

## 50%+ OF ALL FDA-APPROVED DRUGS SUPPORTED BY ERT OVER PAST 4 YEARS

# **CONFIDENCE AT EVERY TURN**

Minimize risk and uncertainty from clinical research to move ahead quickly — with confidence

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# MOVING AHEAD REQUIRES A SURE PATH FORWARD

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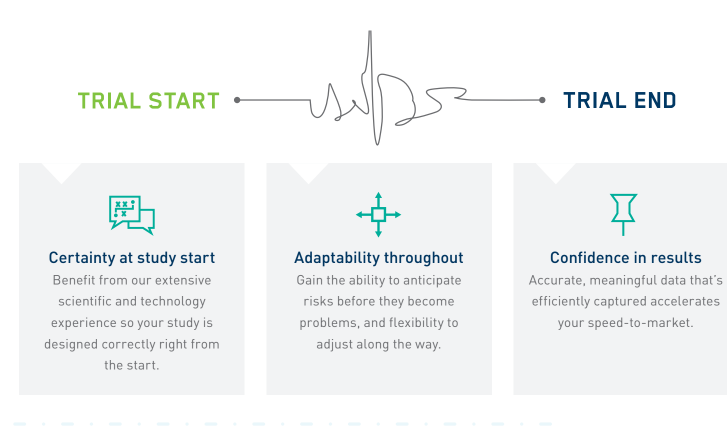
# PROACTIVELY MANAGE RISK INHERENT IN EVERY CLINICAL TRIAL

In clinical research, every detail matters. Trials themselves are inherently risky, and as trial complexity increases, new risks emerge. Mis-steps and inaccurate data can mean more expensive trials and delays in bringing life-saving treatments to the patients who need them most. Spotting trial risks before they become problems means you can bring clinical treatments to patients quickly — and with confidence.

AVOID COSTLY DELAYS BY SPOTTING RISKS BEFORE THEY BECOME PROBLEMS

## **REDUCE RISK & UNCERTAINTY TO MOVE AHEAD QUICKLY**

Every trial runs into unexpected challenges. With more than 9,500 trials under our belt, we've seen most of them before and have the expertise to put your trial back on course. This provides the flexibility and adaptability you need to accelerate through hurdles and get to market faster.



# **CONFIDENCE AT EVERY TURN**

#### Support for every phase of your trial powered by our proprietary ERT EXPERT® platform

#### ERT Trial Oversight

Run more efficient trials by unlocking hidden insights. With ERT Trial Oversight, you can assess site feasibility by leveraging historical performance data, mitigate risk faster without the burdens of a typical CTMS and achieve a new level of day-to-day trial management efficiency.

#### **ERT Site Optimization**

Generate high-quality data by optimizing site selection and performance. With ERT Site Optimization, you can prevent waste caused by low-performing sites, improve the reliability of assessments with customized rater training and drive quality results with flexible and comprehensive education and training programs.

#### **ERT Patient Engagement**

Put patients at the center of every clinical program. With ERT Patient Engagement, you can enroll patients faster, achieve a more comprehensive view of your patients and drive better compliance and retention. Help patients adhere to their treatment regimen by putting power directly in their hands to actively control their condition.

#### ERT Safety & Efficacy

Ensure patient safety and measure compound effectiveness by driving high-quality data. With ERT Safety & Efficacy solutions, you can accelerate your research with electronic clinical outcome assessment (eCOA) technology that doesn't get in the way, generate higher respiratory data quality and monitor every heartbeat with precision.

#### The ERT EXPERT technology platform

Improve trial visibility, performance and reduce time to market through confidence, agility and acceleration.



- > Confidence: Secure CRF Part 11 system ensures data is captured and stored in a repeatable manner that protects the patient, site and sponsor.
- > Agility: Ability to scale and adjust to growing complexity through systematic automation of repeatable processes while providing appropriate hooks to new data collection methods in a secure manner.
- Acceleration: Wide range of therapeutic area expertise to accelerate meeting your trial's unique needs.



WATCH THE VIDEO AT ERT.COM/CONFIDENCE TO LEARN MORE ABOUT HOW WE PROVIDE CONFIDENCE AT EVERY TURN.

## **PROVEN SOLUTIONS. PROVEN RESULTS.**

We're a data and technology company helping bring life-changing medicines to market faster by removing uncertainty and risk from clinical research. Partner with us to gain certainty at study start, ensure adaptability throughout and have confidence in your results.

#### Experts in navigating complexity

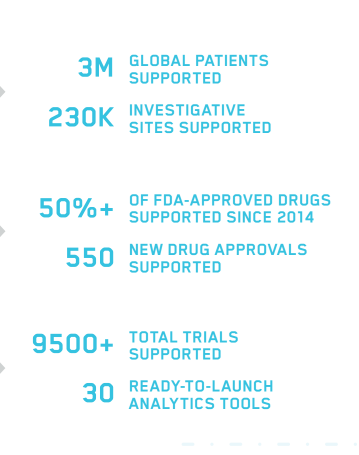
We can tackle the industry's most complex challenges. Seamless integration of any data source enables visibility into trial performance, risk identification and trends.

#### Adaptive problem solvers

Visibility to site performance across studies and programs highlights trends and actionable insights.

#### Anticipatory oversight

Our role is to protect the success of your trial before there's ever an issue. Near real-time access to study performance enables proactive risk mitigation.



### **BENEFIT FROM OUR EXPERTISE**

With more than 40 years of combined clinical, therapeutic and regulatory experience as well as technology and process-related insights, we balance knowledge of what works with a vision for what's next. This allows us to adapt to your needs without compromising standards. Rely on our expertise to help you bring clinical treatments to market faster — and with confidence. ERT

Learn how to minimize risk and uncertainty so you can move ahead quickly and with confidence. For more information, email <u>sales@ert.</u> <u>com</u> or visit <u>ert.com</u>.

### **ABOUT ERT**

ERT is a global data and technology company that minimizes uncertainty and risk in clinical trials so that customers can move ahead with confidence. With more than 40 years of clinical and therapeutic experience, ERT balances knowledge of what works with a vision for what's next, so we can adapt without compromising standards.

Powered by the company's EXPERT® technology platform, ERT's solutions enhance trial oversight, enable site optimization, increase patient engagement, and measure the efficacy of new clinical treatments while ensuring patient safety. Over the past four years, more than half of all FDA drug approvals came from ERT-supported studies. Pharma companies, biotechs, and CROs have relied on ERT solutions in 9,500+ studies spanning three million patients to date. By identifying trial risks before they become problems, ERT enables customers to bring clinical treatments to patients quickly — and with confidence.



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