



Advancing Research to Enhance Outcomes

Driving innovation that transforms patient care

Clinical Research Advancement is dedicated to transforming healthcare by delivering innovative and seamless execution of clinical research initiatives globally.

Embracing our philosophy of personalized solutions, we provide our clients with a comprehensive "full-service CRO" experience, emphasizing strategic recruitment and retention approaches that drive the successful advancement of clinical studies and achieve exceptional outcomes.

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Our Expertise & Focus

At Clinical Research Advancement, our vision is to accelerate clinical research worldwide, transforming scientific innovation into accessible, life-changing therapies. We are committed to creating a landscape where clinical trials are efficient and rooted in scientific excellence.

To realize this vision, we prioritize innovative recruitment and retention strategies that ensure trials are driving successful study outcomes.

- **Recruitment and Retention Excellence:** Delivering customized, data-driven strategies to effectively engage diverse patient populations worldwide. Our proven success rate includes enrollment increases of 40% and above in global studies, driving faster trial progress and better outcomes. Contact us today to discover how we can accelerate your clinical research success.
- **Strategic Site Identification & Feasibility Assessment:** Advanced stakeholder mapping and disease-state insights optimize site selection, enhancing activation speed and patient enrollment quality.
- **Key Opinion Leader (KOL) Development:** Deep engagement with KOLs and healthcare providers drives patient referrals, study advocacy, and informed protocol design.
- **End-to-End Clinical Trial Operations:** Comprehensive capabilities across study start-up, regulatory affairs, clinical monitoring, medical affairs, safety/pharmacovigilance, data management, biostatistics, medical writing, and quality assurance. Our global sites are regulatory-ready and initiated 30% faster, optimizing the study start-up phase to deliver maximum efficiency and accelerated timelines worldwide.

Clinical Research Advancement offers comprehensive, end-to-end clinical trial services designed to support every phase of your drug development journey.

Our clinical trial operations include streamlined study startup procedures and clinical monitoring to ensure data integrity and patient safety throughout each trial. The CR Advancement leadership team brings over 25 years of combined experience with leading global pharmaceutical corporations and top-tier CROs, having held roles at Eli Lilly, Sanofi, Takeda, Boehringer Ingelheim and AstraZeneca, as well as at organizations such as IQVIA, ICON, PRA, Labcorp, PPD and Medpace. CR Advancement employs specialized medical affairs teams to support scientific communication and strategic engagement.

Our therapeutic expertise covers Oncology, Respiratory, Cardiovascular, Endocrinology, Infectious Diseases, and Rare Diseases.

Our experienced global medical affairs team enhances these capabilities by maintaining rigorous standards of documentation and operational excellence, ensuring your studies are executed with efficiency.





Our Approach

The foundation of successful clinical trials rests on the timely and effective selection, recruitment, and retention of study participants. At Clinical Research Advancement, we understand that each step—from identifying eligible subjects to maintaining their engagement throughout the study—is critical to achieving superior trial outcomes within targeted timelines.

Our proprietary Recruitment and Retention Model leverages deep community engagement, strategic partnerships, and data-driven methods to ensure participant enrollment targets are met swiftly and ethically. We prioritize building trust with geographically diverse patient populations, including underserved and minority groups, by collaborating closely with recruitment specialists, faith leaders, and key opinion leaders to foster meaningful connections and awareness.

Our approach is embodied in the "Five Cs" of Clinical Excellence that guide every trial:

- Coordination – Seamless integration of all stakeholders, ensuring smooth operational flow.
- Control – Proactive risk management to protect milestones and timelines.
- Compliance – Rigorous adherence to regulatory and ethical standards worldwide.
- Collaboration – Cultivating strong partnerships among sponsors, sites, KOLs, and patient communities.
- Communication – Transparent, open, and consistent information exchange to accelerate problem-solving and decision-making.

We employ a comprehensive, end-to-end subject engagement strategy that includes:

Community Outreach: Mobilizing recruitment specialists and partnering with faith-based organizations and local health networks to reach potential participants where they live and seek care.

Site-Centric Coordination: Conducting meticulous site identification, feasibility assessments, and qualifications that align with the study's therapeutic focus and target demographics.

KOL and Healthcare Professional Involvement: Engaging Opinion Leaders and key stakeholders to advocate for each study, enhance patient referrals, and support retention efforts.

