

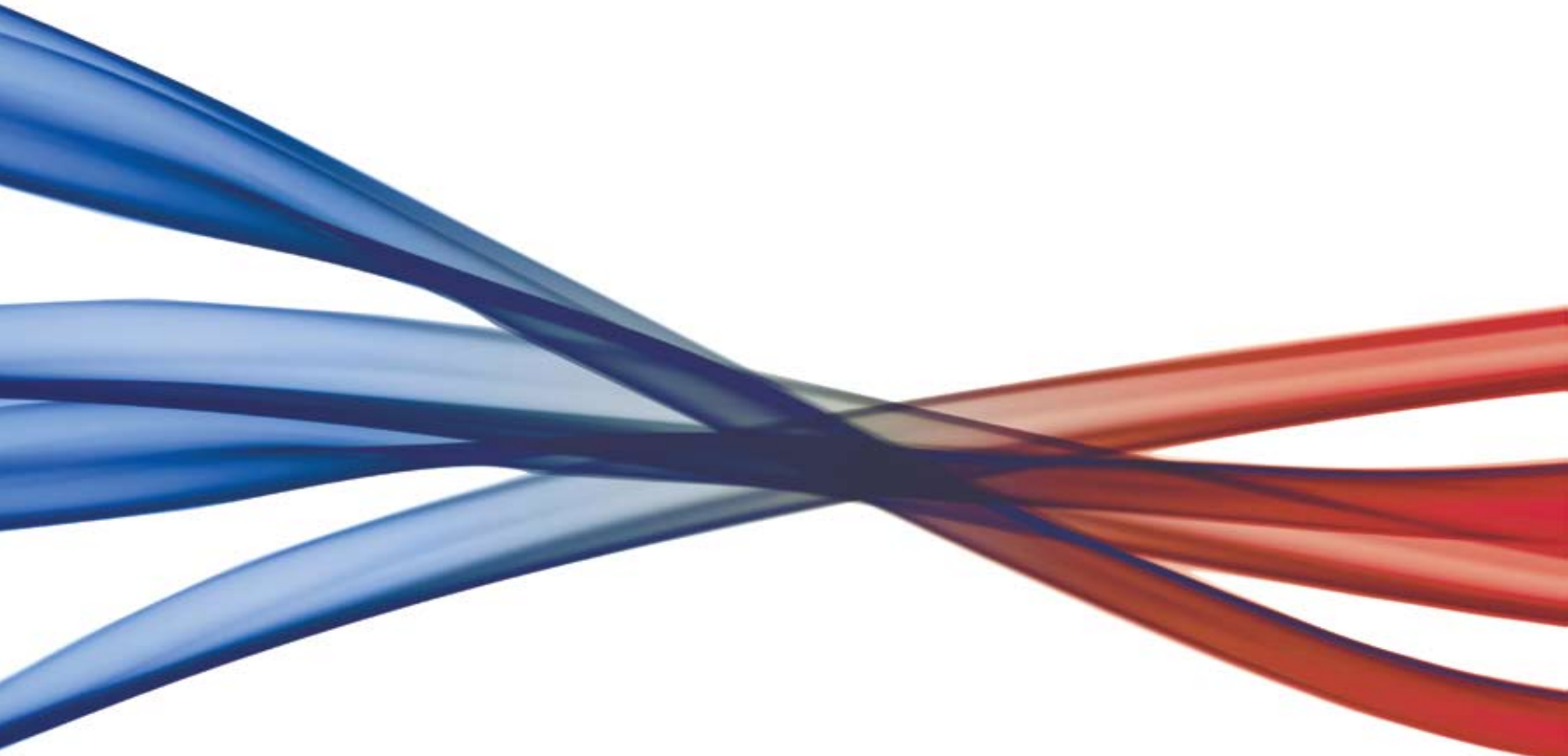


integrated intelligence



United BioSource Corporation

Evidence Matters[®]



integrated intelligence

Our Mission

United BioSource Corporation (UBC) is a global medical and scientific affairs organization that partners with life science companies to develop and commercialize their products.

We help generate authoritative, real-world evidence of product effectiveness, safety and value, to assist health care decisions and enhance patient care. UBC integrates scientific and industry experts, operations professionals and leading-edge technologies to provide innovative solutions across the product life cycle.

