

# EDC Product overview

Owned, Developed and Qualified by Replior





# Trial Online EDC

Trial Online's EDC is feature-rich, fulfilling the needs of even the most complex trials. Clinical sites appreciate the system's easy to use and intuitive approach. Quick study setup gives a short lead time for you to start collecting data.

Clinical Trial EDC systems are an important part of every clinical trial. Selecting the right EDC helps address inefficiencies on the operational side of research, from clinical trial planning stages through preparation, performance, and reporting.

There is a growing need to address the complex process of electronic data capture implementation and select systems with proven interoperability as more and more pharma and biotech sponsors start to recognize the potential opportunities that exist with EDC-ePRO integration.

Our EDC allows the customer to configure their solution entirely through the front end with our easy to use study builder

# Configuration includes:

- √ Role-based access
- √ Edit check validators
- √ Skip/add logic
- √ Derived calculations
- √ and more

## Introducing the new EDC Blue

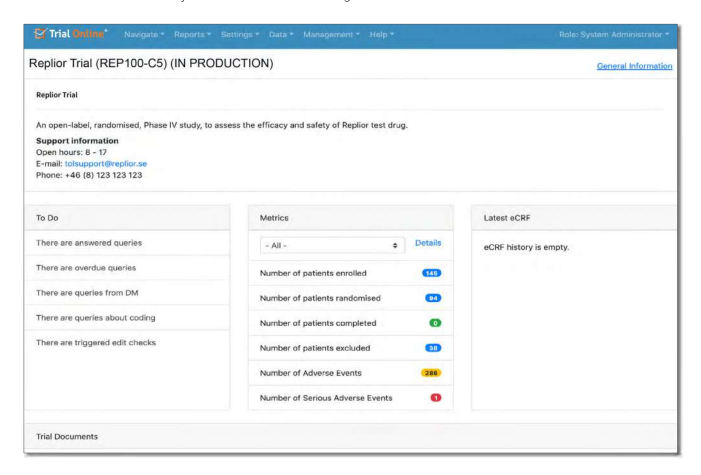
We have updated the design and layout of our EDC system, so now you will see a modern EDC system, which still has its easy to use and intuitive approach.

The new system is furthermore responsive, which means you will get the full experience no matter how you view Trial Online EDC, which also enables better tablet usage.



# Simple User Interface

Enable your site staff to enter data rapidly and accurately with logical user flow and form rules like edit checks. Calculated variables ensure that figures (BMI, imperial to metric, etc.) are always reliable. Data is saved automatically once it is entered ensuring no data is lost.



#### Web based

No special hardware or software is required to run the service, which utilizes standard internet browsers. Since the start-up time and creation of electronic Case Report Forms is very short, Trial Online can be used for all trials, regardless of size.

### Construction of eCRFs and edit checks

The eCRFs are created in a construction module that is a stand-alone software.

This module offers a fast and easy way to build eCRFs and edit checks, with flexibility in design and format. The module includes a library function where you can store and reuse standard forms. The eCRFs are easy uploaded in the Trial Online interface. Changes in the eCRFs during an ongoing trial is also possible and easy to handle.

## **Functions in Trial Online EDC**

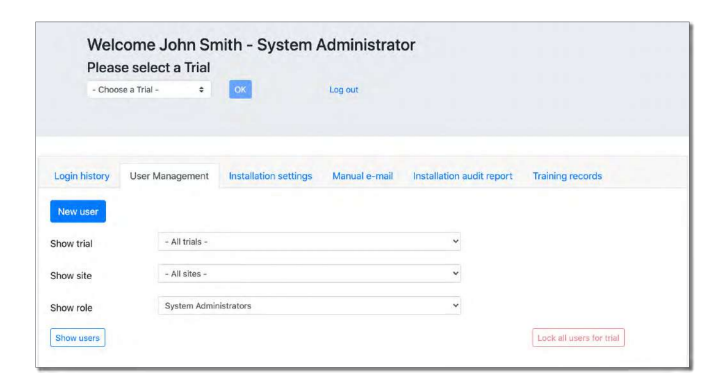
Trial Online EDC ensures the validation of data entry through edit checks that can be set to trigger directly when an "incorrect" entry has been entered, or in batches at regular intervals. Edit checks are easily specified in the Construction Module.

Trial Online offers a paperless query tool to generate, resolve and track queries online.

CRAs, Data Managers and Medical Coders can create queries and forward them to the investigator/coordinator who will be alerted within the system or via a system-generated email.

Laboratory files can be imported into Trial Online. Excel, SAS, ASCII and PDF are standard export formats in Trial Online.

