

Support for Title 21 CFR Part 11 and Annex 11 Compliance: Agilent OpenLab ECM

Valid for OpenLab ECM versions 3.5 and 3.6

Overview

US FDA Part 11 in Title 21 of the Code of Federal Regulations (CFR), and its EU analog, Eudralex Chapter 4, Annex 11, describe the requirements for electronic records and electronic signatures for regulated pharmaceutical organizations. Released in 1997, 21 CFR Part 11 has been enforced since 1999. The intent of these guidelines is to ensure that all appropriate electronic records are attributable, legible, contemporaneous, original, accurate, and maintained with integrity.

This white paper is a resource for users of OpenLab ECM in organizations that must comply with these regulations. OpenLab ECM provides a central secured repository for data acquired and processed by Agilent's data systems (e.g., OpenLab CDS) and non-Agilent's data systems (e.g., Water's Empower), and any other software data (e.g., Microsoft Office files, PDF). It is the responsibility of the user and their organization to ensure that the functionalities provided by OpenLab ECM are used appropriately to achieve compliance-readiness for securing storage of laboratory data. In addition to the technical controls OpenLab ECM provide, the user organization must establish procedural controls through standard operating procedures (SOPs) to address relevant non-technical requirements. Governance, for example, as an internal audit program, must also be established to assure that system operators follow the SOPs.

Appendix 1 provides a detailed description of how OpenLab ECM support users and their organizations in achieving the requirements of each section of 21 CFR Part 11 and the related sections of EU Annex 11 and regulations of other countries. The descriptions assume that system access, including instrument hardware and software, is controlled by the staff responsible for the electronic records contained on the system. Thus, the system is designed as a “closed system” as defined in 21 CFR Part 11.3(b)(4).

This document does not describe how functions of applications that generate data like OpenLab CDS, other data systems and software applications like Microsoft Excel meet the requirements for electronic records and signatures. These applications are referred to as primary applications. Any new versions of the files are securely stored and tracked in OpenLab ECM. However, it does not process data or track the specific details of changes within files.

“Separation of duty, as a security principle, has as its primary objective the prevention of fraud and errors. This objective is achieved by disseminating the tasks and associated privileges for a specific business process among multiple users.”

21 CFR Part 11

21 CFR Part 11 covers three specific elements of a regulated laboratory’s operation:

- Security of electronic records,
- Attribution of work,
- Electronic signatures (if used)

Security

Security refers to the “right people, having the right access, to the right information.” Regulated organizations must be able to both verify the identity of system users and limit system access to trained, authorized individuals (11.10(d), (i) and (g); 11.100(b)). Because laboratory staff have different responsibilities based on their job assignments, data access must be able to be segregated and defined such that certain users have certain types of access to certain sets of data while having different access to other data sets.

Attribution of work

Attribution of work refers to documenting the “Who, what, when, where and why?” of work performed. This is usually done via the use of automated audit trail functionality. Automated audit trails independently record user’s actions

thus connecting laboratory staff to the work they perform. Audit trail entries enable staff and regulatory inspectors to reconstruct the complete history of an electronic record.

- Who: clearly identifies the person responsible for the particular action that creates, modifies, or deletes a record.
- What: is the action that took place, including, if applicable, the old value and the new value contained in the record.
- When: unambiguously declares the date and time the action took place.
- Where: clearly identifies the impacted record.
- Why: explains the reason for a change to a regulated record. The reason may be selected from a list of pre-defined reasons to provide consistency and to enable searching and sorting of entries.

eSignatures

While 21 CFR Part 11 does not require the use of eSignatures, it does provide regulations for their use when they are used. In this case, the system must ensure that eSignatures:

- Are irrevocably linked to their respective records.
- Show the full name of the signer, date and time, as well as the meaning of the signature (such as review, approval, responsibility, or authorship).
- Are present whenever the signed records are displayed or printed

– Botha, Eloff, IBM Systems Journal¹

Appendix 1. Satisfying the requirements set forth in US FDA Title 21 CFR Part 11 and related global regulations using OpenLab ECM

Appendix 1 Table: Notes

Column one

The table addresses 21 CFR Part 11 requirements in the order that they are presented in the US FDA reference document.²

Column two

For completeness, column two lists all requirements of 21 CFR Part 11 and other related global requirements. "System" refers to the analytical system used to acquire and process data.

