



THE FIRST TRULY UNIFIED CLINICAL TRIAL SOFTWARE

Understand why omnia beats the competition with ease!

Save time and money while increasing data quality, simplifying study tracking, and achieving better outcomes.

INFINITE CAPABILITIES IN ONE SOFTWARE SOLUTION

More than just a clinical trial management system



All-In-One Solution

All professional clinical trial tools you will ever need fully integrated into one web-based software solution.



User-Friendly Interface

One simple user interface, no coding skills needed, drag-and-drop-functionality with very minimal training.



Real-Time Reporting

Enter data only once for real time data analysis. Gain instant insights for infinite biomedical statistics.



True Interoperability

Fully interoperable with other clinical trial systems. Import and export data with ease & connect third party solutions.



Infinite Trial Types

Realize all trials - Hybrid, decentralized, synthetic, real world data, global and even custom trial projects.



Infinite Scalability

Broaden research with limitless trials, storage, patients, users, sites, data points, documents, queries, ensuring expansion.



Data Security

You determine data storage and access preferences. We meet all regulatory standards, ensuring maximum security for data storage.



Access From Everywhere

Access data anywhere & work offline. Minimal software costs with our browser-based SaaS solution, ensuring flexibility & efficiency.



Professional Assistance

Professional trial services to ensure optimal trial setup, management and trial member training.



Convenient Customization

Customize EDC with drag-and-drop eCRF-Builder in omnia for effortless creation of documents, variables, and functionalities.

WATCH YOUR RESULTS

Cost reduction

-50%

Data quality improvement

+95%

Better patient outcomes



Time saving

+50%

Time to market reduction

-30%

EXPERIENCE COMPREHENSIVE SUPPORT ALONG YOUR MEDICAL DEVICE LIFE CYCLE

All solutions and services in one place

EDC | RTSM | eTMF | ePRO | eCOA | CTMS | eConsent | Lab Management | Imaging | eSource | Biostatistics | Medical Writing

Simple and competitive pricing. Pay per trial per month.

PROFESSIONAL CLINICAL TRIAL TOOLS

Discover the first all-in-one software solution enabling infinite clinical trials.

CTMS

Customized dashboard per user role · Standard study templates and logs · Key performance indicator (KPI) reports · Monitoring visit tracking · Timeline and milestone tracking · Contract management · Budget and payment management

EDC

Configuration · User access · Automated data integration and processing · IMP randomization directly within the eCRF document · Expert support team

eSource

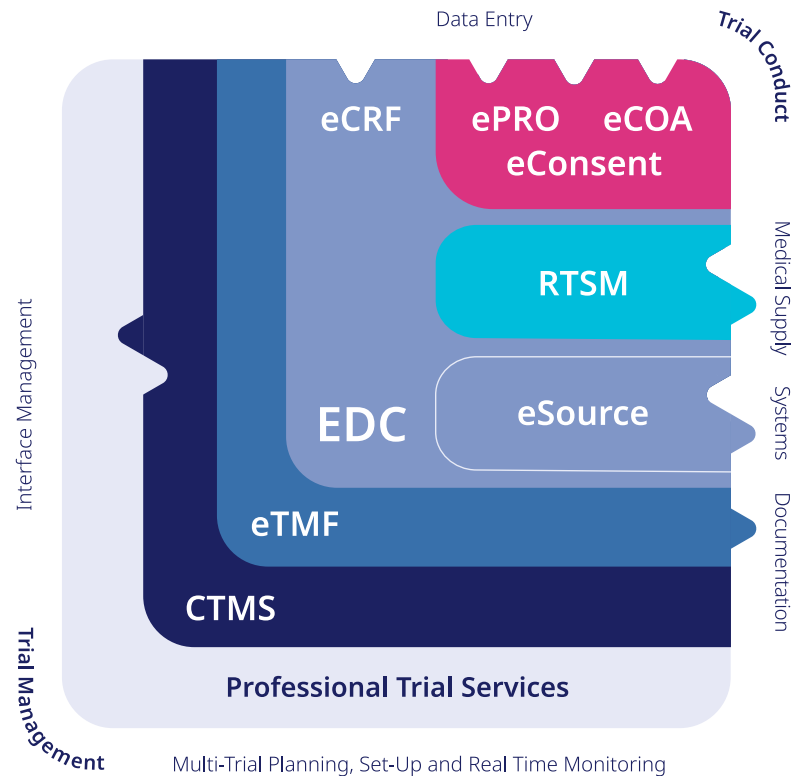
Digital data sources connectable to omnia EDC · Real-time data capturing · High data accuracy and enhanced clinical workflows · File management · Extensive configurability

RTSM

Complete lifecycle coverage for kit shipment and current status · randomization, allocation and accountability · Reduced manual data entry · Automated real-time reconciliations · Export of end-of-trial randomization report

eTMF

Insights through real-time reports · Rapid startup timeline · Exportable to fully validated eArchive · Convenient document management · Enhanced versioning and change tracking · Reliable access control and security



ePRO

Possibility for patients to directly report data · Involving electronic devices such as smartphones, tablets, or web-based platforms · Higher involvement of patients · Improved data quality

eCOA

Collection and documentation of clinical outcome assessments from patients, physicians or nurses · Involving electronic devices such as smartphones, tablets, or web-based platforms · Improved data integrity and efficiency

eConsent

Support of electronic and paper-based consent methods · Seamless eCRF and EDC integration · Multilingual support · Interactive Q&A module for trial staff communication · Digital signatures · Improved participant engagement

PROFESSIONAL TRIAL SERVICES THROUGHOUT THE COMPLETE TRIAL LIFE CYCLE

Plan

- ∞ Protocol Development
- ∞ CRF Development
- ∞ Clinical Advisory and Scoping

Conduct

- ∞ Clinical Data Management
- ∞ Pharmacovigilance
- ∞ Risk-Based Monitoring

Close

- ∞ Biostatistics
- ∞ SDTM and ADaM datasets and define.xml
- ∞ Medical Writing