n*—a team made up of individuals who are knowledgeable in computer system engineering, auditing practices, computer system quality methods, regulatory compliance, validation practices, business and legal policies and procedures (applicable only to computer hardware and

software procurement and related services). (1)⁶

3.1.23 *version control, n*—control of all associated software and document versions. This also includes all documents associated with implementation, validation, or operation of a LIMS.

4. Significance and Use

4.1 Validation is an important and mandatory activity for laboratories that fall under regulatory agency review. Such laboratories produce data upon which the government depends to enforce laws and make decisions in the public interest. Examples include data to support approval of new drugs, prove marketed drugs meet specifications, enforce environmental laws, and develop forensic evidence for trial. This also extends to LIMS used in environmental laboratories. In some cases these systems may need to be interoperable with CLIMS and computer-based patient records (CPR) for reporting environmental exposures and clinical laboratory testing for biologic measure of stressor exposure. The enormous financial, legal, and social impact of these decisions requires government and public confidence in laboratory data. To ensure this confidence, government agencies regularly review laboratories operating under their rules to confirm that they are producing valid data. Computer system validation is a part of this review. This guide is designed to aid users validating LIMS and incorporating the validation process into their LIMS life cycle.

4.2 Validation must provide evidence of testing, training, audit and review, management responsibility, design control, and document control, both during the development of the system and its operation life (2).

5. The LIMS Life Cycle and the Validation Process

5.1 The process of validation should start at the beginning of the LIMS life cycle as defined in Guide E 1578. Adding validation to the end of the LIMS implementation could add from three to twelve months to the LIMS project. Further, adding validation to the end of the process would prevent the organization from using the LIMS during validation. Fig. 1 represents points where validation may impact the procurement of LIMS. Validation will not have an impact on all of the LIMS life cycle, and the amount of interaction with the validation team will vary during each life cycle phase.

5.1.1 *Validation Team Formation Phase*—This phase is typically not a separate phase in the LIMS life cycle, however, it is a critical part of the validation process. A typical team consists of representatives from the laboratory, MIS group, and QAU. There may be other team members depending on the scope of the project and resources within the organization. If required, the identified validation team members should begin to identify training courses on computer systems validation at this time. No training should take place until those who have been selected for the validation team have their management's full agreement to participate in this activity. These courses can be either in-house or outside-developed courses. The vendor audit team may consist only of the validation team or it may be

a specific subgroup within the organization. It is recommended that the vendor audit team should include organizational members from the QAU, MIS, and the laboratory (1).

5.2 *Business Requirements Definition Phase*—The business unit, specifically the laboratory, shall contact the QAU to determine current good manufacturing practices (cGMPs), good manufacturing practices (GMPs), good automated laboratory practices (GALPs), and other requirements that shall be addressed with this project. An initial selection of validation team members is made at this time.

5.3 *Project Definition Phase*—Final agreement and management acceptance for all validation team members should be obtained. Because validation is complex and can take a long time, each team member should have the full support of their management. It is critical that management understands and agrees to the time commitment for these individuals. Without agreement from each member's management chain, the probability for developing and validating the LIMS successfully will diminish. Once formed, the validation team can start to address high-level issues such as the existence of corporate standard operation procedures (SOPs) needed for validation. Time constraints and inexperience of team members can be a limiting factor in the validation process. This is when the team should identify outside consultants that may be needed in the validation process and begin developing the validation plan. Appropriate training of validation team members also should be carried out during this phase of the LIMS life cycle.

5.4 *Model of Current State of Laboratory Practice*—The validation team typically is not part of this process.

5.5 *Model of Future State of Laboratory Practices*—The validation team typically is not part of this process.

5.6 *Functional Requirements Development Phase*—The validation team should work with the group responsible for developing functional requirements. At this time the team can also begin to develop and revise, as necessary, a high-level draft of the organization's validation plan for this project. The validation team may want to begin developing the high-level test protocols during this phase. Further this activity begins to focus attention on validation at the start of the project. Each identified functional requirement should be the subject of one or more test protocols.

5.7 *Request for Proposal (RFP) Phase*—The validation team shall ensure that the RFP includes both a request to audit the vendor and their validation requirements. People using this document for acceptance testing who are in unregulated industries may not require this audit process. Also, the validation team should request that the vendor's development process and LIMS application have undergone independent evaluation/validation. If another company, that is, a third party consultant or another corporation, has validated the vendor operation and LIMS development process, it does not mean that the prospective buyer can assume that the software is validated. During this time the team should specify what actions to take if a LIMS vendor denies them the right to an audit. The validation team should review the RFP prior to its submission to the vendor.

5.8 *Evaluation and Selection Phase*—The validation team should identify those people who will participate in vendor

⁶ The boldface numbers in parentheses refer to the list of references at the end of this standard.

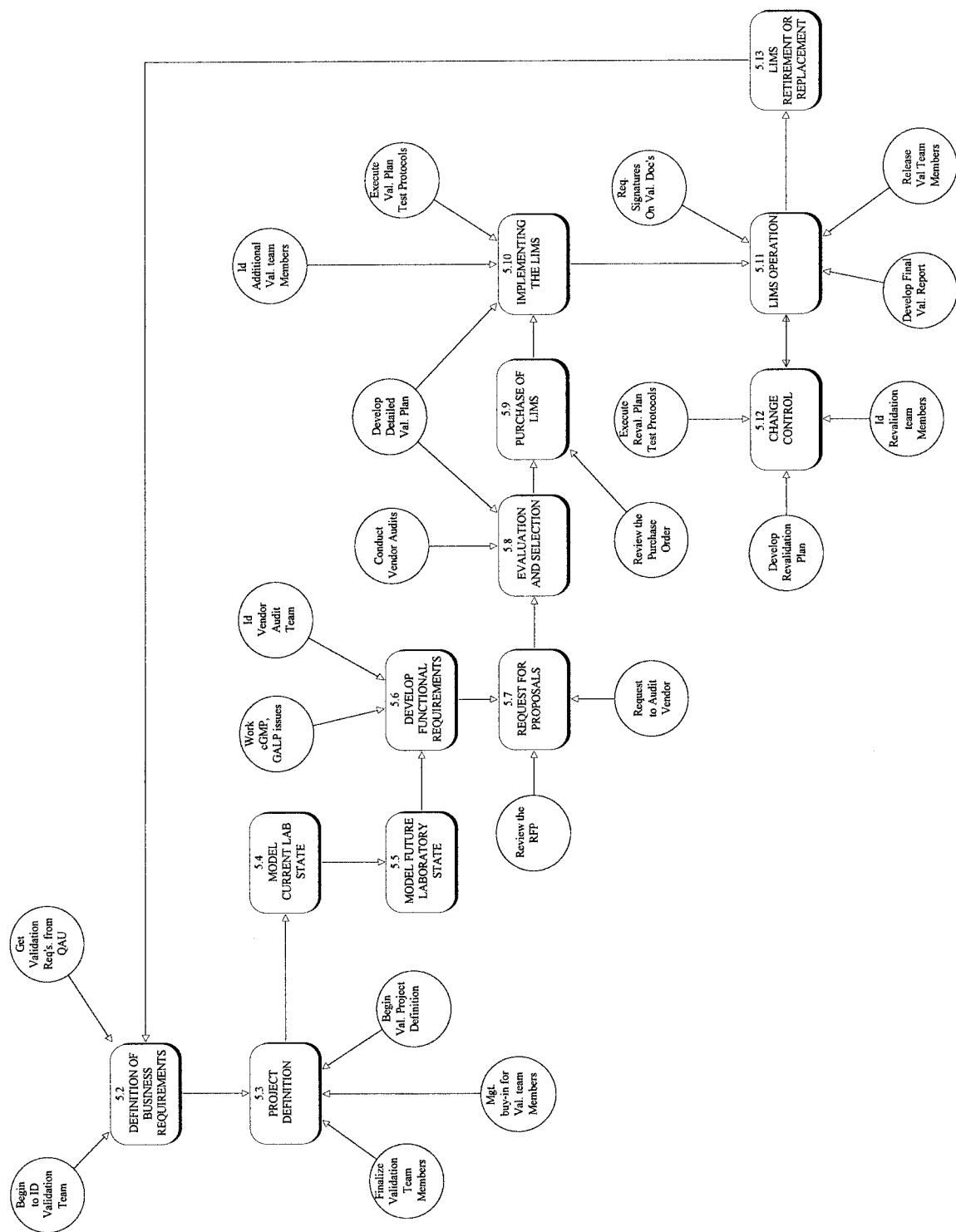


FIG. 1 LIMS Life Cycle

reviews. Since this process can take from one to several days, only those LIMS manufacturers targeted by LIMS team should be visited. The prioritized selection of LIMS shall be based upon the vendor's answers to the RFP. The RFP answers will normally emphasize the stated functional requirements. Perform a vendor audit to find the built-in quality. Continue vendor audits until an acceptable vendor for both quality and function is found. The audit results are useful in assessing the buyer's exposure to risk when system functionality is balanced against quality of system development. See Section 6 for more auditing of the LIMS vendor.

5.9 Purchase—Validation team members should review and be part of the purchase order approval process to ensure validation issues and criteria outlined in 5.8 are met and to begin the early stages of configuration management.