Most requirements are fulfilled by either technical controls (i.e. software functionality) or procedural controls (i.e. SOPs). Technical controls are controls provided by the software and hence the software supplier, while procedural controls are the responsibility of the user organization. 21 CFR requirements listed in bold are requirements addressed by technical controls. Other global requirements are listed in regular font.

Requirements that must be addressed by procedural controls are listed in blue.

Column three

Responsibilities for each requirement are listed in column three. "S" refers to analytical system vendor. "U" refers to the user organization. Use of "S" and "U" implies a combination of both technical and procedural controls.

Column four

If available and where appropriate, related global requirements and comments are provided in column four.

Column five

Column five indicates with a "yes", "no", or "partially" whether the requirement can be satisfied using the technical controls provided in OpenLab ECM. A "partially" response denotes that the requirement is only partially met by technical controls and may rely on procedural controls to be fully satisfied. "N/A" indicates that the requirement is not applicable to the CDS.

Column six

Column six explains how the regulatory requirement can be satisfied using the technical controls provided by OpenLab ECM. Column six also provides additional recommendations for the user organization when relevant.

1. Validation

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Part 11 11.10(a)	1.1 Is the system validated to ensure accuracy, reliability, consistent intended performance, and the ability to discern invalid or altered records?	S, U	<p>Required by all regulations.</p> <p>This is a typical example of shared responsibility between the system supplier and the user organization. While the user organization has ultimate responsibility for validation, some tasks can only be done and must be delivered by the software supplier, e.g., validation activities during development and related documentation.</p> <p>第五章系统</p> <p>第十三条在计算机化系统使用之前,应当对系统全面进行测试,并确认系统可以获得预期的结果。当计算机化系统替代某一人工系统时,可采用两个系统(人工和计算机化)平行运行的方式作为测试和验证内容的一部分。</p> <p>第五章系统</p> <p>第十三条在计算机化系统使用之前,应当对系统全面进行测试,并确认系统可以获得预期的结果。当计算机化系统替代某一人工系统时,可采用两个系统(人工和计算机化)平行运行的方式作为测试和验证内容的一部分。</p>	Yes	<p>Agilent Technologies has verified the performance of OpenLab ECM using tests that evaluate accuracy, reliability and consistent performance. However, the user organization is required to validate their sample preparation system according to regulatory expectations.</p> <p>The system supports all file types.</p> <p>The system tracks unique files and create new version with every subsequent upload of a unique file.</p> <p>The software confirms that a file uploaded to the secure repository is the same as the file created by the primary application (e.g., CDS).</p> <p>Validity check of a file is the primary application responsibility.</p>

1. Validation (continued)

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Annex 11	1.2 Is infrastructure qualified?	U	Annex 11. Principle B Brazil GMP 577	N/A	Qualification of infrastructure such as servers and networks are the responsibility of the user organization.

2. Accurate Copies and Secure Retention and Retrieval of Records

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Part 11 11.10(b)	2.1 Is the system capable of generating accurate and complete copies of records in both human readable and electronic form suitable for inspection, review, and copying by the FDA?	S, U	第五章系统 第十九条以电子数据为主数据时,应当满足以下要求: (一)为满足质量审计的目的,存储的电子数据应当能够打印成清晰易懂的文件。	Yes	Records are available printed on paper or electronically as a PDF file.
Annex 11	2.2 Is it possible to obtain clear printed copies of electronically stored e-records?	S, U	Annex 11.8.1 Brazil GMP 583	Yes	Records are available printed on paper or electronically as a PDF file.
Brazil	2.3 Are there controls to make sure that the data backup, retrieving and maintenance process is duly carried out?	S, U	Brazil 585.2 第五章系统 第十九条以电子数据为主数据时,应当满足以下要求: (三)应当建立数据备份与恢复的操作规程,定期对数据备份,以保护存储的数据供将来调用。备份数据应当储存在另一个单独的、安全的地点,保存时间应当至少满足本规范中关于文件、记录保存时限的要求。	Yes	While backing up data is the responsibility of the user organization, the system is designed to allow backup of all relevant files.
Part 11 11.10(c)	2.4 Does the system protect records to enable their accurate and ready retrieval throughout the records retention period?	S, U	China GMP 163	Yes	All data uploaded into the system is stored in a protected location. It's the user organization's responsibility to manage the physical security and controlled access to the secured storage.
Annex 11	2.5 Are data checked during the archiving period for accessibility, readability and integrity?	U	Annex 11.17	N/A	It is the responsibility of the user organization to ensure data are checked during archival for accessibility, readability, and integrity. The system is designed to ensure that archived data is accessible, readable and cannot be modified.
Annex 11	2.6 If relevant changes are made to the system (e.g. computer equipment or programs), is then the ability to retrieve the data ensured and tested?	S, U	Annex 11.17	Yes	The system is designed to read data from legacy versions. The user organization is responsible for ensuring readability of this data during their implementation and validation processes.
Annex 11	2.7 Are data secured by both physical and electronic means against damage?	S, U	Annex 11.7.1 Brazil GMP 584 第五章系统 第十条系统应当安装在适当的位置,以防止外来因素干扰。 第五章系统 第十九条以电子数据为主数据时,应当满足以下要求: (二)必须采用物理或者电子方法保证数据的安全,以防止故意或意外的损害。日常运行维护和系统发生变更(如计算机设备或其程序)时,应当检查所存储数据的可访问性及数据完整性。	Yes	All records uploaded into the system are stored in a protected location. Physical security is the responsibility of the user organization.

