

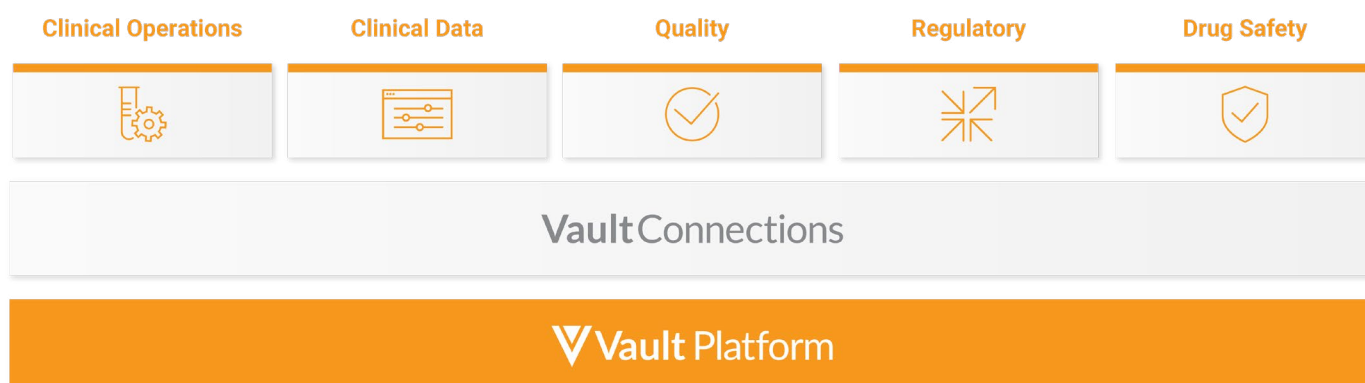
# Veeva Development Cloud

## Technology Foundation for Product Development

Veeva Development Cloud is the technology foundation for product development that brings together applications for clinical, quality, regulatory, safety, and commercial to help organizations drive end-to-end business processes. Today, product development and manufacturing systems are not well integrated, which creates inefficiencies and slows down critical operations. Veeva is the first and only company to offer unified suites of applications that are connected on a single cloud platform. This enables organizations to centralize content and data across global departments for greater efficiency and compliance.

### Vault Connections

Vault Connections are Veeva-delivered integrations that seamlessly transfer data and documents between clinical, quality, regulatory, safety, and commercial Vaults. They are designed to streamline cross-functional business processes by breaking down silos, providing greater visibility, and automating manual tasks. View the [Vault Connections Resource Hub](#) for a full list of available Vault Connections.



## Vault Clinical Operations

**The only suite of unified clinical operations applications on a single cloud platform.**

The industry's first and only suite of unified clinical operations applications—including study start-up, eTMF, CTMS, payments, and site connect on a single cloud platform—to accelerate trial execution and deliver real-time visibility.

### Vault Study Startup

Accelerate time to site activation.

### Vault eTMF

Enable active eTMF for real-time inspection readiness.

### Vault Study Training

Streamline and automate training.

### Vault CTMS

Enable proactive trial management.

### Vault Payments

Pay clinical research sites faster.

### Veeva Site Connect

Automate information sharing.

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## Vault Clinical Data

**Accelerate study timelines with modern, innovative applications for clinical data.**

Vault Clinical Data Management helps clinical teams collect, aggregate, clean, and manage trial data with agility and speed.

### Vault EDC

Collect, clean, and review study data.

### Veeva RTSM

Randomize subjects and manage trial product supply.

### Veeva CDB

Manage complete and concurrent study data.

### Veeva ePRO

Capture data directly from clinical trial participants.

### Veeva eClinRO

Collect clinical measurements by healthcare professionals.

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## Vault Quality

**Modernize quality management while driving compliance and operational innovation.**

Seamlessly manage your quality processes and content with the Vault Quality suite of applications. All parties have access to a single authoritative source, enabling greater visibility and control.

### Vault QMS

Easily manage all quality processes.

### Vault QualityDocs

Document control for all GxP documents.

### Vault Training

Ensure compliance and role-based qualification.

### Vault Validation Management

Execute paperless validation.

### Veeva Learn GxP

Accredited GxP eLearning courses.

### Vault LIMS

Accelerate batch release with a unified lab and quality ecosystem.

### Vault Batch Release

Accelerate GMP release and market-ship decisions.

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## Vault RIM

**Bring speed and agility to your regulatory team with unified RIM.**

Manage the planning, execution, and oversight of all regulatory activities within a single, unified RIM platform.

### Vault Registrations

Manage product registrations globally.

### Vault Submissions Publishing

Automate publishing during submission development.

### Vault Submissions

Speed submission development.

### Vault Submissions Archive

Archive published dossiers securely in the cloud.

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## Vault Safety

**Veeva Vault Safety is the only modern application for the collection, management, and real-time oversight of adverse events.**

### Vault Safety

Real time management and oversight for adverse events.

### Vault SafetyDocs

Centrally manage pharmacovigilance content.