5.10 Implementation Phase—The validation team shall finalize the validation plan and other documentation that must be approved by the system owner and authorized by QAU before the plan is carried out. A schedule of events is developed. Testing protocols will be executed and the results documented. When all test protocols have been executed and documented, the final validation report is developed and the required signatures are obtained to approve this report. The final approval will be obtained from the system owner as authorized by QAU.

5.11 Operational Phase—When all validation tasks have been completed, the validation team can be disbanded. Tasks in this area include the following:

5.11.1 Ongoing training of new users.

5.11.2 Modification of SOPs to address necessary changes to the LIMS or its operational environment.

5.11.3 Review of procedures and their adherence to existing SOPs, documenting compliance with SOPs.

5.11.4 Maintenance of change control procedures for the existing system.

5.11.5 Maintenance of the system.

5.11.6 Upgrades to the LIMS hardware or software. This also includes all associated hardware or software in the LIMS operating environment, that is, the LAN, computers' operating system, etc. See the change control phase in 5.12.

5.12 Change Control—The LIMS Manager will face change control issues often during the normal operation of a LIMS. The LIMS Manager must understand that all minor and major changes to the system shall be subject to change control, assessment of consequences, and revalidation after the change takes place. Upgrades in software as well as changes in how the system is used may require revalidation. The change control committee may determine the system changes require revalidation. All changes shall be documented, as well as assessment of the need to validate the change and the extent of the revalidation. The level of detail for the revalidation process depends upon the type of change. A new validation team may be needed. This team may wish to include some test protocols from the original validation process. The degree of revalidation is highly dependent upon the impact of the identified change. Change requests and problems should be documented (see Appendix X6) (3).

5.13 Retirement/Replacement of the LIMS—The process

starts over with the establishment of a new validation team.

6. LIMS Vendor Assessment/Audit

6.1 Industry regulators require laboratories to ensure that computer applications, such as LIMS, are validated. It is the responsibility of the laboratory owner to demonstrate that specific applications are developed, tested, operated, and maintained according to accepted quality practices.

6.2 The regulatory authorities expect that organizational personnel will follow the formal policies governing operations, as well as, comply with the proper levels of control and documentation. Further, they expect vendors to use the same level of quality control and quality practices as the customers they are supplying. It is the system owner's responsibility to investigate the vendor's operation and verify that they have accepted practices in place and that they are using them. The system owner can use the vendor audit to inspect and evaluate the vendors quality assurance programs, practices, and documentation procedures.

6.3 An organization may want to outsource vendor audits when they lack the organizational expertise, see it as a more cost effective, or they want a more objective or thorough audit. The use of audit results from a third party not associated with the user's organization, or those performed by another corporation, may not be used as a substitute for auditing the vendor. Alternatively, an audit that is jointly conducted by a consortium of corporations all looking to use a particular vendor's application has been used in the past with regulatory authority approval.

6.4 Vendor assessment should occur during the evaluation and selection phase of the LIMS life cycle and before final vendor selection. If the organization already has a vendor audit team established, this group should review their system functional requirements with the LIMS validation team. If the organization does not have such a team already established, they may want to have members of the LIMS validation team perform the vendor auditing. The audit team should be comprised of an experienced software auditor internal or external to the company and one or more individuals from the LIMS team. In general, there should be someone on the audit team responsible for the long-term relationship with the vendor. Typically, this person is the system or application owner.

6.5 The primary goal of the audit is to ensure that the vendor's software development and management procedures are consistent with the accepted practices, that is, those which are traceable back to a reference point and to which these practices adhere. This means that the audit team shall assess the vendor's quality measures, which affect the product they sell and the quality support they provide in the future. The audit team can meet this objective by gathering evidence, which demonstrates that the LIMS vendor is adhering to well-defined and documented software development and maintenance standards or practices (4).

6.6 In addition to these objectives, the auditing organization should evaluate the vendor's financial health and stability (1). It should be noted that even though a LIMS vendor organization is registered as meeting national or international requirements, for example, ISO 9001, the vendor is not exempt from being audited by their customers. The purchasing organization

is still responsible for auditing the prospective LIMS vendor. See Fig. 2 for the GAMP 96(5) guideline on the auditing process flowchart.

6.7 The vendor assessment should cover software development, software maintenance, quality and control issues (4).

Key areas that should be targeted for inspection include documentation that supports system testing, preventive maintenance, operation and maintenance manuals and administrative procedures (1). The source code review process should be limited to a random sampling of the source code modules that

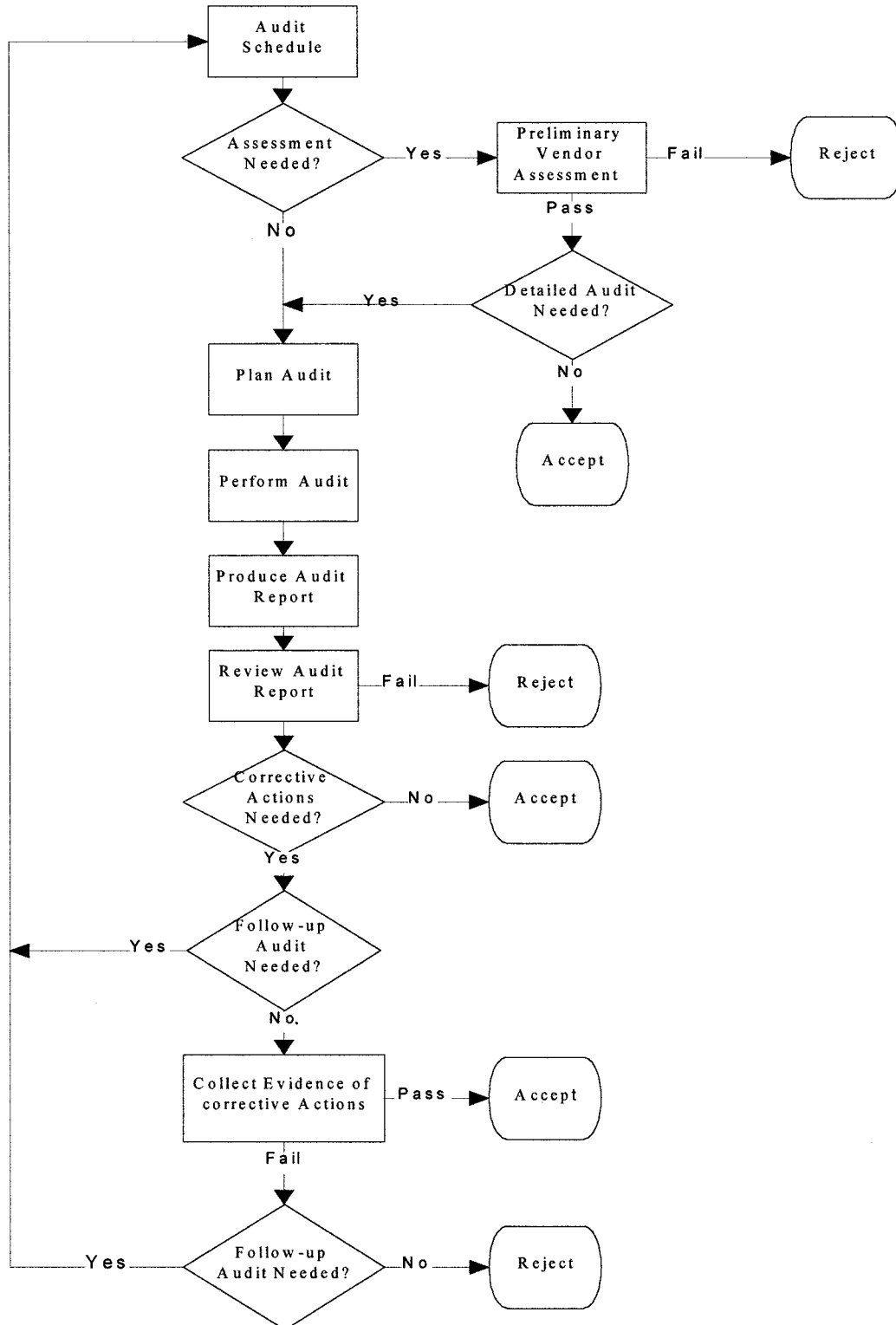


FIG. 2 Auditing Process

the customer selects. Each item should be ranked for the vendor's ability to meet that particular audit point. For example, a major discrepancy would indicate that the vendor had little or no compliance to the audit point/area. A minor discrepancy indicates that the vendor has some compliance. Both ISO 9000-3 and IEEE standards are detailed and may be used to create individualized checklists. It is important to remember that there are many different ways to accomplish compliance, and the auditor must take great care to understand how the audited company works and compare that to the standard instead of comparing it to his or her own quality system. See Appendix X1 for an overview of software items that should be investigated.

6.8 The organization should have established corporate auditing guidelines that describe in detail the procedures to which the vendor audit team shall adhere. These procedures should cover all activities from the initial vendor contact to the final meeting with the vendor. The overall auditing cycle can be divided generally into four stages: preliminary audit, detailed audit, follow-up audits, and surveillance audits (5). Each of these stages has its place within the overall auditing process.

6.8.1 *Preliminary Audits (Preaudit Activities)*—The goal of this stage is to gather enough documented evidence to determine if a detailed audit is required. The tool used to perform this auditing stage is typically a questionnaire. The questionnaire can be divided into the majors areas of concern, such as general corporate background information, sales information on the LIMS application (version-specific), vendor's software development life cycle (SDLC) procedures, and the product development history. Specifically, the buyer should request that the vendor supply, in advance, those standards, procedures, and plans that are associated with the LIMS application being investigated (1). The audit team should look for technical standards, manuals, or guides covering the following: development methodologies, software quality assurance practices, change control procedures, configuration management procedures, personnel training procedures, user support documentation, testing procedures, technical review practices, and security procedures (1).

