Interventional Studies

Evidera’s interventional study team, made up of dedicated peri- and post-approval experts, can design and execute efficient, fit-for-purpose studies that help you meet regulator requirements and achieve optimal commercialization. With deep knowledge of post-market requirements and endpoints, our experienced team can navigate those requirements to position your product for future success.

Dedicated Team, Global Power

Our team leverages the global power, therapeutic expertise, and deep clinical development experience of PPD to seamlessly transition earlier phase studies to late-phase studies. We utilize PPD’s network of physicians and product development experts to execute studies focused on operational excellence. With extensive experience across all interventional study types, we can execute cost-effective trials across the globe drawing on our experience in more than 70 countries, including China and Japan.

Positioning Your Product for Future Success

Our peri- and post-approval experts can help you conduct efficient, optimized studies in order to:

- Build a larger body of evidence around a product or specific patient population to better inform clinical practice
- Enable optimal commercialization and market access for your product
- Generate evidence to address peri- and post-approval evidence gaps
- Meet regulator and payer requirements for post-marketing effectiveness and safety

Our Offerings

- Phase IIIb/IV studies
- Early access programs (EAP)
- Digital and virtual trials
- Compassionate use programs (CUP)
- Extended access programs (XAP)
- Pragmatic/adaptive trials
- Investigator-sponsored/initiated trials (IST/IIT)
Deep Therapeutic Expertise
Our capabilities span 16 therapeutic areas and 115+ disease areas in the past five years with deep expertise in oncology, neurology, infectious diseases, and endocrinology.

![Therapeutic Expertise Chart]

Our Experience

290+ Interventional Studies executed in the past five years

115+ Disease areas in which studies were conducted

Approximately 113,000+ patients have been involved in past interventional studies

Conducted studies across 70+ countries

Leveraging Real-World and Patient-Centered Data
When designing and executing studies, we leverage Evidera’s multi-disciplinary team of scientists and evidence generation strategists to leverage relevant real-world data and high-quality patient-centered data. Our Patient-Centered Research team is the largest dedicated team of its kind in the field and can work with our interventional experts to incorporate clinical outcome assessments (COAs), patient-reported outcomes (PROs), patient preferences, and the patient voice into studies.