2. Accurate Copies and Secure Retention and Retrieval of Records (continued)

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Clinical guide	2.8 Are there controls implemented that allow the reconstruction of the electronic source/raw documentation for FDA's review of the (clinical) study and laboratory test results?	S	Clinical Computer Guide F2 FDA Q&As	N/A	It is the user organization's responsibility to access the primary application to reconstruct electronic source/raw data.
Clinical guide	2.9 Does the information provided to FDA fully describe and explain how source/raw data were obtained and managed, and how electronic records were used to capture data?	U	Clinical Computer Guide F2 FDA Q&As	N/A	It is the responsibility of the user organization to describe how source/raw data were obtained and managed, and how electronic records were used to capture data.
Annex 11	2.10 Does the system allow performing regular back-ups of all relevant data?	S, U	Annex 11.7.1 China GMP 163 Brazil GMP 585 Part 211, 68 b	Yes	While backing up data is the responsibility of the user organization, the system is designed to allow backup of all relevant files.
Annex 11	2.11 Is the integrity and accuracy of backup data and the ability to restore the data checked during validation and monitored periodically?	U	Annex 11.7.2 China GMP 163 Brazil GMP 585 Part 211, 68 b	N/A	It is the responsibility of the user organization to ensure the integrity and accuracy of the backed-up data, and to check, validate and monitor restored data periodically.
Clinical Computer Guide	2.12 Are procedures and controls in place to prevent the altering, browsing, querying, or reporting of data via external software applications that do not enter through the protective system software?	S, U	Clinical Computer Guide E	Yes	OpenLab ECM is a closed system and can only be accessed with privileged user credentials. Applications access to files are controlled via APIs.
Clinical Computer Guide	2.13 Are there controls implemented to prevent, detect, and mitigate effects of computer vi-ruses, worms, or other potentially harmful software code on study data and software?	S, U	Clinical Computer Guide F	N/A	Agilent has tested OpenLab ECM in conjunction with industry standard anti-virus applications. However, it is the responsibility of the user organization to implement anti-virus software.

3. Authorized Access to Systems, Functions, and Data

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Part 11 11.10(d)	3.1 Is system access limited to authorized persons?	S, U	China GMP 183 163 Brazil GMP 579, ICH Q7.5.43	Yes	OpenLab ECM user-based access controls require a unique username and password combination. It is the user organization's responsibility to configure and manage these users.
	3.2 Is each user clearly identified, e.g., through his/her own user ID and Password?	S, U	Several Warning Letters	Yes	Each user is identified by a unique ID and password combination. Entry of both is required to access the system.
Clinical	3.3 Are there controls to maintain a cumulative record that indicates, for any point in time, the names of authorized personnel, their titles, and a description of their access privileges?	S, U	Clinical Computer Guide 4	Yes	Creation and updates of users' information, including their roles are recorded in the activity log, and it is possible to search changes to specific user record.