6.8.2 *Detailed Audit*—When conducting these audits the organization should cover all aspects of interest relating to the application of LIMS. The validation team should plan their audit before actually performing it. The plan should establish the scope of the audit, who will be auditing, and the timing agreed to with the LIMS vendor. The audit notification should specify the purpose, timing, targeted system, scope, and the measurement criteria of the audit (1). The audit process itself can be divided into three major steps: the opening meeting, the review and inspection, and the closing meeting (5).

6.8.2.1 *Opening Meeting*—The opening meeting establishes the basic ground rules of the audit. Items to be addressed include, but are not limited to, introductions of everyone involved in this audit activity, the scope, purpose, agenda, schedule, location of the validation team meeting room, arrangements for accessing specific documents, and the signing of any confidentiality agreements by the LIMS vendor or the validation team members.

6.8.2.2 *Review and Inspection*—The audit team examines

the LIMS vendor's records and their practices in accordance with these documents. The goal is to establish documented evidence that the LIMS vendor operations show adherence to their quality procedures during the LIMS development. The audit team can perform the audit using a checklist based on the scope of the audit. A successful auditor should use a "show me" approach when auditing. The required depth for coverage of each audit item will vary, but in general the audit team should identify one or two items that they will cover in great detail (5). The audit team may want to hold daily wrap-up sessions designed to capture that day's activities. Any observations made and their impact on quality issues should be addressed at this time. The audit team also should begin developing a list for tracking follow-up action items (1). This guide will aid in creation of the final audit report.

6.8.2.3 *Closing Meeting*—The lead audit team member will list all observations that the team noted during their audit. This should include positive results as well as issues of concern (5). The vendor's response to the observations should be included in the documentation used to develop the audit report. The audit report is important because it serves as documented evidence of the audit and its findings, as well as the basis for determining corrective actions required by the vendor. As such, the report shall present the data accurately and objectively. Because it is sensitive, the audit report should be treated as a confidential document. The audit team should close the audit with the following next steps: (1) the lead auditor will produce an audit report, (2) the audit report will be reviewed by the audit team and management, who will devise a set of corrective actions, and (3) the LIMS vendor should be contacted by the lead auditor and devise a plan to implement the identified corrective actions (5). Individuals receiving the audit report will be identified. Expected response times to address the identified weakness shall be included in the audit report (1).

6.8.3 *Follow-Up Audit*—Follow-up audits review the progress made by the LIMS vendor on those items identified as areas of concern on the previous audit. The organization looking to purchase the LIMS has a few options they can pursue based on the outcome of the audit report. These options include the following (5):

6.8.3.1 Use the LIMS supplier unconditionally.

6.8.3.2 Use the LIMS supplier for certain LIMS products only, for example, specific versions.

6.8.3.3 Use the LIMS supplier only after specific corrective actions have been carried out.

6.8.3.4 Prohibit the use of the LIMS vendor.

6.8.4 If the LIMS vendor agrees to make the necessary corrective actions outlined in the audit report, the organization purchasing the LIMS should obtain the necessary documentation from the vendor for the changes made.

6.8.5 *Surveillance Audit*—These audits focus on weaknesses found during previous audits and any new features or LIMS products, for example, a new stability study module. These audits should follow the same general guidelines adhered to by the original audit. The frequency of these audits will depend on previous audit results and criticalness of the issues that need to be addressed.

6.9 The validation team, in concert with management,

should establish an action plan for those instances in which the LIMS vendor refuses to allow an audit. The LIMS validation team must remember that it can not test quality into the system. Further, the amount of testing is proportional to the level of risk the organization will take for implementing the LIMS. Options available to the organization include the following:

6.9.1 The organization accepts the business risk and performs a much greater degree and depth of validation for the LIMS.

6.9.2 The organization rejects the LIMS vendor and moves the selection process towards alternate LIMS vendors.

7. Validation of LIMS Installed at Customer Site

7.1 The customer shall validate their use of the LIMS, independent of any vendor audit, in the operational environment in which the LIMS will be residing. The fact that a vendor's LIMS development process has been validated by the vendor or other organizations has little bearing on validating the organization's LIMS application. Further, the fact that a vendor's LIMS software has been validated by one of their other customers does not obviate the need for an organization to validate their implementation of the application.

7.2 As key functional requirements are identified and evaluated during the product evaluation phase, their results should be recorded. These results may be used in development, execution and documentation of the official LIMS test protocols. Any testing done during development of the LIMS test protocols or overall validation plan should be further refined once a specific LIMS has been selected. It should be noted that the level of testing and evaluation done during the evaluation and selection process generally will not contain enough detail to replace the test protocol used in the validation plan documentation.

7.3 The LIMS validation team may begin to identify additional resources to test the LIMS. Any new individuals selected should be familiar with the laboratory's requirements and its operation. Further, they should be knowledgeable about cGMP, GMP, GLP, GALP, or other requirements that the laboratory shall follow.

7.4 The LIMS typically is delivered as an empty database, that is, devoid of site-specific data. Configuration data and fixed laboratory information must be entered before the system can be validated. At this point, the organization starts to model their laboratory practices in LIMS. This includes test and workstation definitions and laboratory and customer personnel data. It should be noted, that during this step the laboratory may encounter additional functional requirements that were not captured initially. If the organization chooses to implement such functionality, the LIMS requirement document shall be revised to reflect these changes. Further, during this step the organization may uncover requirements that the LIMS cannot meet. The organization should document these facts and include what actions, if any, they will take to solve this problem. There are several strategies that can be used to validate a LIMS. These include, but are not limited to, the following:

7.4.1 Configure the LIMS specifically for testing with only enough configuration data to permit testing. In this case, the test system is identical to the production system, specifically, it

is functioning in the same operational environment as the production system. Generally, this means that it is operating on the same computer on which the production system will reside. The configuration used in the test system shall exactly match the production system. Specifically, all reports, entry screens, queries, etc., must be identical. Furthermore, all features that are to be used in the production LIMS shall be checked for proper operation in the test system.

7.4.2 Configure the LIMS for regular operations, then isolate it from normal service while testing it. A system configured for use is called the production system. This can be accomplished by copying the production database to the test system. The LIMS program executables are the same, for example, the validation data may be part of a separate set of database tables that use the same program executables as the production LIMS, or the validation data may be part of different data group that uses the same database tables and executables as the production LIMS. The difference is in the sample data tables. If there are no problems, this approach saves time. The LIMS does not have to be configured twice, once for testing and again for production. If problems are found, partial or complete reconfiguration may be required after repairs are made. Documentation verifying that the production system is equivalent to the test system shall be provided, and the data generated during the validation process should be retained and identified as validation data.

7.4.3 A separate computer system may be used for testing.

7.4.3.1 The separate computer may be configured specifically for validation, as in 7.4.1, or it may be a copy of the production system, as in 7.4.2.

7.4.3.2 If a separate computer is used, it should have identical hardware, software, and operating system. The operating environment shall be identical to the one used for the production system. Instrument interfaces may be difficult to install on such a test system, but if they are part of the production system, they must be part of the test system as well. Ultimately, the test system could provide backup hardware for the production system.

7.4.3.3 The production and test systems may exist on the same computer, if it is sufficiently powerful, running independently. In this case, both software systems may have access to the instrument interfaces.

7.4.3.4 A subset of tests is needed when the test system is converted to the production system. These tests are used to confirm that the system still functions properly in production mode. No artificial data needs to be loaded into the active system. This subset of tests may consist of vendor-supplied diagnostic routines and little more, as long as they reliably test all parts of the proposed system. While some vendors supply these types of tools, many do not. There is no standard for their construction and execution. The use of such tools should not be the only means of testing the LIMS, but rather augment a more rigorous set of test protocols. In some cases the organization may require the tools themselves be validated prior to their use.

7.4.4 Parallel testing may be used. For a new LIMS, the manual systems can be used simultaneously with the LIMS and the results compared. If the new LIMS is a replacement, both old and new systems can be used in parallel for some period of

time to compare them. The existing validated system is the production system, while the new LIMS is the test system. Validating interfaces to instruments are an issue with a parallel testing approach, since they cannot usually be connected to both systems at the same time. In this case, the organization shall develop an approach that allows for the testing of these interfaces. The organization may want to connect these interfaces to the system undergoing validation after all other tests have been executed and just prior to the development of the final validation report. Another approach is to incorporate these interfaces as their own validation project conducted after the initial validation has been concluded.

7.5 *Response to Errors:*

7.5.1 Error handling and acceptance criteria shall be defined and described in the validation protocol and followed during the testing and reporting of results. The definition shall include criteria to be used to assess severity of errors.

7.5.2 Critical errors, such as system crashes or fatal errors, located during validation tests should be corrected or repaired immediately, before additional testing is done. Often the correction of such errors requires that most or all of the validation tests be run again. These are errors for which there is no work-around. These errors seriously threaten the integrity of the LIMS data.

7.5.3 Noncritical errors should be accumulated during the validation tests. When testing is complete, the team may decide these errors do not compromise the integrity of the information. These are errors which could result in the possibility that unacceptable result data would be accepted by the LIMS. There may be an acceptable work-around for such errors.