Continuous Patient Follow-Up: Maintaining participant involvement from enrollment through to last patient last visit (LPLV) with dedicated follow-up visits, education, and communication that reduce attrition.

By focusing on these core principles, Clinical Research Advancement consistently achieves recruitment and retention results that ensure clean, robust data and accelerate drug development processes. Our experience, combined with innovative decentralized trial methods and culturally tailored engagement strategies, allows us to overcome common enrollment challenges and foster patient-centric clinical studies that enhance access and engagement.



Driving Global Impact Through Strategic Insight & Collaboration

A Leader in Clinical Research Innovation:

We pioneer personalized, data-driven recruitment and retention strategies that address the complexities of modern clinical trials — ensuring access to patients and trial feasibility are seamlessly integrated.

Global Reach with Local Expertise:

Our expansive global footprint enables effective trial execution across North America, Asia Pacific, the Middle East, Africa, and beyond. **Clinical Research Advancement** operates extensively across the US, Asia Pacific, Middle East, and Africa, maintaining local offices and affiliates in Korea, Taiwan, Hong Kong, Malaysia, the Philippines, Australia/New Zealand, China, Thailand, Indonesia, Vietnam, India, Japan, and Pakistan. This local presence ensures compliance with evolving regulatory landscapes while respecting cultural nuances critical to patient engagement and site performance.

Therapeutic Versatility:

From Oncology to Rare Diseases, and Cardiovascular to Infectious Diseases, our talented clinical teams bring depth in therapeutic knowledge and operational excellence that adapts to the scientific demands of each protocol.

Experienced Leaders and Specialty Teams:

With over 25 years of collective clinical research experience, our multidisciplinary experts — including clinicians, recruitment specialists, regulatory affairs professionals, and data scientists — collaborate with sponsors to deliver pragmatic solutions that advance drug development priorities.

Technology-Enabled Solutions:

Leveraging cutting-edge digital health technologies, including decentralized and remote trial models, we streamline clinical operations, reduce patient burden, and accelerate data collection in a secure and compliant framework.

At CR Advancement, we build the future of medicine — faster and with an unwavering commitment to quality and patient-centricity.

Early & Late Phase Research Expertise

Clinical Research Advancement specializes in Tailored Solutions for Early Phase Studies

Early-phase research requires rapid adaptation, precision, and operational scalability. Recognizing these needs, **Clinical Research Advancement** provides scalable solutions designed to maximize efficiency while advancing key study objectives such as dose-finding and signal detection. Our collaborative approach ensures that we develop and implement strategies best aligned with your organizational goals and scientific requirements.

Scalable Late Phase Expertise

Translating lessons learned during early development into effective late-phase strategies is crucial for success. Our integrated model combines global reach with regional insight to facilitate seamless scaling of your trials. This hybrid expertise leads to measurable improvements in patient enrollment, retention, and data quality, ensuring regulatory readiness and faster timelines.

Comprehensive Recruitment & Retention Solutions

The cornerstone of successful clinical trials is effective recruitment and retention. **Clinical Research Advancement** offers a comprehensive, end-to-end approach combining strategic outreach, community engagement, site support, and ongoing patient communication to optimize participant involvement and study completion rates.



Our Ethos

Central to our organization is an unwavering dedication to advancing clinical research. We apply a rigorous, evidence-based methodology to enhance recruitment and retention efforts effectively.

Empowering Participation

Expanding access to patients across diverse geographic regions is fundamental to the future of clinical research. By amplifying patient experiences from multiple communities, we strive to transform the clinical research landscape and ensure broad participation in clinical trials.

Your Partner For Comprehensive Clinical Research Services

Founded in 2010, Clinical Research Advancement brings over 25 years of combined clinical research expertise, having supported more than 60 clients and sponsors across over 200 clinical projects worldwide.

Accelerate your clinical development and elevate healthcare outcomes by partnering with Clinical Research Advancement. With a proven history of building trusted relationships with key opinion leaders and clinical sites, we deliver comprehensive, end-to-end CRO services tailored to your unique needs—from study startup and regulatory support to clinical monitoring, study closeout, and final report completion.

Leverage our global expertise, personalized strategies, and seamless execution to ensure efficient patient recruitment, retention, and high-quality data across every phase of your clinical development program.

Engage with Clinical Research Advancement today to explore how our end-to-end solutions can drive your next study forward with the highest standards of precision, integrity, and operational excellence.





To Accelerate Your Next Clinical Trial, Contact Us

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