4. Electronic Audit Trail

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Part 11 11.10(e)	4.1 Is there a secure, computer-generated, time-stamped audit trail to independently record the date and time of operator entries and actions that create, modify, or delete electronic records?	S	China GMP 163 第五章系统 第十六条计算机化系统应当记录输入或确认关键数据人员的身份。只有经授权人员,方可修改已输入的数据。每次修改一个已输入的关键数据均应当经过批准,并应当记录更改数据的理由。应当根据风险评估的结果,考虑在计算机化系统中建立一个数据审计跟踪系统,用于记录数据的输入和修改。	Yes	All operations performed on files stored in OpenLab ECM are recorded in a secured, system generated, time stamped Activity Log which also includes a reason for the change.
FDA GLP	4.2 Does the audit trail record who has made which changes, when and why?	S	FDA 21 CFF 58.130 e Clinical Computer Guide 2 Clinical Source Data 3	Yes	OpenLab ECM activity log lists modifications, date and time of the change, the user ID, and the reason for change.
Annex 11	4.3 Can the system generate printouts indicating if any of the e-records has been changed since the original entry?	S, U	Annex 11, 8.2	Yes	It is the primary application's responsibility to track the details of the changes inside the e-record audit trail. Once an e-record is uploaded and stored in OpenLab ECM, it is assigned a unique id and new file version. When change(s) are made to the file and it is uploaded again to OpenLab ECM, the system detects that and stores the file with new sequential version. Every operation on the file's (e.g., upload, move) is recorded in an activity log which can be exported and printed out.
FDA GMP	4.4 Does the audit trail include any modifications of an established method employed in testing? 4.5 Do such records include the reason for the modification?	S	Part 211.194 8b	Yes	OpenLab ECM does not control test equipment.
	4.6 Is the audit trail function configured to be always on and can it not be switched off by system users?	S, U	Warning Letter	Yes	OpenLab ECM allows users to configure whether the activity log is active or not during account creation. Once an activity log is set to be active, it cannot be de-activated by any user. Further, entries in the activity log cannot be switched off, altered, or deleted by any user.
Annex 11	4.7 Is audit trail available to a generally intelligible form for regular review?	S	Annex 11, 9	Yes	OpenLab ECM activity log has been designed to be easily reviewed and can be exported and printed out.
	4.6 Is the audit trail function configured to be always on and can it not be switched off by system users?	S, U	Warning Letter	Yes	OpenLab ECM allows users to configure whether the activity log is active or not during account creation. Once an activity log is set to be active, it cannot be de-activated by any user. Further, entries in the activity log cannot be switched off, altered, or deleted by any user.
Annex 11	4.7 Is audit trail available to a generally intelligible form for regular review?	S	Annex 11, 9	Yes	OpenLab ECM activity log has been designed to be easily reviewed and can be exported and printed out.
	4.8 Can audit trail contents be configured such that only relevant activities are recorded for realistic and meaningful review of audit trail information?	N/A	Implicitly required by Annex 11 and many warning letters related to review of audit trail.	Yes	OpenLab ECM captures all file related operations in the Activity Log. OpenLab ECM allows users to set filtering of the activity log prior to displaying its content to address user preferences for reviewing the information it contains.

4. Electronic Audit Trail(continued)

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Part 11 11.10(e)	4.9 Is previously recorded information left unchanged when records are changed?	S		Yes	Primary application (e.g., OpenLab CDS) records all data entry additions, changes, and deletions. Changes to files stored in OpenLab ECM are saved as new revisions of the original file, which is left unchanged. When opening files for further processing, the user chooses the version of the file used (based on their permissions).
Part 11 11.10(e)	4.10 Is audit trail documentation retained for a period at least as long as that required for the subject electronic record?	S, U		Yes	Primary application record's Audit trail information is stored within the electronic record. Activity log information for files stored in OpenLab ECM is linked to the electronic record and cannot be separated from it.
Part 11 11.10(e)	4.11 Is audit trail available for review and copying by the FDA?	S		Yes	Activity log can be viewed, filtered, and printed.
Annex 11	4.12 Is it possible to obtain clear printed copies of electronically stored e-records (e.g., e-audit trail?)	S	Annex 11, 8.1	Yes	It is the responsibility of primary applications like OpenLab CDS to enable access and print out of the audit trails. OpenLab ECM allows access to view and print record's activity log.

