

EVADO INFORMED CONSENT

Evado Informed Consent service is a fully compliant paperless app that ensures that participants are fully informed before they decide whether to enroll in a trial or study. Evado eConsent has built in compliance monitoring that can be run on either a desktop or a tablet computer. Audio visual and cartoon graphics can be included to enhance participant comprehension. Evado's compliant signature collection has inbuilt auditing and meets current EU and USA regulatory and privacy requirements including GCP, GDPR and HIPAA. Evado's Informed Consent can be run as a stand-alone app or integrated into Evado OnDemand.

The eConsent service includes:

- Evado supports versioned consent forms to manage any changing consenting requirements throughout the execution of the study.
- Design tools assist in the creation of the informed consent page. A form can be broken into one or more sections or groups with each section or group containing one or more page fields.
- The eConsent page supports rich text, videos, images and quizzes.
- If preferred, the participant consent form can be completed at a clinic aided by local staff and signed off by the participant.
- Registered participants can complete the informed consent online remotely.
- Sites can monitor the progress of each participant's consenting process.
- Full audit-trails generated of all the participant consent data.

Evado e-Informed Consent

- All participant personal information is encrypted and time date stamped.
- Full privacy management ensures that only site users can view participant data.

The FDA Informed Consent guidelines state that the purpose of the research must be explained to participants including the participant's role in the study and how the study would run.


Evado Informed Consent isn't just a signature confirming that a participant will participate in a study. Informed consent opens up opportunities for participant recruitment and retention that keeps participants interested and involved in the study. This also benefits participants with low health literacy as the paper based systems can be confusing, alternatively eConsent ensures that participants are fully informed. There are long-term financial advantages for sponsors and participants benefit by being fully invested in the studies they participate in. Best of all, participants will comprehend and agree to participate in a study in an informed and meaningful way.

Informed Consent is suitable for remote studies as well as studies that are conducted in the clinic or hospital. Some participants prefer to upload their study data in their own home and Evado has provided a way for home based participants to remain anonymous if they so wish. Industry figures demonstrate that participants are much more likely to complete the trial if they use an Informed Consent application.

Contact us at hello@evado.com to schedule a demo or to find out more about Evado's Informed Consent app.

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