7.5.4 The validation team may wish to use an error grading system that helps to take action when errors are encountered. Each error would be identified by grade, and a decision would be made on what follow-up, if any, is necessary. The following are examples of grades and the errors that fall into those grades (6):

7.5.4.1 *Grade 0*—Typographical errors and other errors not related to the computer system.

7.5.4.2 *Grade 1*—Minor errors such as the use of upper and lower case letters used in fields not constructed for them.

7.5.4.3 *Grade 2*—Tolerable errors that must be communicated to the vendor.

7.5.4.4 *Grade 3*—Major errors that must be immediately reported to the vendor and the QA manager. All validation efforts should be suspended until QA has discussed the problem.

7.5.4.5 *Grade 4*—Disastrous errors such as relational errors in the database. These are reported the same as Grade 3 but the validation effort should be aborted. QA could still decide that the effort continue after thorough discussions.

7.6 *Standard Operating Procedures (SOPs):*

7.6.1 SOPs are necessary for validation and ongoing operation of an organization's LIMS. These documents cover several areas, from the operation of the LIMS application through to the hardware on which the application resides. The SOPs formalize the procedures used to maintain the LIMS in a validated state by describing specific procedures to be followed. These procedure help ensure that the organization

maintains a quality operation. SOPs are detailed in 11.4.

8. Validation Plan Design

8.1 The validation plan provides the overall direction of the validation process. The validation plan includes, but is not limited to, the overall objectives, a description of the system, any test boundaries or assumptions under which the validation team will be operating, the participants' responsibilities, and any general instructions for the execution of installation qualification (IQ) or operational qualification (OQ) test protocols. The validation plan needs to include a listing and description of all software and hardware components. Sometimes software modules associated with the LIMS are changed by the installation of other software. These changes could be from operating system upgrades, an upgrade to the LIMS, or other unrelated software. Further, the addition of hardware components, video cards, modems, sound cards etc., and their associated software can affect the initial LIMS validation state. The detailed listing of software and hardware components associated with the LIMS is essential as it makes up the LIMS initial configuration and describes the beginning state from which all change control is based. All test protocols for both the IQ and OQ of the associated hardware and software components are included in the validation plan. The last part of the validation plan is the signatures of the individuals responsible for ensuring that validation plan meets the organization and regulatory requirements. Typically, these signatures include the QAU validation manager, a laboratory manager, LIMS manager, and others.

8.2 IQ testing should be based on manufacturer's specifications, or recommendations, or both. Application-specific configuration will be verified as part of the IQ/OQ testing.

8.3 Vendor-supplied diagnostics can be used as part of IQ/OQ testing. IQ/OQ protocols based on vendor-supplied diagnostics shall include step-by-step verification of diagnostic procedures, recording of all results, and acceptance criteria for each result.

8.4 IQ/OQ protocol documents and test results should be produced for all hardware and software used with the LIMS, that is, operating system, database, report generators, statistical packages, network, connected instruments, computers including terminals, PCs, clients and servers, printers and plotters, bar code readers, etc. If the LIMS application is being loaded on an existing computer system, the original hardware IQ documentation may be used.

8.5 A suggested format of the IQ/OQ protocol document can be found in Appendix X2.

9. Test Protocol Design

9.1 Each organization should determine which LIMS features may attract the largest amount of attention by the auditing agencies. The organization shall determine what level of risk they are willing to accept. To validate every feature is too costly in terms of resources and time. McDowall has suggested that the organization divide the LIMS functions into one of the following three categories: must validate, should validate, and could validate (2).

9.1.1 The validation test protocols need to identify critical LIMS functions that will be tested. Critical LIMS functions

should be based on core functions and the intended use of the LIMS application. Rationale should be provided for not testing portions of the LIMS.

9.2 The development and execution of test protocols (TP) takes the largest amount of time in the validation effort. This fact often is overlooked when the validation project plan is developed. Many factors affect TP development and execution. First, good familiarity with the new LIMS and how it operates are essential. The less familiar the user is the longer it takes to develop detailed TPs. The validation team should build sufficient time into the project schedule for the personnel developing TPs to develop familiarity with the new system. A second factor affecting TP development is how long the TP developers have to focus upon the validation project. Not focusing enough on the TP development effort will add a significant number of additional months to the validation project. The execution of the TPs also is affected significantly by focusing the testers on the execution of the TP. A third factor affecting TP development is the number of resources available to work on the TPs. Last, the experience level of the individuals writing and executing the TPs will affect the time necessary for these activities. If possible, the organization should have at least one experienced individual working with those developing and executing the TPs.

9.3 The number of TPs necessary for validating the LIMS depends on the complexity of the LIMS and the level of detail required to adequately test the key features. TPs can be as simple as one or two lines of execution instructions or as complex as several hundred lines. The level of complexity will depend on the direction that the organization takes in the design of their TPs. Each organization should have an organizational SOP that describes how TPs are to be designed. The design can be as simple as very high level and general instructions on what testers should do and what they should expect as their acceptance criteria. TPs designed in this manner generally require the tester to write down, in detail, what they have done. At the opposite end of the spectrum are those TPs that instruct testers step by step on what to do. TPs designed in this manner typically require the testers to answer yes/no or true/false to the acceptance criteria. In either case, complex TPs can take several days to execute and document. The detail captured by testers for each TP should be sufficient enough to ensure that the LIMS function or the process being tested is under control. See Appendix X3.

9.4 In addition to execution of the TP, the validation team shall incorporate the time necessary to review TP results and to solve any identified problems. The review process can take almost as long as the execution of the TP, if the test is extremely complex. The time necessary to carry out this validation step often is underestimated. The review of each TP is necessary to ensure that the content makes sense and that it adheres to GMP documentation requirements. Specifically, all errors should have a single line drawn through them; the tester should initial, date, and give a reason why the word or group of words were crossed out. In some cases the reviewer may be responsible for deciding if the TP has met its acceptance criteria successfully, and thus, either passes or fails.

9.5 The validation team should address in the validation

plan how they will handle failed TPs. This shall be addressed before the testing begins. They also should address early on how they will allow changes to the TPs after approved by the QAU. There are times when testers will need to make changes to the TP during the execution phase of a TP. Testers should be provided a way to incorporate these changes into the existing TP. The procedure shall be approved by the QAU and incorporated into the validation plan. It is essential to give testers freedom to further design and follow additional test steps when executing the TP. This freedom allows them to explore why a particular step or set of steps did not meet its acceptance criteria. Without this freedom the entire validation project can be delayed.

9.6 All TPs shall be designed to test the given LIMS feature or function. The actual design of TPs will vary from organization to organization. The designer of the TP may wish to include any or all of the following in the design of the TP:

9.6.1 *Test Protocol Header Information*—This section contains the name of the corporation using the LIMS, the department name of the LIMS owner, date the TP was designed, statement if the TP is for IQ or OQ, TP revision number, and what system is being tested (for example, ABC LIMS Version 7.1).

9.6.2 *Test Protocol Identification Number*—Each TP should have a unique identification number. This number is only unique to the associated validation plan for the TP.

9.6.3 *Purpose*—What the TP is designed to test. For example, the purpose is to verify that new users can be added, modified, or deleted from LIMS.

9.6.4 *Requirements Under Test*—These are the functional requirements that are being tested by the TP. The TP may be designed for more than one functional requirement. Any functional requirement that was not included into the validation plan should not be included in the development of the TPs.

9.6.5 *Special Needs/Requirements*—This section lists special items that are needed to execute the TP, including specific skills the testers must have or links to other test protocols or other applications.

9.6.6 *Test Step Procedures*—Each test step should include a step number, a test procedure, and acceptance criteria for that step. Further, the test steps should be divided into and have a set of test steps for each of three categories: normal testing, stress testing, and robustness testing. Normal testing steps test the LIMS function using all common user commands. Test steps that test the function at its boundaries are stress testing. An example would be entering 20 characters into a 20 character field. Robustness testing represents testing the feature outside its boundaries. For example, a user's password may only accept character and numbers, so testers are instructed to enter special characters or punctuation characters for a newly created user's password. Testers shall identify if the test step passed or failed acceptance criteria. Typically, this is a simple yes/no statement.

9.6.7 *Comments Section*—This section is used by testers to enter their comments on any unexpected results obtained while executing the TP. Users also can capture how these unexpected results were resolved.

9.6.8 *Tester Sign-off*—The tester should sign and date the

TP at the end of the testing process. If the TP covers several pages, the tester only should sign and date the page when they have completed the test steps on that page. In some organizations testers are responsible for determining if the TP passes or fails. If the TP fails, testers should document in the comments section why the TP fails. If they have identified a possible resolution, testers should document this as well.

9.6.9 Reviewer Sign-off—The TP reviewer should sign and date the TP only after reviewing the data and concurring that the TP has been completed. If questions exist, the reviewer should not sign the TP until the questions are answered. In some cases, it is the responsibility of the reviewer to determine if the TP passes or fails. If the TP fails, the reviewer should use the comments section to explain why. The reviewer should not make changes to the document. If changes need to be made, the original tester should be contacted to make the changes.

9.6.10 Attachments—All attachments that are part of the execution of the TP should contain the following pieces of information: the TP identification number, the step number, initials of who created the attachment, and the date the attachment was created. Furthermore, the tester may want to highlight or explain certain items on the attachment. Any handwritten item requiring change shall follow the same criteria as the TP and include a single line through the item, initials of the person making the change, date, and a reason for the change.

