Your Clinical Research Organisation for Clinical Studies

As an independent company for Clinical Research Organisation (CRO), we have been offering high-quality cooperation in carrying out studies since 2006. Up to now we have supported more than 1000 medical studies and are happy to witness a continuously increasing interest in our services.

CRO Dr. med Kottmann focuses on non-interventional studies (NIS), case series as well as clinical studies in the phases II to IV and hereby achieves high-quality and cost-effective solutions for research projects with pharmaceuticals or medical devices.

Basically, the entire project management – ranging from study planning, writing of monitoring plan / test plan to database creation (compiling of CRF or eCRF, e.g. with secuTrial, OpenClinica or xclinical), determination of sample size, statistical analysis to clinical monitoring and medical writing – is carried out under the direction of a medical specialist. Our literature research identifies high-quality studies featuring high evidence levels. Thus, the company’s main focus is clinically orientated research – thanks to many years of clinical experience, particularly significant results can be achieved. Here we also gladly give impulses in Investigator Initiated Trials (IIT), in order to create scientifically sound and appealing study designs together with the study doctors, offering excellent feasibility.

We work according to Good Clinical Practice (GCP) and observe the declaration of Helsinki as well as the required laws such as MPA or MPG. Our CRO attaches great importance to quality standards – thus, we are ISO 9001 and DEKRA certified and furthermore hold a data protection certificate. Please find more information regarding our quality management here.

How can we help you?