



#### FOR MEDICAL DEVICE COMPANIES

# Zelta: a clinical trials platform

Control in every stage. Confidence in every outcome.

With the Zelta clinical trials platform by Merative, you are in full command of every aspect of your medical device trials and research — from designing workflows and forecasting costs, to building diaries for your participants. We empower you to take control in every stage and our solution is designed to help you accelerate trial outcomes with confidence.



## Choose technology designed to scale and accelerate your trials

Zelta<sup>™</sup> clinical trial platform is a unified clinical data management and acquisition platform with customizable modules that can be tailored to the unique needs of your clinical trials.

#### Scalable

- Control research of all types regardless of phase, therapeutic area or geographic location
- Host and scale thousands of trials around the world
- Create, standardize and scale processes to optimize cross-study control and reporting
- Help maximize international sites and patient engagement, supporting 60+ languages and dialects

#### Intuitive

- Make it easier to implement and execute research, manage participant compliance, perform routine tasks and report results to stakeholders through a single, user-friendly interface
- Optimize for sites and users
- Design trials with zero programming knowledge
- Take direct control of study go-lives, protocol amendments, study design changes and study closeouts

#### Unified

- Access modules and reports through a unified platform from anywhere in the world with single sign-on and one code base
- Remain current and ensure all of your trials and users are on the latest version of code with our single-instance technology
- Streamline clinical trial processes and help maximize patient, caregiver and provider engagement with clinical operations and patient and provider modules
- Eliminate managing and cross-checking multiple external systems with fully-integrated DICOM capabilities

#### Why Zelta?

The essential combination of trusted technology and human expertise.

#### Services

Whether you prefer self-service or full-service solutions, Zelta offers technology and deep medical device industry knowledge to help you overcome common challenges in clinical trials, whether it would be MDR, 510(k), or other medical device needs.

#### Data security

Our unified platform is hosted on a secure and flexible HIPAAenabled cloud.

#### Flexibility

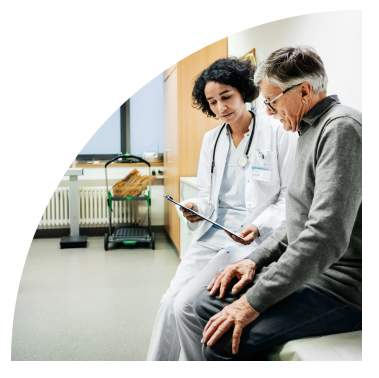
From small local studies to complex trials on a global scale, Zelta offers flexible functionality and pricing plans that can be customized based on your needs.

#### **Platform support**

Our team of certified, experienced designers are here to support you 24/7/365.

#### Trusted global partner

Experience supporting all phases of clinical research for over 10 years with study sites in more than 109 countries and across 23 therapeutic areas.



### Powerful modules and features

Built with all users in mind, Zelta modules are fully integrated and share one code base with the rest of the unified platform. This streamlines clinical trial processes and helps you maximize patient, caregiver and provider engagement to accelerate clinical trial outcomes. One client built their complete study database in as little as 4 days.

Read case study here  $\rightarrow$ 

Build and automate data connectors with minimal

Electronic Clinical Outcome Assessment (eCOA)

Engage directly with participants and caregivers via in

Increase efficiency by leveraging AI to build



#### Electronic Data Capture (EDC)

Design, validate and launch studies and apply amendments without database migration



#### Reporting and Analytics

Use pre-built or custom reports to derive single and cross-study insights



**Randomization and Trial Supply Management (RTSM)** Design and manage randomization and trial supplies in a single interface with minimal coding required



#### eConsent

Deliver quick and easy remote participant consenting without additional EDC integration



Digital Imaging and Communications in Medicine (DICOM) Transfer, view, take measurements on one fully Integrated system

app assessments and real-time analysis

## Choose technology designed to scale and accelerate your trials

Life science companies are using insights to improve outcomes:

600+

**Data Integration** 

Medical Coding with Al

consistency and reduce errors

coding

Medical device trials have been executed with Zelta

#### About Merative

Merative is a data, analytics and technology partner for the health industry, including providers, payers, life sciences companies and governments. With trusted technology and human expertise, Merative works with clients to drive real progress. Merative helps clients orient information and insights around the people they serve to improve decision-making and performance. Merative, formerly IBM Watson Health, became a new standalone company as part of Francisco Partners in 2022.

Learn more at <u>www.merative.com</u>

© Copyright Merative 2023

Produced in the United States of America, June 2022.

Merative, the Merative logo, merative.com are trademarks of Merative, registered in many jurisdictions worldwide. Other product and service names might be trademarks of Merative or other companies.

The performance data and client examples cited are presented for illustrative purposes only. Actual performance results may vary depending on specific configurations and operating conditions. THE INFORMATION IN THIS DOCUMENT IS PROVIDED "AS IS" WITHOUT ANY WARRANTY, EXPRESS OR IMPLIED, INCLUDING WITHOUT ANY WARRANTIES OF MERCHANTABILITY, FITNESS FOR A PARTICULAR PURPOSE AND ANY WARRANTY OR CONDITION OF NON-INFRINGEMENT. Merative products are warranted according to the terms and conditions of the agreements under which they are provided.

MCD-3077987612 Rev 5.0

Take the first step to boost the efficiency of your clinical trials.

Speak with your sales representative or read more about Zelta at <u>merative.com/clinical-development</u>