5. Operational Device Checks

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation to customers?
Part 11 11.10(f)	5.1 Are there operational system checks to enforce permitted sequencing of steps and events, if required?	S, U		N/A	It is the responsibility of the user organization to designate and enforce procedural controls.
Part 11 11.10(g)	5.2 Are there authority checks to ensure that only authorized individuals can use the system, electronically sign a record, access the operation or computer system input or output device, alter a record, or perform the operation at hand?	S	Part 211, 68 b	Yes	The system supports configurable user roles that control access to content and ability to perform specific operations.
	5.3 Is the system designed to record the identity of operators entering, changing, confirming or deleting data including date and time?	S	Annex 11, 12.4	Yes	The identity of operators performing actions are recorded in the activity log.
Part 11 11.10(h)	5.4 Does the system allow to use device checks to determine, as appropriate, the validity of the source of data input or operational instruction?	S, U	There are two equally valid interpretations of this requirement. Systems should be designed such that: 1. Proper communication is confirmed between the computer and the "source" of data input (i.e., the instrument) prior to transmission of instructions to or data from the "source." 2. Regulated records created by the system must unambiguously indicate the "source" of the data (i.e., which instrument or component generated the data.)	Partially	The system is designed to continually ensure a valid connection between the primary application and the server. OpenLab ECM stores information/records acquired and processed by primary applications. It is not connected directly to the instrument components (e.g., LC modules) that generates the data and unaware of the data source.

5. Operational Device Checks (continued)

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation to customers?
Part 11 11.10(i)	5.5 Is there documented evidence that persons who develop, maintain, or use electronic record/electronic signature systems have the education, training, and experience to perform their assigned tasks?	U	China GMP 18 Brazil 571	N/A	It is the responsibility of the user organization to maintain documented evidence that the persons who develop, maintain, or use electronic record and electronic signature systems have the education, training, and experience needed to perform these tasks. Agilent software professionals involved in development of OpenLab ECM have received training in relevant aspects of data integrity.
Part 11 11.10(j)	5.6 Is there a written policy that hold individuals accountable and responsible for actions initiated under their electronic signatures, in order to determine record and signature falsification?	U		N/A	It is the responsibility of the user organization to establish a written policy (SOP) that holds staff responsible for the actions initiated under their electronic signatures.
	5.7 Have employees been trained on this procedure?	U	Implied requirement of Part 11 11.10(j)	N/A	It is the responsibility of the user organization to train their staff.
Part 11 11.10(k)	5.8 Are there appropriate controls over systems documentation including: 1. Adequate controls over the distribution of, access to, and use of documentation for system operation and maintenance? 2. Revision and change control procedures to maintain an audit trail that documents time-sequenced development and modification of systems documentation.	U	China GMP 161 第五章系统 第十一条应当有详细阐述系统的文件 (必要时, 要有图纸), 并须及时更新。此文件应当详细描述系统的工作原理、目的、安全措施和适用范围、计算机运行方式的主要特征, 以及如何与其他系统和程序相接。	N/A	1. It is the responsibility of the user organization to establish systems documentation. 2. Agilent maintains development and testing documentation for OpenLab ECM. Upon request, this documentation is available for review.
Part 11 11.10(i)	5.9 Are there revision and change control procedures to maintain an audit trail that documents time-sequenced development and modification of systems documentation?	S, U	第五章系统 第十七条计算机化系统的变更应当根据预定的操作规程进行, 操作规程应当包括评估、验证、审核、批准和实施变更等规定。计算机化系统的变更, 应经过该部分计算机化系统相关责任人员的同意, 变更情况应有记录。主要变更应当经过验证。	Yes	Agilent maintains development and testing documentation for OpenLab ECM. Upon request, documentation is available for review. The user organization is expected to maintain documentation of their system and associated changes in situ.

6. Data Integrity, Date and Time Accuracy

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Annex 11	6.1 Do computerized systems exchanging data electronically with other systems include appropriate built-in checks for the correct and secure entry and processing of data?	N/A	Annex 11.5	N/A	In this context, OpenLab ECM does not exchange data with other systems. It simply stores data generated and process by other systems. OpenLab ECM ensures secure and faithful transfer of files.
Annex 11	6.2 Is there an additional check on the accuracy of the data? (This check may be done by a second operator or by validated electronic means.)	U	Annex 11-6 Brazil GMP 580 ICHQ7-5.45 第五章系统 第十五条当人工输入关键数据时 (例如在称重过程中输入物料的重量和批号), 应当复核输入记录以确保其准确性。这个复核可以由另外一个操作人员完成, 或采用经验证的电子方式。必要时, 系统应当设置复核功能, 确保数据输入的准确性和数据处理过程的正确性。	N/A	It is the responsibility of the user organization to define procedures for additional checks and to ensure accuracy of the data. In OpenLab ECM, when the data is transferred over HTTP/HTTPS, checksum of data is calculated on server side and client side whose values should match. In case these values do not match, the file is not uploaded, and considered corrupted.