We strive to be an intelligent and nimble customer-focused organization that offers a unique combination of strategic vision, scientific expertise and operational excellence. With experts and operations in strategic locations worldwide, UBC offers the global reach and local expertise necessary to support the largest and most complex multinational product development programs.

Our clients can expect UBC to design and deliver high-quality solutions tailored to meet their unique needs.



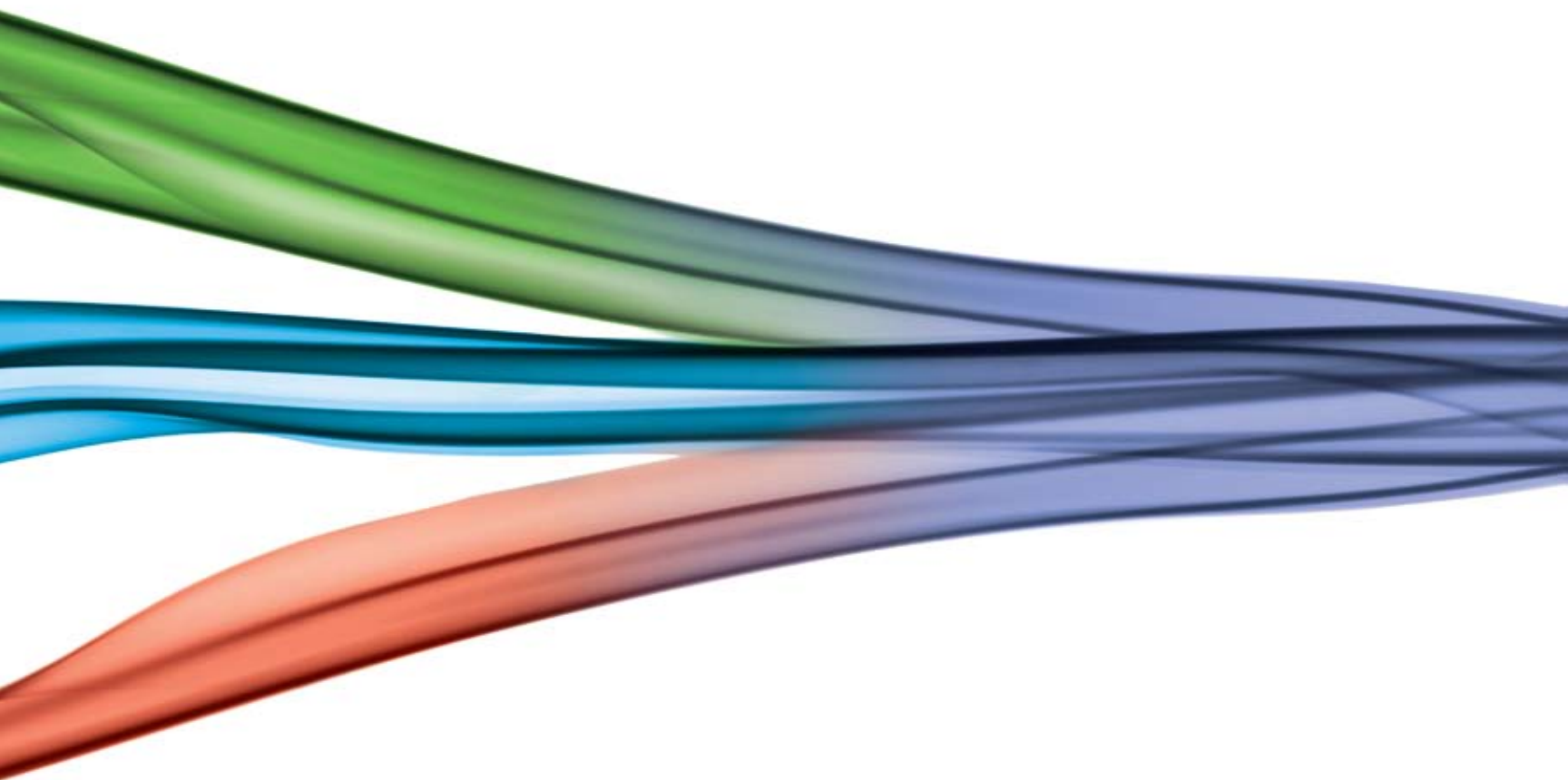
UBC designs and implements clinical development programs, including peri- and post-approval programs, large streamlined safety studies and registries. In addition, we offer specialized research services and technologies to help clients generate real-world evidence of product efficacy and safety.

