

Medrio CDMS/EDC

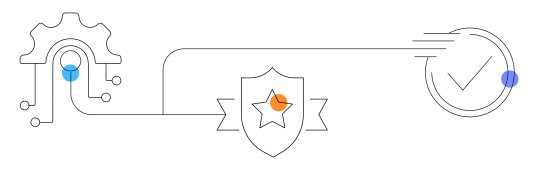
Maximize clinical trial efficiency and operational excellence with the *only* choice for clinical data management technology.

More than just a data collection tool, Medrio's intuitive user interface and robust back-end keep you in control of your study - from database build to your final report.

Advanced controls for improved data quality and regulatory readiness, all purpose-built for ease of use and efficiency, ensure you can conduct your study with confidence. The best part? Our unrivaled expert support and adherence to industry-leading compliance standards guarantee your data is in safe hands.

Plus, at Medrio, our customers come first. We actively engage with our customers, listening to their feedback and incorporating their needs into our innovative features and functionality, which continue to evolve alongside the industry.





Quality

Maintain autonomy and control of your study from end to end.

Control

Advanced controls for improved data quality and regulatory readiness.

Speed

Purpose-built for the ease of use and efficiency needed in today's environment.

Easy to learn, fast to deploy

Our intuitive user interface and eLearning modules allow users to feel comfortable after just one training session. With our code-free base and study copy functionality, set-up is quick and build speed increases over time. With Medrio, you can go live within hours to days, not weeks to months.

Mid-study changes made easy

Medrio's multi-tiered validation environment is the ultimate tool for managing mid-study changes. Medrio allows users to quickly develop and test how changes impact existing workflows before pushing live, ensuring seamless implementation without data migration or downtime. Choose Medrio for efficient, hassle-free mid-study change management.

Accessible data in near real-time

Medrio delivers near real-time access to data, empowering quick, informed action. Interactive dashboards give visibility into study metrics, which help teams monitor progress and respond proactively. With AI-enabled natural language processing (NLP), users can ask questions, receive actionable insights, and simplify analysis. Configurable roles and permissions ensure stakeholders access only necessary data. Single-click exports streamline collaboration with casebook creation in multiple file formats.

Ready to learn more?

By transitioning to Medrio, we have eliminated the amount of necessary source data verification by two-thirds. This has allowed CRAs to focus on other critical tasks, such as making sure adverse events and protocol deviations are handled correctly or ensuring that recruitment efforts are on track.

- **Frances Rubenstein, PhD.,**Director of Database

Management

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So BUILD

- Configurable, point-and-click builds
- Customizable user administration
- Pre-built study template library
- Study and form copy functionality

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- Data carry
- Edit checks and queries
- Always-on validation environment

II. CONDUCT

- Near real-time data access
- Targeted monitoring workflows
- Self-service AI-enabled reporting

1 LOCK

- Global regulatory compliance
- Flexible data export options
- Single-click submission casebooks