It is possible to upload images, videos and other binary files in Trial Online. Files may be assessed by an independent investigator in the system



### **User roles**

All roles have access to Trial Online with defined permissions and passwords.

When logged on, the system provides a clear project overview, specific To Do list and email alerts.

- Investigator
- CRA/Monitor
- Sponsor
- Trial/System Administrator
- Assessor
- Coordinator
- Data Manager
- Medical Coder (MedDRA and WHO-DDE coding)



### Reports

Trial Online includes several types of reports, which simplify the overview, monitoring and security of the trial. In Trial Online the user can also create and customize additional reports.

The Status Report Summary gives the user insight to work flow navigation and a visual overview of patient status.

### **Audit trail**

A complete audit trail is maintained from data entry through all changes in the system.

The Trial Online system includes many types of reports that simplify the overview, monitoring and security of the trial and additional, customized reports are readily available.

# Regulatory submissions

From ICH/GCP and 21 CFR Part 11 to GDPR and HIPAA, we provide you with complete confidence that your data is compliant with our tools, so you can focus on the trial. To ensure data security over the internet, Trial Online features 256-bit HTTPS data encryption.











# **About Trial Online**

#### Trial Online EDC and Trial Online ePRO is owned, developed and qualified by Replior AB.

Replior is a privately held Swedish software solutions company focused on clinical trial support systems and offering a full suite of data collection products.

Replior develops in-house software and delivers Software as a Service, SaaS, solutions to Pharma companies and CROs.

The company is headquartered in Stockholm and has offices in Lund, Sweden, and Split, Croatia.

The EDC product Trial Online has been in operation since the year 2000, collecting data in over 1000 trials. Trial Online is a trusted eCRF provider for thousands of clinical studies made over the years, from small Phase I trials to global Phase III/IV trials. Trial Online ePRO provides Patient diary and questionnaire service to clinical studies.

#### About Trial Online FDC and ePRO

In early 2016, Trial Online ePRO was launched. Trial Online ePRO can be used as an integrated service with Trial Online EDC or as a stand-alone service.

In 2019, the Trial Online ePRO app and eConsent tool was launched.

Trial Online is owned by Replior AB as a part of the Hiberion Group.

#### Therapeutic Areas

Trial Online has conducted over five hundred studies in a wide range of therapeutic areas.

Our software can accommodate the most complex of clinical trial designs, with an easy and time-saving set-up. Conducting a study has never been easier.

- √ 100% Web-based EDC and ePRO
- √ Clients are able to perform all activities themselves
- √ Easy and fast set up of the complete system
- √ Fully compliant with 21 CFR part 11 and GCP
- √ Hosted on a secure dedicated server
- √ User-friendly
- ✓ Medical coding: MedDRA and WHO-DD
- √ Flexible and fast eCRF design
- √ Facilitates networking with specialists and partners
- √ Cost-effective even for small MedTech and BioTech studies
- √ Easy construction of Data Sets adaptable data export
- √ E-training



# **Quality and Compliance**

Our services and products shall always meet our customers' and users' expectations and requirements. By understanding our customers' needs today and, in the future, we deliver competitive services and products with the right quality, on time and according to agreed terms.

A cornerstone for our quality work is an engaged and aware staff and the company's collective knowledge and rules.

All systems and processes used to manage a clinical trial are part of a regulated process that must be validated and needs to meet FDA regulations, EMA regulations and ICH guidelines.

All Trial Online's products follows these regulations and guidelines as well as industry standards to ensure product quality. This includes Software Development Life Cycle and System Qualification.

Trial Online is a designed, developed, and tested computer system as defined in PIC/S 'Good Practices for Computerised Systems in Regulated "GxP" Environments'.

Trial online is compliant with FDA 21 CFR Part 11, GDPR and HIPAA. Our Quality Management System is built on quality standards in the industry, e.g. guidelines and directives provided by ISPE, FDA, EMA and ICH.

Our independent Group Compliance Officer performs regular internal audits on Replior and Complior. Clients (CROs and Sponsors) regularly perform audits of Replior and Trial Online.

We are dedicated to delivering a service of the highest quality in all aspects of our operation. Furthermore, we continually try to meet, or exceed, all expectations of our customers.

From development, through general product release, we have a high focus on quality and compliance. We have accomplished this by leveraging the latest technological solutions and according to regulatory standards and guidance.

## **Product Quality**

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# **Replior AB**

Trial Online is owned, developed and qualified by Replior AB.

We have offices in Stockholm-Sweden, Lund-Sweden and Split-Croatia.

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Schedule a demo

