

YOUR TRUSTED PARTNER



800+ FTEs



100%
HCPs on delivery teams



Safety reporting in
100+ countries

OUR SERVICES PORTFOLIO

Assuring drug safety to pharma companies who are looking for pre/post marketing compliance & reporting solutions

Pharmacovigilance Services



Provided end-to-end PV set up for 200+ Biotech and Pharma companies since 2007



287,000+ cases processed in 2024



Provided consulting / advisory support services for 90+ clients in 2024

E2B R3 compliant safety databases

QPPV Services

RMP & REMS Compilation & Implementation

Safety Reporting, Aggregate Writing & Signalling

Industry Standard Processes & SOPs with Quality Overlay

ICSRs Processing & Literature Surveillance

Medical Review

Medical Information Services



24x7 multi-channel Global Contact Centers with resilient infrastructure

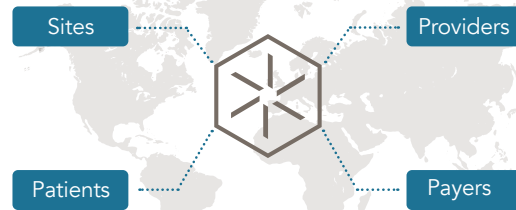


38,000+ inquiries handled in 2024



Medical information coverage in 50+ countries, 30+ languages, scalable to 100+ languages

Integrated PV & MI Response Centers



HCPs available 24X7 AE/MI/PC handling

MEDICAL WRITING

- Clinical Writing
- Regulatory Writing
- Scientific Writing

REGULATORY

- Pre Approval Services
- Submissions
- Life Cycle Management

QUALITY

- Audits • Training
- Inspection Readiness
- CAPA Management • Consulting

TESTIMONIALS

“ We appreciate APCER for its efficiency in responding to matters that they get even at the eleventh hour. Working with APCER has made compliance a lot easier. ”

– AVP, Pharmacovigilance, US-based pharmaceutical company

“ APCER’s invaluable support with clinical trial disclosure deadlines helped us in timely submissions of periodic reports. ”

– Head, Clinical Trial Transparency, Top 10 pharmaceutical company with operations in the EU and US



APCER
LIFE SCIENCES

Together for better health
Part of APC Group

APCER Life Sciences is committed to improving health in partnership with its clients. We bring together safety, medical, regulatory and technology resources to ensure that patients receive the safest, most effective therapies possible.

We are an ISO 9001:2015, ISO 27001:2022, and ISO 27701:2019 certified company

Learn more at www.apcerls.com or contact us at one of our global offices:

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Medical
Information

Regulatory
Affairs

Quality
Assurance

Medical
Writing

Pharmacovigilance



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