(Senior/Principal) Statistician (m/f/d)

We are offering a permanent or freelance position as Statistician (m/f/d) for our locations in Sulzbach (Frankfurt/Main) and Marburg, or home-based.

RESPONSIBILITIES
- Analysis, and reporting of clinical trials and preparation of integrated summaries
- Authoring of Statistical Analysis Plans (SAP) including design of table, listing and graph (TLG) shells
- Input to dataset specifications and review of outputs for analysis and reporting of pre-clinical/clinical trials or pooled analyses
- Quality control (QC) of programmed files used for the reporting of clinical trials, preparation of submission dossiers, ad hoc requests by health authorities, or publications
- Statistical input to clinical trial reports, integrated summaries and (e)CTDs
- Collaboration with other trial team functions (such as data management, statistical programming, pharmacovigilance, clinicians, or medical writing) throughout the clinical trial conduct and submission preparation
- Communication and presentation of statistical methodology and analysis results to clients, investigators, regulatory authorities, scientific communities, and non-statisticians (e.g., clinicians)
- Contribution to (medical) publications
- Training of team members on statistical methodology, processes, and tools
- Participation in industry working groups, conferences, and scientific meetings
- Development of standard operating procedures (SOPs) and contribution to continuous improvement

QUALIFICATIONS
- Masters’ or equivalent/higher degree in Statistics, Biostatistics, Mathematics, or similar subject
- Strong knowledge in statistical methodology
- Analytical and problem-solving skills

Interested in joining us? apply@mainanalytics.de

ABOUT MAINANALYTICS
We are a well-known team of highly-motivated statistical programmers and biostatisticians with a wealth of experience.

Founded in July 2019 as an employee-owned CRO, we offer comprehensive expertise for analysis and reporting of clinical studies – from preclinical to phase IV, and in various indications. We also support legacy data conversions, CDISC implementation, and submissions to global regulatory agencies.

Our professionals keep up-to-date on industry and regulatory standards as well as statistical methodology through continuous training. Several members of our team contribute their expertise in international, cross-organizational working groups and committees.

Teamwork, flexibility, timeliness, and accuracy are our strengths at mainanalytics.

We look forward to receiving your application. For any questions or to apply, please contact:
apply@mainanalytics.de

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Qualifications – continued

- Expertise in one or more programming languages (particularly SAS and/or R)
- Experience in clinical research and working in a strongly regulated environment (GCP) preferred
- Strong interest in clinical data analysis
- Flexibility to work independently and as part of a team and to interact with multidisciplinary scientists
- Ability to focus on accurate and timely completion of project-related and competing tasks
- Willingness to (further) develop knowledge of industry standards and new statistical methodology
- Excellent English verbal and written communication skills

BENEFITS

- Flexible work structure
- Option to work remotely
- Flat hierarchy
- Collaborative and positive team spirit
- Dedicated resources for further professional development