9.7 As TPs are finished they should be forwarded for review. After they have been reviewed and signed, they should be given to the validation team leader. This will facilitate the development of the final validation report. Furthermore, if there are identified system outages to be addressed, the validation team leader can start to address these issues without impeding the progress of the testing team members.

9.8 The validation team members can use several approaches to design and test their LIMS implementation. The test team may wish to include the following additional approaches in the TP design:

9.8.1 Running vendor supplied diagnostic tests (supplied tools/test set may need to be validated prior to their use).

9.8.2 Running automated testing tools, if available, for that particular LIMS (supplied tools/test set may need to be validated prior to their use).

9.8.3 Log results manually along with the LIMS for a given time period, and compare the results.

9.8.4 If the LIMS has telephone access, test the associated telephone security measures thoroughly.

9.8.5 Introduce errors deliberately, and determine if the system properly identifies and rejects them.

9.8.6 Stress the system by artificially and completely filling it with data, or running many activities at once.

9.8.7 If operating in a windows environment, for example, open all the windows at once.

9.8.8 Schedule heavy loads.

9.8.9 Test security by trying to break in or use prohibited functions. Look for “back-door” entry points.

9.8.10 Try to abort an input to see if the system behaves as specified.

9.8.11 Visually observe interfaces and other aspects that

produce a discernible action.

9.8.12 Vary load sequences of automated instruments.

9.8.13 Review every output screen for completeness, correct data in every field, and adherence to specification.

9.8.14 Use screen capture or keystroke capture techniques to review system operation.

9.8.15 Test all event triggers by forcing them to happen. Include scheduled events and, as much as possible, exception events.

9.8.16 Disconnect the power to interfaced instruments, servers, and other parts of the system.

9.8.17 Use protocol testers for network performance, including adherence to protocol, timing, and data integrity.

9.8.18 Use instrument simulators, if available, to test exceptions and errors in interfaces.

10. LIMS Operation

10.1 Once the LIMS has been validated, operational system maintenance begins. At this stage the validation team members can be disbanded. From this point on those responsible for daily operation have the responsibility to maintain it in a validated state. The critical issues that face the organization, and more importantly the LIMS Manager, are as follows (**2, 7**):

10.1.1 Configuration Management—The purpose of configuration management is to ensure that any changes to the hardware, firmware, network, LIMS executable code, or any other component that was part of the initial LIMS validation process are identified and controlled. All LIMS applications and the hardware platforms on which they reside will change. It is essential that the organization controls and documents these changes. The procedures for managing these changes fall under configuration management. Configuration management starts during the development and execution of the LIMS hardware and software IQs and OQs. At that time the validation team established a listing of all hardware and software in the LIMS setup and configuration, including part numbers, release numbers, serial numbers, and software version numbers. All these items together make up the initial LIMS configuration. This is the baseline for configuration management. Additional items that should be considered are DLLs used by the LIMS application and any associated software. In this case the user shall track DLL names, dates installed/written, and versions. This is crucial to ensure that no other software changed the DLLs used by the LIMS. The objective is to show that the organization is in control of the LIMS. The organization must show that once the initial configuration has been established, all changes to that configuration are authorized, tested, and documented. It is essential that if responsibilities for the various parts of the LIMS configuration are shared by other organizational groups, for example, information services, maintains the network infrastructure, they must be aware that they cannot make changes to their area of responsibility without first checking with the LIMS manager and the organization.

10.1.2 Change Control—Changes to the LIMS are a fact of life. It is imperative that all the implemented changes go through change control. Change control involves several steps: change request, analyze impact, review/approve, implement,

and validate (2). The organization should have an SOP describing the procedures to be followed by those requesting the change. Furthermore, the organization should have a change control board that reviews all proposed change controls. The membership of this board will vary from organization to organization. Key members include QAU personnel familiar with validation of computerized systems and personnel from the various business areas. The board's role is to review all proposed changes and determine if the approach adheres to both the organizational and regulatory requirements. In addition, the board reviews and assesses the impact of the change on operations. When preparing to make the decision to request a change to the LIMS, consider the following: will the changes provide big enough benefits to offset the time and resources needed to revalidate the LIMS; what other systems will be impacted by the change; how much time will be required to successfully implement the change; what resources shall be made available to implement the change; and, what effects will not implementing the change have on both the laboratory and the organization. In addition, LIMS applications that reside on a PC-based server must be controlled carefully because a user's PC may use different DLLs and update versions of the LIMS DLLs in a noncontrolled manner.

10.1.3 *SOPs*—See 11.4 for details on SOPs.

10.1.4 *Operational Log Records*—The organization should use log books to document the proper ongoing operation of the LIMS. Records can be as simple as a predefined form that is filled out and filed, to as complex as a specialized recordkeeping application. In general, the organization's goal in utilizing these logs is to show evidence of control over the LIMS operation. In some cases, these records can be used to show trends in the performance of software or hardware components. Operational logs should cover the following areas (7):

10.1.4.1 *Backup of Data Log*—This document provides evidence that the LIMS application is being backed up in accordance with the organization's SOP. This log should contain, but not be limited to, such items as who performed the backup, the time of the backup, to what extent the LIMS was backed up (for example, full system backup including the LIMS and the operating system it resides on or partial where only the LIMS data directories are backed up), where the tapes are stored, and if the backup was successful.

10.1.4.2 *Error and Error Resolution Log*—The organization should maintain an error and error resolution log. This log helps to determine if there are trends in the errors, as well as provide evidence that errors are addressed as they are captured. The organization shall determine if it can resolve the error. If there are ways to fix it in-house the organization should contact the LIMS vendor. In either case, the organization should state what they have done to resolve the error. When errors are identified as bugs, the organization should obtain a time commitment from the vendor for resolving the bug. This data should be entered into the log. If the LIMS vendor states that the bug has been fixed in an upgrade of the software, the organization should record this data. Revalidation of the fixed bug should be a key area addressed in the revalidation effort upon implementation of the upgrade.

10.1.4.3 *Hardware Maintenance Logs*—These logs deal

specifically with the hardware components of the LIMS, including any associated networking peripherals. The user should track the serial/part number of the component being replaced, the manufacturer's name of the replaced board, if known, the serial/part number of the new part, the manufacturer of the new board, the printed name and signature of the serviceman, the date the replacement took place, a reason why the component was replaced, and other data that will be helpful to debug problems later (6).

10.1.5 *Revalidation*—All changes must be assessed for their impact on the validation of LIMS. Changes that impact the integrity or accuracy of data in LIMS require the LIMS to be revalidated. The revalidation effort need not be as major as the original validation effort, assuming the changes are minor in nature. The effort involved can be shortened by using some of the original TPs from the initial validation effort. The design and amount of documentation will vary from one organization to the other, as each has their own change control SOP. The user can design a shortened version of the original LIMS validation plan.

10.1.6 *Periodic Audits*—The organization should conduct periodic audits of their LIMS. This audit verifies that the LIMS complies with the established policies and procedures. These audits, typically, are not carried out by the LIMS or laboratory personnel. Generally, they are handled by QAU personnel. While these audits are not part of the LIMS manager's direct responsibilities, this person does not have responsibility for maintaining the LIMS in a validated state. The areas of greatest concerns for those auditing include: security procedures, error logs, maintenance logs, change control procedures, training records, operational logs, if used, back-up and recovery procedures, disaster recovery procedures, and documentation management procedures.

11. Documentation

11.1 There are many types of documents associated with validation. Each document must be version-controlled to ensure that users can identify the specific versions they used in their validation process.

11.2 The validation documentation should include, but is not limited to, the following (8):

11.2.1 *Validation Plan (see Appendix X2)*—The master plan that outlines roles, responsibilities, and the course of action to be followed by the validation team.

11.2.2 *Functional Requirements*—Contains the requirements the LIMS is expected to meet. This is a key essential document used in the validation process of LIMS (see 11.5 for more details).

11.2.3 *Prevalidation Systems Acceptance Test Documents*—This document can be used to determine the validity of the LIMS, based on the functional requirements document. The difference in this case is that the functional requirements are not tested as stringently as in a normal protocol testing environment.

11.2.4 *Complete System Specifications* (database schema, user interface designs, wiring diagrams, etc.).

11.2.5 *IQ, OQ Protocol Documents (see Appendix X2)*—These documents comprise the bulk of the validation activity. The goal in each case is to design a test protocol that tests one

or more functional requirements. Each set of tests shall contain what the user considers the acceptance criteria for that test step.

11.2.6 *Test Protocols (see Appendix X3)*—These are part of the IQ/OQ document. Each test protocol will test one or more functional requirement. All test protocol attachments, that is, hard copies of screen layout, paper reports, etc., all become part of the IQ/OQ documentation package.

11.2.7 *SOPs*—See 11.4.

11.2.8 *LIMS System Manual*.

11.2.9 *Final Validation Report-Qualification Report (see Appendix X4)*—This report completes the validation plan, which has been executed. It shall document any system limitations identified during the execution of the associated testing protocols. It must record the formal decision to accept the system with sign-off. It should note if acceptance is for limited operation because some tests failed. It should document how the identified limitations are to be handled.