We provide industry-leading scientific and operational expertise, advanced technology and global capabilities to deliver comprehensive evidence generation solutions for our clients' products.

Our experience:
10+ years of product, disease and pregnancy exposure registries.
20+ years of clinical development in peri-approval programs.
25+ years of governmental and industry studies.

streamlined science

- Peri-/Post-Approval Programs
- Large Streamlined Studies
- Phase IIIb-IV Solutions
- Registries
- Comprehensive Risk Management Solutions
- Adaptive Trial Solutions
- Biostatistics & Programming
- Investigator Services
- Patient Recruitment & Retention
- Site Selection & Retention



UBC designs and implements an array of training programs and offers many unique research support products to help clinical research sponsors ensure data integrity, reduce variability and improve signal detection across all phases of product development.

Our objective is to help study sponsors collect accurate and reliable data that will meet study objectives and are relevant to clinicians and regulators.

measurement precision

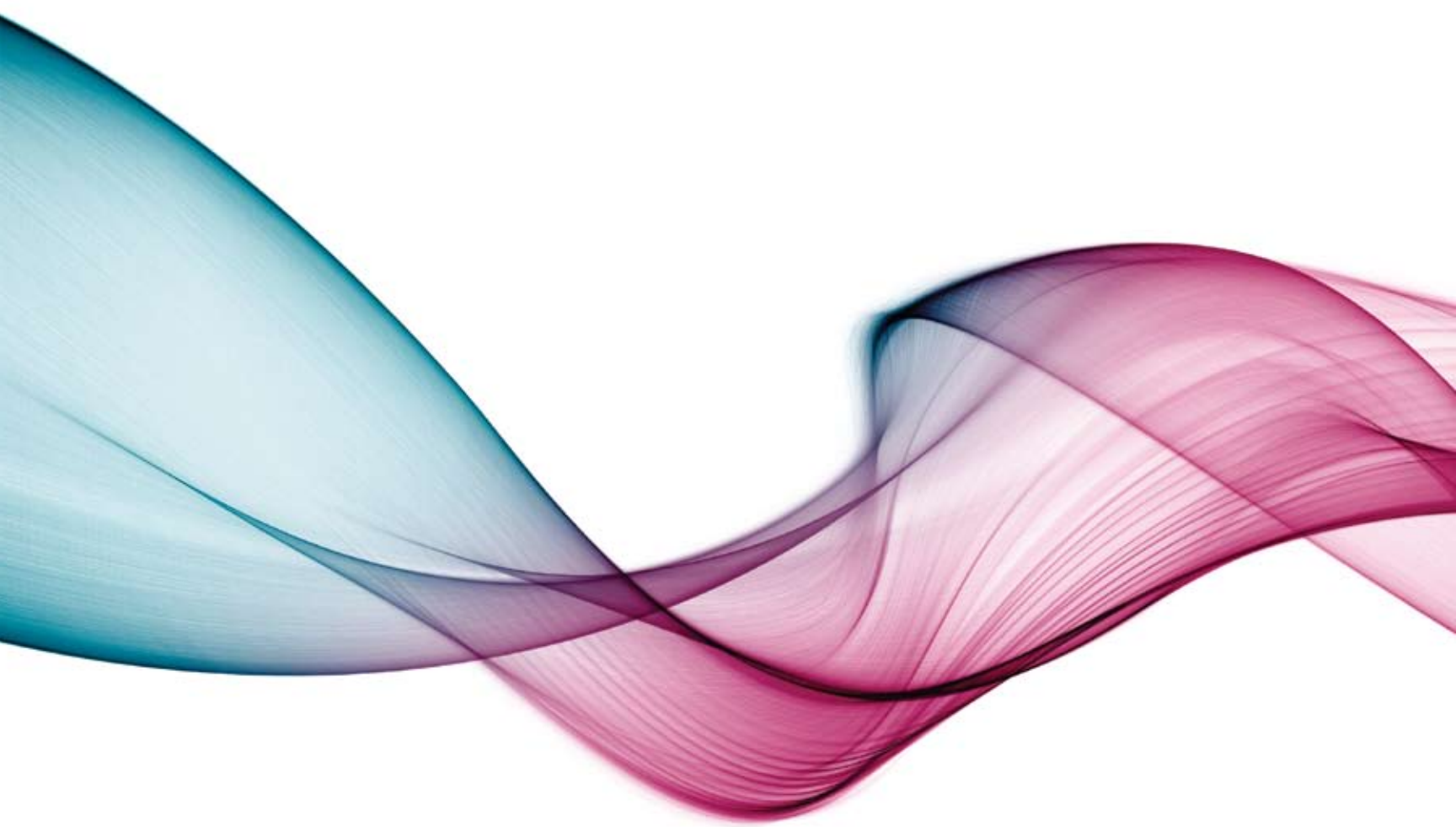
Inter-Rater Reliability Measuring
Rater Training, Certification & Rater Monitoring
Scale Translation & Acquisition
Computerized Cognitive Testing
Centralized Rating
Research Site Profiling & Credentialing
Training of Investigators, Physicians & Site Personnel
Web-Based Investigator Meetings
Patient Recruitment & Retention
NDA/MAA Preparation



