



SiteVault 26R1 Validation Release Certification

This document confirms and attests that the 26R1 release of **Veeva SiteVault** on April 17, 2026, has been successfully validated by Veeva Systems.

Approvals

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|----------------------------|---|---|-------------|
| Doc ID: | QV-56441 | | |
| Name | Title | Signature | Date |
| Business Approval | | | |
| Cooper VanLare | VP Product, Site Solutions | <i>Approval by Electronic Signature in EDMS Vault</i> | |
| Validation Approval | | | |
| Harneet Kaur | CSV Program Manager, Clinical Systems | <i>Approval by Electronic Signature in EDMS Vault</i> | |
| Quality Approval | | | |
| Melissa Woodruff | Senior Quality Operations Program Manager | <i>Approval by Electronic Signature in EDMS Vault</i> | |

The validation package is published to Veeva Compliance Docs. In the case of an audit or inspection, the validation package can be made available to SiteVault customers by contacting [Veeva Site Support](#). The validation package includes the following documents:

| Deliverable | Description |
|---|--|
| System Overview | Overview of the system and the validation boundaries |
| Validation Project Plan | The plan of activities, deliverables, and resources |
| Qualification Protocol | Overview of the testing environment, strategy, assumptions, and test design |
| Requirements & Specifications¹ | System requirements and specifications |
| Installation Qualification Testing¹ | Qualification of the test team, devices used, test environment, and configurations required for the system to perform its intended use |
| Operational Qualification Testing | Testing and verification of the functional requirements |
| Performance Qualification Testing | Testing and verification of business processes and performing its intended use within a normal operating environment |
| Requirements Traceability Matrices | Traceability of system requirements to test cases |
| System Release Memo² | Early summary of validation results to support customer pre-release testing |
| Validation Summary Report | Summary of all the validation activities performed and confirmation the system is fit for use |

System validation will be maintained through the use of approved standard operating procedures. All future modifications to the core functionality will be addressed through change control procedures. Veeva has validated the technical controls of SiteVault, accessed via web or Veeva Vault Mobile³, in alignment with industry-recognized frameworks including:

- ISPE GAMP 5 A Risk-Based Approach to Compliant GxP Computerized Systems
- PIC/S Good Practices for Computerized Systems in Regulated “GXP” Environments
- FDA General Principles of Software Validation

¹ Specifications and Installation Qualification Testing are only available during audit of Veeva, or in the case of Health Authority Inspection/Audit review with Veeva personnel supporting.

² Only applicable for SiteVault Enterprise customers.

³ Veeva Vault Mobile functionality validated as part of the Veeva Vault Platform.

Signature page for SiteVault 26R1 Validation Release Certification (v1.0)

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