medrio



SOLUTION SHEET

Medrio eConsent

Efficiently manage consent, enhancing compliance and participant experiences.

Empowering organizations to streamline all aspects of the consent process, Medrio eConsent ensures improved process oversight and regulatory readiness.

The flexible, web-based solution supports both electronic and paper-based processes, allowing you to prioritize the patient experience and the sites' needs. Efficiently deliver consent forms, capture electronic signatures, and implement in-clinic, remote, or hybrid workflows.

Robust features like optional signatures, multi-signer support, and paper consent uploads add flexibility to consent processes for sites and patients. Medrio eConsent also provides a dashboard for tracking participant consent statuses and supports complex hybrid workflows within a study.

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| For the Participant: | |
| | s form. I have read it or it has been read to me. I have had my questions answered to my satisfaction. |
| | ned of any new findings developed during the course affect my willingness to stay in this research study. |
| Subject Name Subject One | × |
| Please sign here to consent | |
| 1 | |
| Please pass the device to the L | AR before 'Consenting'. |
| I have fully explained to the par | ticipant the nature and purpose of the above- iat are involved in its performance. I have answered |
| described study and the risks ti | ticipant the nature and purpose of the above- iat are involved in its performance. I have answered |
| I have fully explained to the part described study and the risks the all questions to the best of my a LAR/Witness Name | ticipant the nature and purpose of the above- nat are involved in its performance. I have answered billty. |
| I have fully explained to the part described study and the risks to all questions to the best of my a LAR/Witness Name LAR One | ticipant the nature and purpose of the above- nat are involved in its performance. I have answered billty. |
| I have fully explained to the part described study and the risks to all questions to the best of my a LAR/Witness Name LAR One | topant the nature and purpose of the above- at are involved in its performance. I have answered bolly. |



From consent deployment to re-consent, reduce manual tasks and improve oversight.

Quality

Ensure data meets ALCOA++ standards for easier audits and inspections.

Speed

Support faster time-tomarket with shortened enrollment timelines and regulatory findings.

Empowering flexible workflows

Built-in flexibility means minimal workflow disruptions for sites and more autonomy for participants. You can upload paper consent, download and print unsigned consent forms, and collect electronic consent across geographies with different regulations, making it easy to customize the consent process to any participant's preference.

Simplifying operations management

Medrio saves time and money with digital, often automated, consent processes, creating smoother study operations. You can easily automate participant consent requests and site document assignments, ultimately reducing site burden. Not to mention, builtin consent and IRB-approval statuses and dashboards facilitate hassle-free tracking and study oversight.

Improving participant experiences

Our easy-to-use interface enables better participant comprehension, improving retention rates and reducing study costs. Email and text notifications aid enrollment and engagement, while multimedia options, like videos and quizzes, enhance participant learning and understanding. It was very easy to use Medrio's system. In less than two weeks, and after just a couple of calls with the Medrio implementation manager, I felt comfortable creating and deploying eConsent on my own. It was very quick."

> – **Lucia Cesnakova,** DiMe Program Lead Digital Medicine Society

BUILD

- Upload paper consent forms for hybrid consenting approaches
- Consent deployment triggered by subject self-registration
- Document labeling and version control
- Require participant login credentials to input signature, if needed to comply with regulations in some EU countries

ADMINISTRATION

- In-clinic and remote workflows within a single study
- Single and multiple participant signatures workflows
- Email or SMS notifications

II. DATA REVIEW

- Built-in monitoring workflow
- eConsent data audit log
- eConsent status dashboard
- Single and multiple staff signatures

Ready to learn more?