6. Data Integrity, Date and Time Accuracy (continued)

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Clinical Computer Guide	6.3 Are controls established to ensure that the system's date and time are correct?	S, U	Clinical Computer Guide D.3	Yes	Agilent recommends that the system be configured to reference a time server to ensure accuracy of the system date and time. This is configured in and controlled by the operating system.
Clinical Computer Guide	6.4 Can date or time only be changed by authorized personnel, and is such personnel notified if a system date or time discrepancy is detected?	U	Clinical Computer Guide D.3	N/A	OpenLab ECM is designed to synchronize with local Windows time. It is the user organization's responsibility to: <ul style="list-style-type: none"> – Limit access controls of Windows time settings to only authorized personnel. – Maintain procedural controls for setting and maintaining the accuracy of Windows time.
Clinical Computer Guide I	6.5 Are timestamps with a clear understanding of the time zone reference used implemented for systems that span different time zones?	S, U	Clinical Computer Guide D.3	Yes	All time data is time stamped in Coordinated Universal Time (UTC)/Greenwich Mean Time (GMT) and displayed in the local time of the computer used.

7. Control for Open Systems (Only applicable for open systems)

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Part 11 11.30	7.1 Are there procedures and controls designed to ensure the authenticity, integrity, and, as appropriate, the confidentiality of electronic records from the point of their creation to the point of their receipt?	N/A		N/A	OpenLab ECM is not intended to be deployed as an "open" system as per 21 CFR Part 11.3(b)(9).
Part 11 11.30	7.2 Are there additional measures such as document encryption and use of appropriate digital signature standards to ensure, as necessary under the circumstances, record authenticity, integrity, and confidentiality?	N/A		N/A	OpenLab ECM is not intended to be deployed as an "open" system as per 21 CFR Part 11.3(b)(9).

8. Electronic Signatures – Signature Manifestation and Signature/Record Linking

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Annex 11	8.1 When electronic signatures are used, do they have the same impact as hand-written signatures within the boundaries of the company? Are they permanently linked to their respective record? Do they include the time and date that they were applied?	N/A	Annex 11.14 ICH Q7.6.18 第五章系统 第二十三条电子数据可以采用电子签名的方式, 电子签名应当遵循相应法律法规的要求。	Yes	The user organization must establish the legal impact of electronic signatures Signatures are permanently linked to their respective records. Signed electronic records show the name of the signer, and date and time the signature was executed, and the meaning of the signature.
Part 11 11.50 (a)	8.2 Do signed electronic records contain information associated with the signing that clearly indicates all of the following: 1. The printed name of the signer? 2. The date and time when the signature was executed? and 3. The meaning (such as review, approval, responsibility, or authorship) associated with the signature?	N/A		Yes	Signed electronic records show 1. the name of the signer, 2. the date and time the signature was executed, 3. and the meaning of the signature.

8. Electronic Signatures – Signature Manifestation and Signature/Record Linking

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Part 11 11.50 (b)	8.3 Are the items identified in paragraphs (a)(1), (a)(2), and (a)(3) of this section subject to the same controls as for electronic records and are they included as part of any human readable form of the electronic record (such as electronic display or printout)?	N/A		Partially	All electronic signature components are displayed in human readable form. Only PDF electronic signatures can be printed.
Part 11 11.70	8.4 Are electronic signatures and handwritten signatures linked to their respective electronic records to ensure that the signatures cannot be excised, copied, or otherwise transferred to falsify an electronic record by ordinary means?	N/A		Yes	Handwritten signatures are not addressed by the system and must be managed procedurally by the user organization. Electronic signatures are embedded in the electronic record and cannot be modified, overwritten or deleted.
Part 11 Preamble	8.5 Is there a user specific automatic inactivity disconnect measure that would “de-log” the user if no entries or actions were taken within a fixed short timeframe?	U	Part 11 Preamble section 124	Yes	Automatic session locking enables the user organization to configure a time after which the user is automatically logged-out.

