Veeva Vault Clinical Suite is the industry’s first and only cloud platform that streamlines clinical data management and clinical operations with modern and fast applications for EDC, eSource, CTMS, eTMF, and study start-up. Veeva’s suite of clinical applications is built on the Vault Platform, the only content management platform with the unique capability to manage both content and data, eliminating system silos, and streamlining end-to-end clinical trial processes.

**Vault EDC**
Veeva is delivering a better EDC that’s modern, adaptive, and fast to dramatically reduce cost and complexity of clinical trials. Veeva’s unique approach to addressing the entire data ecosystem ensures that each contributor / consumer will see their role enhanced and simplified through the use of guided intelligence and considered process design. This innovative and integrated approach to clinical data management improves data speed and quality. Vault EDC is planned for availability in April 2017.

**Vault eSource**
Veeva Vault eSource transforms site data collection and management for immediate data quality. This eliminates wasted time and cost. Vault eSource delivers real-time site collaboration across sites, sponsors and CROs. Veeva’s unique approach to deliver eSource and EDC together on the same platform eliminates multiple steps, including data transcription and source data verification. Vault eSource is planned for availability in December 2017.

**Vault CTMS**
Veeva Vault CTMS is the only true multitenant cloud solution that unifies information and documentation for a single source of truth across clinical operations. Sponsors, CROs, and investigators can have a single source for clinical master data with one system of record for study, study country, and study site information. With a comprehensive, real-time view of trial information, organizations can make faster, more informed decisions into the performance and efficiency of clinical trials. Vault CTMS is planned for availability in the first quarter of 2017.

**Vault eTMF**
Vault eTMF provides real-time inspection readiness, full visibility into TMF status, and access for all study partners. Sponsors get the clarity they need to oversee trials more effectively. CROs gain the flexibility and control required to operationalize their SOPs and efficiently populate the eTMF. Auditors get easy online access with a dedicated role. And sites receive a simple and efficient means to interact with CROs and sponsors. Vault eTMF promotes the highest levels of TMF quality, access, visibility, and control.

**Vault Study Startup**
Vault Study Startup accelerates time to site activation by connecting global teams and enabling best practices for managing country and site start-up processes. Content-intensive start-up processes and milestone maintenance activities are managed in a single system, providing unparalleled insight and efficiency. Study startup and TMF content and data are seamlessly accessed across teams, systems, and studies.

**Vault SiteExchange**
Vault SiteExchange makes it easy for sponsors, CROs, and investigator sites to collaborate on study activities by providing a single point of access and consistent processes for document exchange during clinical trial execution.

Sponsors, CROs, and investigator sites are one log-in away from securely accessing the latest study documents, workflows, alerts, and notifications across multiple trials all in one place. With a single view and consistent processes for document exchange across trials, Veeva Vault SiteExchange streamlines information sharing among clinical teams, increases visibility across studies, and improves operational efficiency.
**Key Capabilities of a Unified Application Suite**

The only clinical suite that combines EDC, eSource CTMS, eTMF, and study start-up delivering clinical trial excellence across clinical operations and data management.

**Single application platform**

Veeva’s suite of clinical applications is built on the Vault platform, the only content management platform with the unique capability to manage both content and data, eliminating system silos, and streamlining end-to-end clinical trial processes.

**Single source of truth for content and data**

Enter trial data and content once and leverage it across Vault eSource, Vault EDC, Vault eTMF, Vault Study Startup, and Vault CTMS applications. Access and share the same content and data with all study partners, ensuring timeliness and greater accuracy.

**Single source for clinical master data**

Ensure high quality data across clinical applications with one system of record for master study, study country, and study site information.

**Unified view**

Optimize your development portfolio by making faster and more informed decisions with a comprehensive and accurate view of trial status.

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**Vault Platform**

**Proven Platform for Regulated Content and Data Management**

Veeva Vault is the first cloud platform built from the ground up to meet the rigorous usability, scalability, performance, validation, and security requirements of the life sciences industry.

Uniquely designed for both content and data on a single platform, organizations can quickly use the Vault applications to manage end-to-end processes and associated content. The Vault Platform leverages the latest in cloud technology, and is delivered and accessed through the web for greater ease-of-use. Hosted at SOC 1 Type II and ISO 27001 certified global data centers, every release is IQ and OQ qualified reducing the validation effort.

With a modern user experience and cloud pace of innovation, Vault Platform is the next generation of regulated content and data management.