Transforming clinical study operations and management

The Veeva Clinical Suite delivers a unified, cloud-based solution to manage complex studies common among consumer goods, cosmetics, and crop science companies. With applications focused on managing study documents (eTMF), data and processes (CTMS), as well as tracking payments to sites, company sponsors, CROs, and other study partners can now have full visibility into the process and a single source of truth for all study information. The Veeva Clinical Suite delivers speed, visibility and accuracy to a critical business capability.

Veeva eTMF

Veeva eTMF provides real-time inspection readiness, full visibility into TMF completeness, and access for all study partners. Sponsors get the clarity they need to oversee studies more effectively, CRO’s gain the flexibility to efficiently populate the eTMF, and sites can efficiently interact with sponsors and CRO’s. Veeva eTMF delivers quality, access, visibility, and control.

Veeva CTMS

Veeva CTMS unifies study information and processes, streamlines study management and monitoring, and delivers visibility across the study portfolio. Manage the entire end-to-end clinical study process and gain a global view into tasks in one unified system. One seamless system of record for study data and information improves operational efficiency and enables faster, higher quality study execution.

Veeva Payments

Veeva Payments speeds payments to research sites and provides complete financial visibility to all study partners. Seamless integration with Veeva CTMS enables sponsors and CROs to streamline payment tracking processes within their existing trial management workflows, ensuring sites get paid on time with greater visibility and accuracy.
Key Capabilities of a Unified Application Suite

The Veeva Clinical Suite’s applications focus on managing study documents (eTMF), data and processes (CTMS), as well as tracking payments to sites, company sponsors, CROs, and other study partners. These applications provide authorized users full visibility into the complete study management process and a single source of truth for all study information.

Single application platform
Veeva’s suite of clinical applications is built on the Veeva 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 study processes.

Single source of truth for content and data
Enter study data and content once and leverage it across Veeva eTMF, Veeva CTMS, and Veeva Payments 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 study portfolio and processes through improved visibility, tracking and reporting. Make more timely, informed decisions, and accelerate timing from study to business impact.

About Veeva

Founded in 2007, Veeva NYSE (VEEV), is a global provider of industry-specific cloud software solutions that address the unique operating challenges and regulatory requirements of the consumer products and chemical industries. Our solutions help R&D, quality, regulatory and commercial teams eliminate inefficiencies and bring high-quality, safe, sustainable products to market without compromising compliance. Veeva is headquartered in the San Francisco Bay Area, with offices throughout North America, Europe, Asia, and Latin America. Learn more.