11.2.10 The following is other miscellaneous supporting documentation the user may want to include (8):

11.2.10.1 All purchase orders associated with the LIMS application, hardware, software, consulting services, etc.

11.2.10.2 The vendor audit status report.

11.2.10.3 The escrow agreement for the LIMS source code.

11.2.10.4 Source code maintenance requirements for any in-house customization work accomplished.

11.2.10.5 Structural testing documentation for the source code.

11.2.10.6 Service contract and support agreements.

11.2.10.7 User and LIMS administrator training records.

11.2.10.8 The LIMS implementation plan.

11.2.11 The user should have the following additional documentation for customization work (8):

11.2.11.1 System development life cycle.

11.2.11.2 Programming standards and conventions document.

11.2.11.3 Configuration management records created during the development of the system.

11.2.11.4 Documented evidence of structural testing on the source code.

11.2.11.5 Procedure to release the system from development phase to validation phase.

11.2.11.6 Documented evidence verifying the adherence to procedures.

11.2.11.7 Procedure to address problems found after the system is implemented.

11.3 *Documentation Strategies*—Several schemes exist for tracking progress during validation.

11.3.1 All activities should be documented, especially tests that fail and must be subsequently repeated.

11.3.2 A logbook may be kept, where all tests are recorded chronologically along with their results and dispositions. Each entry should record when the test was done, who did it, what results were obtained, and how problems were resolved.

11.3.3 There may be a protocol opened when each test is begun. If a test fails, the protocol must be closed with unsatisfactory results. After repairs, a new protocol for that test may be opened and the test repeated.

11.4 *Standard Operating Procedures That Are Specific to*

the Operation of the LIMS:

11.4.1 SOPs shall be in place to ensure that the organization has well defined procedures. The number, the design, and the focus of SOPs will vary considerably across organizations and LIMS applications. For example, if the LIMS runs on a server versus a stand-alone PC, the organization will need a different SOP for each. The following list gives general SOPs that organizations may wish to develop. The user of this guide should not assume that the list below is complete or required.

11.4.2 *SOP on SOPs*—Describes how SOPs shall be designed, including specific required sections and types of information, who has responsibility for what, and a numbering system for all corporate SOPs.

11.4.3 *Validation of Computerized System*—This is a corporate level SOP that describes the ins and outs involved in the development of a validation plan for a computerized system. This SOP should be targeted at a specific class of computer systems.

11.4.4 *Training*—Covers who shall train, who shall be trained, what is to be covered, and version control of training material. Include who has responsibilities for informing trainers and trainees. The extent of the training depends upon what access the person needs in the system. Changes in access, or responsibilities, or both, may require more training. The training should include theoretical and practical use of the system, and how to document training records, etc.

11.4.5 *Backup and Restore*—Includes procedures for backup, use of the journal log, off-site copies, policy on keeping earlier versions of the database (for missed errors), and restoration procedures.

11.4.6 *Disaster Recovery*—This SOP covers those procedures that should be followed in case of major disasters, such as fire, flood, sabotage, and major system or equipment failures. These procedures include defining the interim laboratory operation for how the business will be conducted during the loss of the LIMS. Further, this SOP should cover how to resume business once the LIMS is operational again.

11.4.7 *Security*—Includes system policy, corporate policy, and enforcement policy. Minimum password policy should be specified, such as maximum lifetime of a password, avoidance of trivial passwords, who assigns passwords, expiration dates, maximum number of tries before lock-out, and access logs. Policy should include when security reviews are conducted, who performs them, who reviews the results, how security policy revisions are made, and who assigns responsibilities and rights. The need for securing physical access to the system also should be incorporated into the SOP. Other procedures may exist, such as keyboard locking, biological identifications, etc. These should be addressed as necessary in this SOP.

11.4.8 *Change Control*—Includes version identification, maintenance of static data still needed to document older results, change policies, sign-offs required, retesting and re-validation needed, and the documentation required. The effect of changes on more general information should be considered, such as research studies, material specifications or formulations, analysis techniques, and method parameters. Change control is needed any time LIMS performance may be affected, for example, changes to the operating system, the local area

network, the database engine, the server hardware or software, the LIMS software, any interface, all major repairs, and many minor repairs.

11.4.9 LIMS Operation—This SOP should include the operating policies and responsibilities of each user from the LIMS manager down to the end user. If the LIMS operates over a local area or wide area network, these functions may be under different management. The LIMS manager and the organization shall ensure that the SOP addresses these issues in order to provide proper support for their LIMS. Further, the SOP shall address other LIMS items, such as start-up and shut-down procedures, ownership of supplies, routine problem resolution, etc. Some organizations have specific job descriptions for their personnel. As such, these job descriptions may be referenced in this SOP.

11.4.10 Maintenance—Includes who did the service, when it was performed, what was done, and what documentation that should be created. This should apply to both routine and unscheduled maintenance. Documentation is also required for who approved on completion of service, what retesting was done, and if necessary, what level of revalidation was performed and documentation required. In some industries, repairs shall follow the organization's established change control procedures.

11.4.11 LIMS Usage—Emphasizes responsibilities. This may refer to the manual(s), if they exist, but a user handbook or manual should not be written in this SOP. The understanding is those using this SOP will already be trained and know how to use the LIMS. It is not necessary to rewrite the SOP if the system changes appearance, for example if “log the samples by batch” appears instead of, “Log - <return> <return> <down-arrow> highlight “by batch” and <return>.”

11.4.12 Error Handling—Addresses how LIMS errors are to be handled. This can be a separate SOP or it can be incorporated into the LIMS Operational SOP. In either case, LIMS errors should be documented. Further, the SOP should describe the course of action that LIMS users should take when they encounter a problem.

11.4.13 Building Static Data Templates—Includes nomenclature to be used in the design of the various static tables. For example, this SOP may state that all test methods will be coded into LIMS using a specific method numbering system or that all test result templates will track certain data elements (test initials, tester lab notebook number, etc.).

11.4.14 Instrument Interfacing—Describes how new instruments are connected to the LIMS, how they are tested, how they are validated, and how they are to be used.

11.5 Functional Requirements Document:

11.5.1 This is a key document in the validation process of LIMS. This document is used to ensure that the LIMS does what it purports to do and will continue to do so once validated (9). This document should outline the business and regulatory needs and policies. While the development of the functional requirements document is the responsibility of the LIMS project team, it is essential that the LIMS validation team know and understand what should be contained in this document.

11.5.2 The functional requirements document puts into common language the required LIMS functionalities and LIMS

performance issues (9). It is a communication device for conveying requirements to the LIMS vendor. In addition, the functional requirements document aids in the development of the qualification documents and their associated test protocols. When the test protocols are executed they will be compared to requirements detailed in this document.

11.5.3 This document should contain detailed information that covers the system description, systems constraints, vendor-related requirements, detailed system information, general systems performance requirements, system implementation and other operational requirements, and other documentation for custom-developed software (9).

11.5.3.1 The system description should include, but is not limited to, a main purpose, essential features system environment and associated interfaces critical to the system operation, and projected completion schedule (9). Additional items also include a glossary of terms, acronyms, and abbreviations specific to the LIMS and references to other corporate standards.

11.5.3.2 System constraints should include, but are not limited to, a preferred platform for the hardware and software, system interfaces to other systems (instruments, LAN, WAN, etc.), future system expandability requirements, environmental requirements, life expectancy of the system, scheduling requirements, source code availability, and maintenance requirements (9).

11.5.3.3 Vendor-related requirements should include, but are not limited to, vendor audit requirements, vendor systems deliverables (hardware, source code, etc.), user manuals, training manuals, vendor service deliverables for bug support, maintenance, and training (9).

11.5.3.4 The overall objective and task requirements are outlined in the detailed system information document. Systems functionalities should be divided into three main blocks: input, processing, and output requirements. Each block should describe subfunctionalities specific to each area. For example, an input subfunctionality would include requirements for migrating data from the current system to the new LIMS (9).

11.5.3.5 General system performance requirements should cover the expected response time for specific tasks using the system, expected maintenance downtime, error handling requirements during start-up and shut-down, and backup and recovery requirements (9).

11.5.3.6 System implementation and other operational-related requirements should cover support and service needs, supporting documentation, such as user's manual, an administrator's manual, as well as archival and data-retention requirements (9).

11.5.3.7 For customized LIMS, the functional requirements document should include the systems development life cycle used, the SDLC phase and required deliverables for each phase, the quality assurance plan, documentation for prototyping, requirements for configuration management and items to be included, requirements for change control, required testing and documentation to be performed during development testing, and requirements for any additional documentation for post-implementation activities (9).

12. The Quality Assurance Unit (QAU)

12.1 The QAU conducts or assists quality assurance activities in the interpretation of the various regulatory requirements. This extends to issues relating to the validation of computerized systems, such as LIMS. Additional roles that the QAU has that affect the LIMS and its validation include vendor audits, review and final sign-off of the LIMS validation plan and final validation report, ongoing monitoring of the LIMS via audits and change control requests, and assistance in the development and maintenance of the LIMS related SOPs.

12.2 QAU personnel who are responsible for the validation of computerized systems should have a sound technical understanding of both the regulations and computer technology. QAU individuals can use in-house or industry training course, read technical literature on this subject, or work in conjunction with an experienced computerized systems validation QAU expert to obtain the required level of expertise. It is imperative that QAU personnel stay abreast of the technology changes.