Demonstrating product value is essential in today's competitive, cost-focused health care environment. UBC integrates the best scientific expertise and technologies to help clients build and implement strategies to prove the value of their products. Our experts in health economics, outcomes research, pricing and reimbursement, and epidemiology build strategies and deliver solutions to facilitate appropriate access to medicines, devices and other medical technology throughout the product life cycle.

accelerated value

Health Economics
Health Outcomes Research
Market Access
Pricing & Reimbursement
Patient Assistant Programs
Epidemiology
Health Care Data Capture
Database Analysis



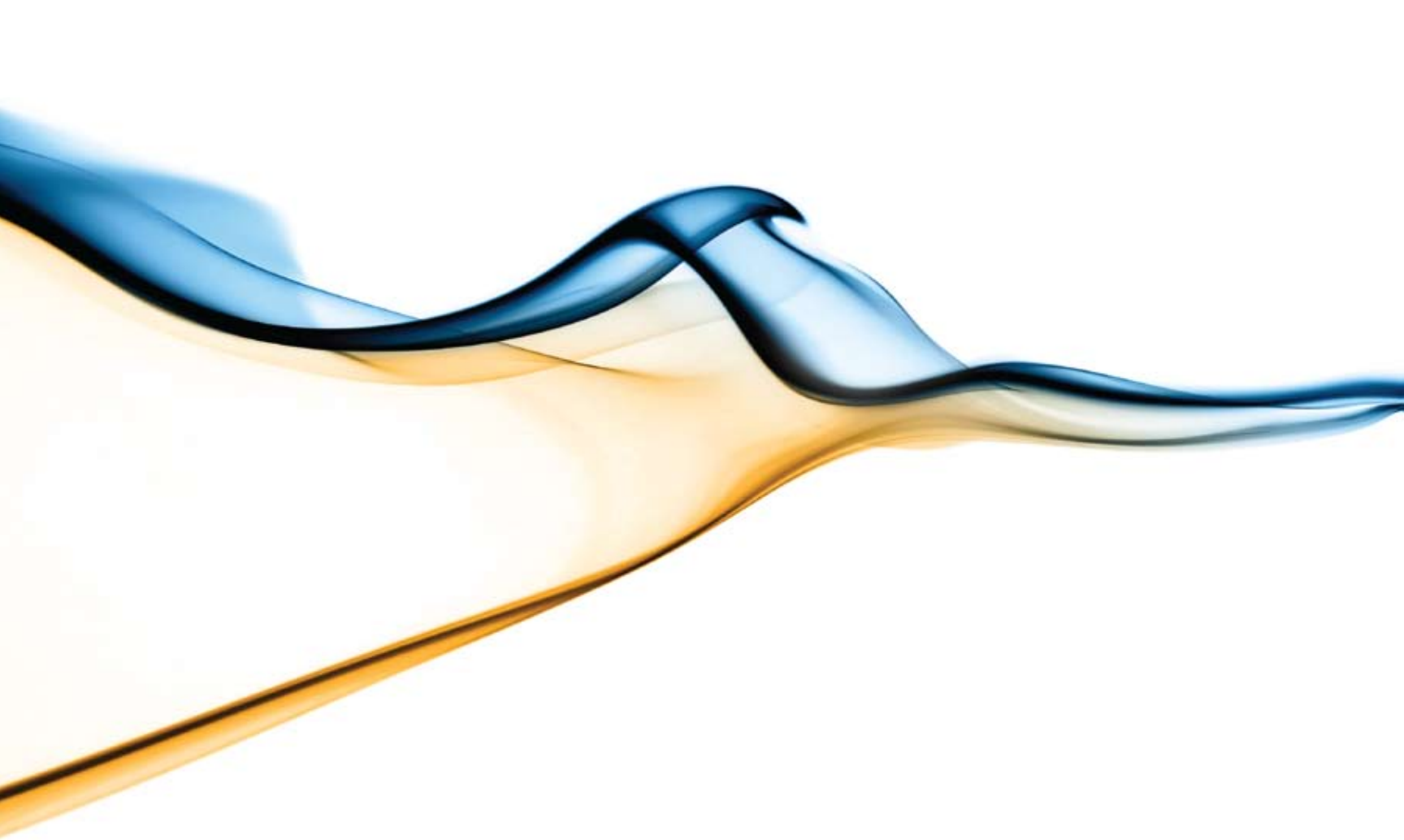
UBC offers world-recognized scientists and industry experts with extensive experience and a depth of knowledge necessary to help clients successfully develop and commercialize their products.

