

## **Feature Checklist**

As trials and protocols increase in complexity, moving beyond Excel trackers to manage study start-up processes becomes critical to ensure on-time site activations. From event-driven automation that triggers downstream updates to unification with other clinical systems, this checklist highlights all of the features you should look for in a study start-up application to optimize start-up activities and get to First Patient First Visit faster.



Study Setup and Planning	Site and Country Activation
Milestone durations Predictive planning	<ul><li>Country start-up templates and workflows for 45+ countries</li></ul>
Site Qualification and Selection  Investigator, organization, and person database Site identification workflow Site identification searching and matching Site qualification and selection workflow Feasibility surveys  Essential Document Collection	Role-specific startup homepages Critical path management IRB/EC submission management Milestone auto-completion Ad-hoc events and milestones Contract and budget negotiation workflow Budget tracking Cycle time metrics
Site package distribution and document collaboration  Document management  Site and investigator document re-use  Auto-filing in Veeva eTMF  Shared workflows and data with Veeva CTMS in a single, unified environment	Site Greenlight Review and Approval  Site activation progress tracking Greenlight package approval workflow  Reporting and Visibility Metrics and cycle time tracking Configurable reports and dashboards  Application Security and Access Controls Granular profile and role permissions Object-level and workflow security

See a demo of all these capabilities in Veeva Study Startup today.



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