12.3 The validation team should have a QAU member at the start of the project. Early involvement will aid validation in many ways. First, the QAU representative can gain an understanding of the LIMS project. Second, they can indicate which regulatory requirements the team needs to work against. This allows the validation team ample time to design these requirements into the validation plan versus reworking the issues later in the project. Third, as the validation plan is created the QAU representative can review and suggest corrections to the document. When QAU is included from the start of the project, the validation team will be better able to meet the project timelines.

12.4 The QAU also is responsible for the ongoing evaluation of the LIMS. They should periodically review procedures for operating the LIMS. The goal is to ensure that the proper controls are used. The LIMS manager and others should be aware of the need to follow the outlined procedures. Areas that draw the most QAU attention are those that directly affect data integrity, its accuracy, or its security. QAU representatives will be checking for prescribed change control procedures and documentation after any changes. Error and operational logs must be kept up to date. This can be a monumental effort if the responsibilities for maintaining the LIMS is spread across several organizational groups (laboratory, IS server operations, IS database manager, etc.).

12.5 Representatives from the QAU may be involved in the following LIMS project steps:

12.5.1 Project definition.

12.5.2 Functional requirements.

12.5.3 *Investigation of Vendors*—The QAU may perform a vendor audit to ensure good software practices were followed while developing the LIMS. They should at least review the vendor audit report if they did not participate in the actual audit.

12.5.4 *Vendor Negotiations*—The QAU may be involved as functional requirements are added, dropped, and revised. This often occurs to resolve differences between ideal requirements and available features in commercial systems.

12.5.5 *Vendor Selection*—A revised validation plan should be part of the contract with the vendor.

12.5.6 *Validation Phase*—The QAU will monitor compliance to the validation plan and review conclusions and approvals. This includes authorizations of the validation IQ/OQ protocol, including the TPs prior to execution. Further, it includes authorization of the qualification report after the execution of the IQ/OQ protocol.

13. Management's Role

13.1 Management's key role is to commit and support the appropriate resources to the validation project, which include both labor and money. Furthermore, management shall help define the level of risk the business is willing to accept. Management should strongly support the quality assurance unit and their involvement from the start of the LIMS project, as well as ensure that all necessary SOPs are in place and that those responsible for the LIMS validation project have received the necessary training to conduct their job.

13.2 Management may act as the project sponsor with overall project sign-off responsibilities, which includes responsibility for being involved in all major decision points during the validation project. Management needs to ensure that the proper resources are allocated to maintain the LIMS and any associated systems in a validated state. Management may be called upon to resolve issues across organization boundaries and to ensure that all those involved in the LIMS daily operation are aware of the organizational and regulatory requirements.

APPENDIXES

(Nonmandatory Information)

X1. VENDOR ASSESSMENT INFORMATION

Vendor Name:	Vendor Contact:
Product Name:	Evaluation Date:
Phone #:	FAX #:
Audited By:	

AUDIT ITEMS	LEVEL OF EVIDENCE 1=NONE, 2= SOME; 3=COMPLETE	COMMENTS / REFERENCE DOCUMENTS
Software Development		
Review standards/guides/procedures (software Life Cycle)		
Evidence of technical reviews		
Evidence of standard coding practices		
Development documentation exists for:		
Requirements phase		
Design phase		
Source Code phase		
Testing phase		
Installation and Checkout phase		
Operation and Maintenance phase		
Source Code Module Structure includes:		
Header Information		
Program name		
Program description		
Development date-time stamp		
Developer(s) name(s)		
Inputs		
Outputs		
Program and subroutine calls		
Data parameters		
Revision-control sections		
Annotated Code		
Testing		
Structural Testing (tests the code)		
Functional Testing (tests the design)		
Validation Test Data Sets		
Documented Test Results/Exceptions		
Software Maintenance		
Customization of Standard Software		
Revision control		
Synchronization between vendor/client		
Configuration Control		
Software Quality and Control Issues		
Software Quality Assurance Group		
Security Controls to Access Software		
Error-Detection/Problem-Resolution Procedures and Outcome Records		
Distribution Controls		
Records Retention Schedule		
Disaster Recovery		
Software Quality Plan		
Software Functional Requirements		

FIG. X1.1 Vendor Assessment Information Sample Form

AUDIT ITEMS	LEVEL OF EVIDENCE 1=NONE, 2= SOME; 3=COMPLETE	COMMENTS / REFERENCE DOCUMENTS
Software Quality and Control Issues Continued		
Software Development Life Cycle Defined		
Software Developer Training Requirements		
Software Validation Plan		
General Facility Issues		
Security Controls for Building Access		
General Cleanliness		
Organizational Quality and Control Issues		
Standard Operating Procedures (organizational & departmental)		
SOP Utilization		
Training Program		
Personnel Qualification Records		
Personnel Training Records		
Contract Programmers Training/Supervision Records		
Internal Auditing Program		
Quality Policy		
Quality Manual		
Documentation Control		
Product Life Cycle Model		
Product Project Plan		

FIG. X1.1 Vendor Assessment Information Sample Form (continued)

X2. VALIDATION PLAN

{Project or Equipment Name under going validation}
{date}

(I) INTRODUCTION

- A) **Objective** (*State the objective(s) of the qualification plan*)
- B) **System Description** (*A description of the purpose, location, and method of operation for the system being qualified*)
- C) **Test Assumptions/Boundaries** (*Narrative describing base assumptions associated with the qualification. Clearly state boundaries of the system and the operating conditions which the qualification will evaluate*)
- D) **Responsibilities** (*Listing of individuals name and what they are responsible for*)
- E) **Development Summary** (*Summary of the developer testing and results ---relates to in-house developed systems*)
- F) **General Instructions** (*Provide detailed instructions, by inclusion or reference, on how to execute IQ/OQ tests*).

(II) INSTALLATION QUALIFICATION (IQ) (*Involves establishing documented evidence that the LIMS is installed and configured to meet design intent and user requirements. IQ does not typically include operation of the system.*)

- G) **Equipment List & Description** (*List of all major hardware and software components with a brief description of the function/construction of each. This list will include the make/model and revision number as appropriate*).
- H) **Engineering Specifications**
 - 1) **Spec's** (*Include any specifications of components needed for the effective installation of the system or sub-system.*)
 - 2) **Drawing list** (*Flow diagrams, data models etc. - whatever gives a clear picture of the system or subsystem*).
 - 3) **Environmental requirements** (*Discuss anything outside the LIMS which it needs to operate effectively. This shall include utility requirements such as power, air conditioning, other software, other hardware, and anything else needed for the LIMS.*)

I) IQ Test Plan/Protocol

- 1) **Test Plan/Protocol** (*Detail here the test plan to verify that the LIMS is installed and configured according to the design and user requirements. The test plan shall provide sufficient instruction to assure that the task is carried out correctly. Each Test plan/protocol shall include the requirement(s) being tested, the test procedure, and the acceptance criteria for each test procedure*).
 - A) **Calibrations**
 - B) **Initial Set-up Procedures**
 - C) **Test Tools**

(III) OPERATIONAL QUALIFICATION (OQ) (*Involves establishing documented evidence that the LIMS operates as intended throughout anticipated ranges. This activity requires evaluation of the LIMS under dynamic operational conditions. Full OQ need not be repeated for each new installation as long as installation is conducted within originally qualified operational ranges and functional requirements. Limited OQ, however, is required*)

- J) **Critical Factors** (*List of critical factors for the LIMS identified during design and development process, vendor information, or technical judgment. The critical factors listed shall be verified via test plans.*)
- K) **OQ Test Plan/Protocol** (*Detail here the test plan/protocol to verify that the LIMS operates according to the design and user requirements. The test plan/protocol shall provide sufficient instruction to assure that the task is carried out correctly. The test plan/protocol must specify expected outcome and acceptance criteria for each critical factor and function. Testing encompasses not only the expected range of input values and volume (normal conditions testing) but also how the LIMS will respond to unusual or extreme operating ranges/conditions (stress testing) and invalid operating ranges/conditions (robustness testing).*)
 - 1) **Standard Operational Procedures(SOPs)**

(IV) Signatures (*This shall include who is submitting the plan for the validation team and will include the name and date of the individuals responsible for reviewing the validation document and concurring that validation was completed correctly. Typically these signatures include someone from the QAU, the laboratory responsible for the LIMS, a representative for the information management group*).

FIG. X2.1 Validation Plan Example

X3. TEST PROTOCOL DESIGN EXAMPLE

ABC Corp., City, State USA	Your Department Name
Date: 06/05/96 Operational Qualification Protocol	Revision: 1.0
Subject: XZY LIMS Application	

K.1 OQ TEST PLAN: Administrator Functions

Verify that the LIMS Administrator Function operates as originally designed. Further, to ensure that LIMS users gain the correct access authority level.

