

[Accountability Logs](#)

Accountability Logs

Device Accountability Log

Investigators are responsible for maintaining strict control over investigational devices to ensure that the device is used only for subjects enrolled in the study. The [Device Accountability Log](#) helps maintain study device inventory and can be included in the regulatory binder.

Drug Accountability Log

Investigators are responsible for maintaining strict control over investigational drugs to ensure that the drug/biologic is used only for subjects enrolled in the study. The [Drug or Biologic Dispensing/Accountability Log](#) helps maintain an inventory of drugs used during the study and can be included in the regulatory binder.

Specimen Log

Safety laboratory assessments are usually part of most clinical trials that involve an investigational test article. Biological materials might be sent to the local laboratory or to a central laboratory. The [Specimen Log](#) helps track samples that are collected from study subjects.

Additional log resources can be found on the Partner's Healthcare website [here](#).

Need assistance or have clinical study management questions? Please contact [ResearchGo](#).

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 - [Clinical Research Information Systems](#)
 - [Clinical Research Business Partners](#)
- Group 2
 - [Office of Research Administration](#)
 - [Jonsson Comprehensive Cancer Center](#)
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