We provide a contemporary and in-depth understanding of the health care landscape to help clients address complex medical, regulatory and health policy challenges facing manufacturers of biopharmaceuticals, medical devices and other life science products.

We leverage this experience and knowledge to help clients make smart, pragmatic development decisions to help ensure the success of their products.

extensive expertise

Evidence Generation Planning
Regulatory Strategy
Meta-Analysis & Systematic Reviews
Safety & Risk Management Strategies
Adaptive Trial Solutions
Medical Communications & Publication Solutions
Health Science Policy



UBC understands that product safety is a top priority and a complex challenge for companies that manufacture and market life science products. Our experts design and implement comprehensive strategies to identify, assess and minimize product safety risks.

Our solutions are designed to protect patients, comply with regulatory requirements, and facilitate appropriate commercial positioning for our clients' products throughout the development life cycle.

minimized risk

Risk Evaluation & Mitigation Strategies (REMS)
Large Streamlined Safety Studies
Registries
Risk Assessment
Pharmacovigilance
Safety Processing
Data Monitoring Committees



UBC offers an array of proprietary technologies and unique research tools designed to accelerate and optimize product development and commercialization. We offer research products and tools that increase clinical research efficiency, automate data collection, accelerate information movement and enhance data accuracy. In addition, UBC provides leading-edge software and solutions to ease medical publications planning and implementation.

Our products and tools are designed and continuously updated based on extensive experience and deep knowledge of clinical research and product development.

innovative tools

Electronic Data Capture (EDC)
IVRS & IWRS
Data Warehouse
eLearning & Virtual Meetings
ePLAS
eRegistries
eROs, ePROs & eDiaries
Adaptive Trial Solutions
Datavision™ & Visiontracker™
iPledge
Technology Integration

UBC at-a-glance:

1,400+ employees
30% hold advanced science degrees
2,000+ peer-reviewed publications
Trials conducted on six continents

Global results:

365,000 patients
82,000 study sites
3,000 clinical protocols
2,500 health outcomes studies
30,000 investigators trained in 70 countries
50 languages

U.S. locations:

Ann Arbor, MI
Bethesda, MD
Blue Bell, PA
Chantilly, VA
Glastonbury, CT
Kansas City, MO
Langhorne, PA
Lexington, MA
McLean, VA
Morgantown, WV
Newtown, PA
Philadelphia, PA
San Francisco, CA
Seattle, WA
Southport, CT
Wayne, PA

International locations:

Budapest, Hungary
Cologne, Germany
Geneva, Switzerland
Goring-on-Thames, United Kingdom
Horsham, United Kingdom
Kiev, Ukraine
London, United Kingdom
Montreal, Canada
Moscow, Russia
Prague, Czech Republic
São Paulo, Brazil
Tokyo, Japan

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