	<u>PROCEDURE</u>	<u>ACCEPTANCE CRITERIA</u>	<u>MET ACCEPTANCE CRITERIA YES/NO?</u>
NORMAL CONDITIONS			
1.	Log into LIMS with Administrator Authority and add a new user to LIMS with User authority. Log out and then log back into LIMS with the User identification just setup.	New user to be added to LIMS with User Authority level. The new user just setup will be able to login LIMS and will have the proper authorities set.	
2.	Log into LIMS with Administrator Authority and change the user from Step 1 from User authority to Stability Group Authority. Log into LIMS with the changed user identification	System will allow this change. User will now have Stability Group Authority rights.	
3.	Change and verify the change of an existing users password. Next log into LIMS using the user id that was just changed and use the changed password.	LIMS will allow the password to be changed and verified. The user will be able to log into LIMS using the changed password.	
4	User added in step 1 logs into LIMS with their current password. Record the screen contents.	The users password will not be displayed and the user will be able to log into LIMS.	
5	User added in step 1 logs into LIMS, changes their password and then verifies the change.	The user will be able to log into LIMS, change their password and verify the change.	
STRESS CONDITIONS			
6.	Add a new user with same name as the user added in step 1.	System will not allow the same user name to be added to LIMS.	
7.	Add a new user with the same name as a the user created in step 1 but with a different authority level.	System will not allow the same user name to be added regardless of the difference in authority levels.	
8.	User logs into LIMS and selects the change password button. They enter their current password and enter the new password and then verify the new password using a password which is different than the new password just entered.	System will accept the current password and the new password entered but will issue an error message and reject the verification of the password.	

All Acceptance Criteria met (circle one) YES NO	
Tested By / Date:	
Reviewed By / Date:	
Comments:	

FIG. X3.1 Test Protocol Design Example

X4. QUALIFICATION REPORT

{Project or Equipment Name under-going validation}
{DATE}

A. CONCLUSIONS

B. DISCUSSION

(1) Compliance within the IQ/OQ Qualification Protocol:

Make reference to the protocol and indicate whether it was followed completely or not. Clearly explain and document any deviations from the protocol and their impact on the system.

(2) Results vs. Success Criteria:

Attach the completed Test Protocols with document signoffs. Discuss results that may not be obvious in the test plan. Identify system limitations and how they will be handled.

(3) Documentation:

Verify that the system documentation is complete and filed for validation purposes.

C. SIGNATURES

Submitted by: _____ for the _____ Team

By signing below, we indicate that we have reviewed the attached Qualification Report and concur that the protocol was followed, all protocol requirements have been satisfied and documented, and acceptance criteria were met except as noted.

(Name / Title) Date

(Name / Title) Date

(Name / Title) Date

(Name / Title) Date

FIG. X4.1 Qualification Sample Report

X5. LIMS VALIDATION ERROR REPORT (6)

User Identification: _____

Date: _____

Error Type: User _____

Program

Other

Evaluation of error: Emergency_____

Normal

Other _____

Error Number (system generated): _____

Error Message:

Error description and suggested solution/actions:

[illegible]

Date _____

Signature

To be filled in by the system manager

Error Type: _____

Degree of Seriousness: _____

Action: _____

Conclusion: _____

FIG. X5.1 LIMS Sample Validation Error Report (6)

X6. CHANGE REQUEST/PROBLEM LOG (3)

(Log ID)

A. Title: System ID: _____ System Name: _____ Version: _____			
B. Nature of the Request or Problem (operational/error, change in business/technology requirements, preventive maintenance, requested change, re-assignment of colleagues, change in documentation or SOPs etc.)			
C. Date of Request/Date Problem Encountered: ____ / ____ / ____			
D. Person Requesting Change/Person Reporting Problem: _____			
E. Findings:			
Assessor: _____ Date: ____ / ____ / ____			
F: Possible Decisions:			
<u>Decisions</u>	<u>Check One</u>	<u>Decided By</u>	<u>Date</u>
Cancel Request/Postpone	_____	_____	____ / ____ / ____
Defer Until Next Version	_____	_____	____ / ____ / ____
Proceed	_____	_____	____ / ____ / ____
Other (Specify Below)	_____	_____	____ / ____ / ____

FIG. X6.1 Sample Change Request/Problem Log (3)

SECTION AREA	SECTION FIELD NAMES	NAME EXPLANATION
A	Log ID	Unique ID assigned to each entry. The assignment should be in some type of order that provides evidence that none are missing.
	System ID	The system's identification number, if it has one.
	System Name	The application's name, for example, ABC LIMS.
	Version	Version of the application, for example, Version 1.2.4.
B	Nature of Request	Description of what is being requested.
C	Date of Request/Problem	The date the problem was discovered or request was made.
D	Person Requesting/Reporting Problem	The person who identified the problem or who is making the request.
	Findings	For change requests, the assessment of findings may include, but is not limited to, requirements, recommendation(s), estimate of work, possible schedule/timeframes, verification testing required, and validation impact. For problems encountered, findings could include what caused the problem, recommended resolution, estimate of work, possible schedule/timeframe, verification testing required, and validation impact. If minor changes are made, then minimal localized and regional tests are recommended.
	Assessor	The person who completed the findings.
	Date	The date the assessor signed.
F	Possible Decision	Check one of the available options.
	Decided by	Signature of the person making the decision.
	Other (page 1)	Other decision made about the event.
G	Resolution	Check all that apply. If the decision was made to proceed, the responsible user party then identifies any one or more items that need to be completed before closing out the entry. If no items are selected, this event could be closed out by having the responsible party sign and date the bottom of the form.
	Completed by	Signed by the person responsible for completing the specific action items.
	Date	Date the action was completed.
	Change to System	This section may have to be completed by the IS professional, either alone, or in close collaboration with the responsible user party for the system. This section could describe briefly and refer to other, more extensive documentation.
	Update/Approve Requirements	This would be checked as a result of finding an error in requirements, an error in the system that impacts the requirements, or an enhancement which requires that the requirements be updated.
	Update/Approve Design Specifications	This could be checked as a result of correcting an error in or clarifying the design specifications, resolving a bug that required a modified design, or an enhancement which requires that the design specifications be updated.
	Update/Approved Validation Plan	If a bug were found or enhancement made that required updating the requirements or design specifications, the validation plan may need to be updated—if only to add/modify test cases.
	Revalidate System	At the time of change, the responsible person in collaboration with the IS person may decide that the system needs to be revalidated.
	Update/Approve/Distribute User Manual	This could be checked due to finding an error in or needing to clarify instructions in the manual, an enhancement, or correcting an error in the system.
	Train Users	This would be checked if the users needed retraining or initial training.
	Update/Approve/Distribute SOPs	During changes or as functional/organization changes are made in the department.
	Move Change into Production	Document exactly what was done to move change into production or reference other documentation.
	Notify Users	Indicate who, how, and when they were notified or reference other documentation.
	Other (page 2)	Indicate here, or reference other documentation, anything else needing to be completed that is not covered elsewhere.
	Responsible Party	Person, typically in the user department, responsible for the validated state of the system. This signature indicates that all resolution actions have been completed and that this log entry is now closed.
	Date	Date the responsible party signed the document.

FIG. X6.1 Sample Change Request/Problem Log (3) (continued)

REFERENCES

- (1) Grigonis, G. J. Jr, and Wyrick, M., "Computer System Validation: Auditing Computer Systems for Quality," *Pharmaceutical Technology*, September 1994.
- (2) McDowall, R. D., "Practical Computer Validation for Pharmaceutical Laboratories," *Journal of Pharmaceutical and Biomedical Analysis*, Vol 14, 1995, pp. 13–22.
- (3) Brodbeck, C. (Parke-Davis), "One Generic Change Control Log Can Serve Multiple Uses," *Warner-Lambert Computer Validation and Training Symposium*, April 1997.
- (4) Double, M. E., and McKendry, M., *Computer Validation Compliance—A Quality Assurance Perspective*, Interpharm Press, IL, 1994.
- (5) Good Automated Manufacturing Practice (GAMP 96), ISPE 3816 W. Linebaugh Ave., Suite 412, Tampa, FL 33624.
- (6) Segalstad, Siri H., Synnevåg, M. J., "A Practical Guide to Validating LIMS, Chemometrics and Intelligent Laboratory Systems," *Laboratory Information Management*, Vol 26, 1994, pp. 1–12.
- (7) McDowall, R. D., "Operational Measures to Ensure the Continued Validation of Computerized Systems in Regulated or Accredited Laboratories," *Laboratory Automation and Information Management*, Vol 31, 1995, pp. 25–34.

(8) Budihandojo, R., "Computerized Systems Validation: A Concept Approach in the Preparation of a Validation Plan Document," *Pharmaceutical Technology*, February 1997.

(9) Budihandojo, R., "Computerized Systems Validation: A Concept Approach in the Preparation of Functional Requirements Document," *Pharmaceutical Technology*, March 1997.

RELATED MATERIAL

Hinton, M. D., *Laboratory Information Management Systems*, Marcel Dekker, Inc., New York, 1995.

Lamprecht, J. L., "Implemementing the ISO 9000 Series," Marcel Dekker, Inc., New York, 1993.

Mahaffey, R., *LIMS Applied Information for the Laboratory*, Van Nostrand Reinhold, New York, 1990.

McDowall, R. D., *Laboratory Information Management Systems*, Sigma Press, Wilmslow, Cheshire, England, 1988.

EPA Good Automated Laboratory Practices (GALP), Scientific Systems Staff, Office of Information Resources Management, U.S. Environmental Protection Agency, Research Triangle Park, North Carolina 27771.

NIST PB-167074, National Institute of Standards and Technology, 820 West Diamond Ave., Gaithersburg, MD 20899.

FDA Glossary of Computerized Systems and Software Development Technology, Taratec Development Corporation, 1170 US Highway 22, Bridgewater, NJ 08807.

The TickIT Guide: A Guide to Software Quality Management System Construction and Certification to ISO 9001, DISC TickIT Office, London, Issue 3.0, 1995.

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