9. Electronic Signatures General Requirements and Signature Components and Controls

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation to customers?
Part 11 11.100(a)	9.1 Is each electronic signature unique to one individual and not reused by, or reassigned to, anyone else?	N/A		Yes	The system does not allow duplicate user IDs. Each user has a unique login and thus a unique signature that cannot be used by another user.
Part 11 11.100(b)	9.2 Does the organization verify the identity of the individual before the organization establishes, assigns, certifies, or otherwise sanctions an individual's electronic signature, or any element of such electronic signature?	N/A		N/A	It is the responsibility of the user organization to verify the identity of staff before it establishes, assigns, certifies, or otherwise sanctions an individual's electronic signature, or any element of such electronic signature.xw
Part 11 11.100 (c)	9.3 Are persons using electronic signatures, prior to or at the time of such use, certified to the agency that the electronic signatures in their system, used on or after August 20, 1997, are intended to be the legally binding equivalent of traditional handwritten signatures? 9.4 Do persons using electronic signatures, upon agency request provide additional certification or testimony that a specific electronic signature is the legally binding equivalent of the signer's handwritten signature?	N/A		N/A	It is the responsibility of the user organization to verify that staff using electronic signatures meet these requirements.
Part 11 11.200(a) (1)	9.5 Do electronic signatures that are not based upon biometrics employ at least two distinct identification components such as an identification code and password?	N/A		Yes	Both identification (user ID) and password are required to make an electronic signature.
Part 11 11.200(a) (1) (i)	9.6 When an individual executes a series of signings during a single, continuous period of controlled system access, is the first signing executed using all electronic signature components?	N/A		Yes	Both identification (user ID) and password are required to make the first electronic signature.

9. Electronic Signature General Requirements and Signature Components and Controls (continued)

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation to customers?
Part 11 11.200(a) (1) (i)	9.7 When an individual executes a series of signings during a single, continuous period of controlled system access, are subsequent signings executed using at least one electronic signature component that is only executable by, and designed to be used only by, the individual?	N/A		Yes	It is the user organization's responsibility to ensure the users are not sharing passwords.
Part 11 11.200(a) (1) (ii)	9.8 When an individual executes one or more signings not performed during a single, continuous period of controlled system access, is each signing executed using all of the electronic signature components?	N/A		Yes	Both identification (user ID) and password are required to make an electronic signature when signing is not performed during a single, continuous period of controlled system access.
Part 11 11.200(a) (2)	9.9 Are controls in place to ensure that electronic signatures that are not based upon biometrics are used only by their genuine owners?	N/A		N/A	It is the user organization responsibility to ensure that user names and passwords are known only by the assigned individuals and are traceable to individual users.
Part 11 11.200(a) (3)	9.10 Are the electronic signatures be administered and executed to ensure that attempted use of an individual's electronic signature by anyone other than its genuine owner requires collaboration of two or more individuals?	N/A		Yes	Misuse of electronic signatures by anyone other than the owner would require intentional cooperation of a user and the System Administrator.
Part 11 11.200(b)	9.11 Are electronic signatures based upon biometrics designed to ensure that they cannot be used by anyone other than their genuine owners?	N/A		N/A	Biometric authentication is not supported in OpenLab ECM

10. Controls for Identification Codes and Passwords

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Part 11 11.300(a)	10.1 Are controls in place to maintain the uniqueness of each combined identification code and password, such that no two individuals have the same combination of identification code and password?	S, U		Yes	OpenLab ECM does not allow duplicate user IDs. Same user IDs can be used across accounts.
Part 11 11.300(b)	10.2 Are controls in place to ensure that identification code and password issuance are periodically checked, recalled, or revised (e.g., to cover such events as password aging)?	S, U		Yes	Password expiration is configurable for built-in users. For NT Domain users, the user organization should configure password expiration based on a documented risk assessment.
Part 11 11.300(c)	10.3 Are there procedures to electronically deauthorize lost, stolen, missing, or otherwise potentially compromise tokens, cards, and other devices that bear or generate identification code or password information, and to issue temporary or permanent replacements using suitable, rigorous controls?	U	第五章系统 第十四条数据的输入或修改只能由经许可的人员进行。杜绝未经许可的人员输入数据的手段有：使用钥匙、密码卡、个人密码和限制对计算机终端的访问。应当就输入和修改数据制订一个授权、取消、授权变更，以及改变个人密码的规程。必要时，应当考虑系统能记录未经许可的人员试图访问系统的行为。对于系统自身缺陷，无法实现人员控制的，必须具有书面程序，相关记录本及相关物理隔离手段，保证只有经许可的人员方能进行操作。	N/A	It is the responsibility of the user organization to establish these procedures.

