

Your Partners in Drug Development and Safety

Specialist services to the pharmaceutical industry spanning all phases of clinical trials, post-approval pharmacovigilance and medical information.



Specialist Drug Development and Safety Solutions

Founded in 1997, Ergomed is dedicated to the provision of specialist services to the global pharmaceutical industry.

Today, Ergomed supports pharmaceutical companies with services spanning all phases of clinical trials, post-approval pharmacovigilance and medical information.

Our business includes a full range of high-quality clinical research and clinical trial management services and internationally recognised expertise in orphan drug development together with an industry-leading suite of specialist pharmacovigilance solutions.

By providing this full-service offering, Ergomed enables emerging and established life sciences companies to meet their regulatory obligations, maximise their drug development success and therefore their product value.

17
offices
worldwide

53
countries with
active clinical
trials

1000+
professionals

600+
studies
completed

100+
supporting
products in over
100 countries

300k+
ICSRs
processed
per year.

Transforming Drug Development

Ergomed, our Clinical Research Organisation (CRO) provides complete Phase I-IV clinical development and trial management services, assisting clients by providing complete solutions tailored to their unique requirements.

With experience in over 600 trials, Ergomed has planned, managed, monitored and reported clinical trials with a range of technologies that include small molecule drugs, monoclonal antibodies and other targeted agents as well as cancer vaccines, immunotherapy, radioactive agents and photodynamic therapies.

Assisting clients with project directorship, project management, regulatory affairs monitoring, safety and medical monitoring, data management, biostatistics, medical writing, site management support and study physician support.

20+
years'
experience

25%
staff with
PhD or MD

- Site management program specifically designed to increase study performance
- Specialist expertise in orphan drug development
- Therapeutic specialisations: oncology, respiratory, neurology and orphan.



Discover the Orphan Advantage

PSR is the leading expert in assisting biotech and pharmaceutical companies in orphan drug development and supports the Ergomed global CRO offering.

Our orphan disease toolkit reflects a genuine family and patient approach to recruitment and retention. Our close links to patient advocacy groups maximise our partner's chances of success.

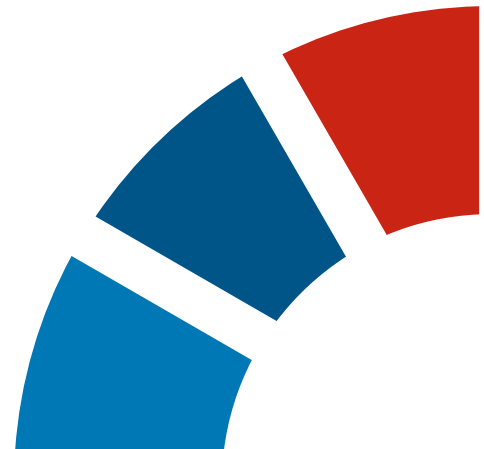
PSR is specialised in designing and executing complex clinical development programs requiring innovative regulatory and clinical approaches in Europe and the US.

Through our site management model and study physician team support, we find hard to locate patients around the globe and work with the investigative sites, creating the best designs to maximise clinical programs and registries.

10+
years'
experience

170+
orphan
projects

- Thought-leaders in the orphan drug community
- True family and patient centricity and Patient Organisation Advisory Board
- Global reach and access to patients with rare and ultra-rare diseases.



Transforming Drug Safety

PrimeVigilance provides global, top quality, cost-effective, innovative Life Cycle Management Services to enable emerging and established life sciences companies to meet their regulatory obligations and to maximise product value.

PrimeVigilance covers the entire product life cycle, assisting clients with the effective management of their drug safety information, and offering expert consulting services from former regulators and opinion leaders.

Our drug safety services include: case management, signal management, risk management, pharmacoepidemiology, audits, services of qualified persons for pharmacovigilance, training, strategic advisory, literature searches and medical information services.

10+
years'
experience

25+
QPPVs

- Global leader in QPPV services
- Automisation in pharmacovigilance expertise
- Choice of leading drug safety databases
- Regulatory experts and key opinion leaders.



Real World Evidence

Strengthen the value of your products through Real World Evidence.

Real World Evidence brings credibility to your product's study results, as it satisfies the industry's ever growing need for more information about the real-life safety and effectiveness of medicines.

Ergomed provides a unique set of services specifically tailored to the specialised needs of pharmacoepidemiology and Real World Evidence generation studies.

Therapeutic Specialties include: oncology/haematology, neurology/CNS, and allergy/respiratory and orphan.

60+
studies

2k+
sites

37k+
patients

- Complete services for observational research, including PASS, registries and other peri-approval programs
- Expertise in various study types addressing clinical, medical affairs and market access objectives
- Unique fit-for-purpose operational strategy using dedicated processes and tailored real-world data collection solutions.

Your guide and support along the drug development continuum.

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