10. Controls for Identification Codes and Passwords (continued)

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation?
Part 11 11.300(d)	10.4 Are there transaction safeguards in place to prevent unauthorized use of passwords and/or identification codes, and to detect and report in an immediate and urgent manner any attempts at their unauthorized use to the system security unit, and, as appropriate, to organizational management?	U	第五章系统 第十四条数据的输入或修改只能由经许可的人员进行。杜绝未经许可的人员输入数据的手段有：使用钥匙、密码卡、个人密码和限制对计算机终端的访问。应当就输入和修改数据制订一个授权、取消、授权变更，以及改变个人密码的规程。必要时，应当考虑系统能记录未经许可的人员试图访问系统的行为。对于系统自身缺陷，无法实现人员控制的，必须具有书面程序，相关记录本及相关物理隔离手段，保证只有经许可的人员方能进行操作。	N/A	It is the responsibility of the user organization to establish these transaction safeguards.
Part 11 11.300(e)	10.5 Are there controls for initial and periodic testing of devices, such as tokens or cards, that bear or generate identification code or password information to ensure that they function properly and have not been altered in an unauthorized manner?	U		N/A	It is the responsibility of the user organization to establish controls to test devices initially as well as periodically to ensure they function properly and have not been altered in an unauthorized manner.

11. System Development and Support

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation to customers?
Annex 11	11.1 Has the software or system been developed in accordance with an appropriate quality management system?	S, U	Annex 11 4.5 Brazil GMP 577 GAMP This is a shared responsibility between the system supplier and the user organization. The user should require the supplier to provide documented evidence that software is developed within the framework of a quality management system (QMS). 第二章原则 企业应当能够提供与供应商质量体系和审计信息相关的文件。	Yes	OpenLab ECM is developed within the ISO 9001 Quality Management Standard (Ref. section 2.2 of the LSCA Quality Manual).
Brazil	11.2 Is there a formal agreement in case of the software supplier subcontracts software and maintenance services. Does the agreement include the contractor's responsibilities?	S, U	Brazil GMP 589 第二章原则 第四条企业应当注重计算机化系统供应商的管理，制定相应的操作规程。供应商提供产品或服务时（如安装、配置、集成、验证、维护、数据处理等），企业应当与供应商签订正式协议，明确双方责任。	Yes	Agilent requires formal agreements with all suppliers. (Ref. section 7.4 of the LSCA Quality Manual).
ICH Q10	11.3 For outsourced (development and support) activities, is there a written agreement between the contract giver and contract acceptor?	S, U	ICHQ10, 2.7 c	Yes	Agilent requires formal agreements with all suppliers (Ref. section 7.4 of the LSCA Quality Manual).

11. System Development and Support (continued)

Part 11 Others	Requirements	S, U	Other regulations or comments	Yes/No/ Partially	If yes, how, specifically, is the requirement satisfied? or If no, what is the recommendation to customers?
ICH Q10	11.4 Are the responsibilities and communication processes for quality related activities of the involved parties (contractors) defined?	S, U	ICHQ10, 2.7 c	Yes	Responsibilities and communication processes for quality related activities are part of Agilent's agreements with suppliers.
Part 11 11.10(i)	11.5 Is personnel developing and supporting software trained?	S, U	第三章人员 第五条计算机化系统的“生命周期”中所涉及的各种活动,如验证、维护、管理等,需要各相关的职能部门人员之间的紧密合作。在职责中涉及使用和管理计算机化系统的人员,应当接受相应的使用和管理培训。确保有适当的专业人员,对计算机化系统的设计、验证、安装和运行等方面进行培训和指导。	Yes	All Agilent personnel are required to be trained (Ref. section 6.0 of the LSCA Quality Manual).

References

1. R. A. Botha and J. H. P. Eloff. Separation of duties for access control enforcement in workflow environments. IBM Systems Journal– End-to-end security. 40 (3), 666-682. (2001).
2. U.S. Food and Drug Administration. CFR - Code of Federal Regulations Title 21. Title 21—Food and Drugs, Chapter I—Food and Drug Administration Department of Health and Human Services, Subchapter A— General. Part 11 Electronic Records; Electronics Signatures [Online] <https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfcfr/CFRSearch.cfm?CFRPart=11> (accessed November 2, 2020.)
3. European Commission Health and Consumers Directorate-General. Public Health and Risk Assessment. Pharmaceuticals. EudraLex. The Rules Governing Medicinal Products in the European Union.
4. Good Manufacturing Practice. Medicinal Products for Human and Veterinary Use. Annex 11. Computerized Systems. [Online] http://ec.europa.eu/health/files/eudralex/vol-4/annex11_01-2011_en.pdf (accessed November 2, 2020)

www.agilent.com/chem/openlab-ecm

This information is subject to change without notice.

DE-002720

© Agilent Technologies, Inc. 2024
Printed in the USA, May 14, 2024
5994-3547EN

