



Position: Medical Monitor, Clinical Science (Contractor – EU)
Reports to: VP, Clinical Development EU
Location: London, UK

Summary

The Medical Monitor (Consultant) will be responsible for working as part of the medical team to design, execute and report clinical trials in food allergy and subsequently pursue a planned MAA. The position would suit a physician who enjoys the phase III study environment and has MAA/BLA experience.

Specific Responsibilities:

- Execute clinical development plans and protocols for clinical trials in food allergy
- Medical monitoring of clinical trials and provision of medical expertise to the team
- Provide clinical trials expertise to support to study teams and participate in multidisciplinary meetings
- Contribute to the analysis and discussion of study results
- Author key clinical documents, including protocols and study reports
- Contribute to the preparation of documents required for regulatory submissions
- Develop relationships with investigators
- Ensuring compliance with GCP across clinical trials
- Work closely with cross functional colleagues, including Clinical Operations, Biometrics, Patient Safety and Regulatory counterparts

Qualifications / Requirements:

- Medical degree and experience in patient care required
- At least 5 years of pharmaceutical or biotechnology industry experience including working in late-stage clinical development and, preferably support of MAAs
- Additional training/experience in clinical immunology and allergy or other immunological disease areas preferred
- Thorough understanding of GCP and familiarity with relevant EMA, FDA and ICH guidance
- Working knowledge of statistics as applied to clinical trial design and analysis
- Experience in writing and editing scientific research reports

Preferred Experience, Special Skills, Knowledge:

- Dedication to the conduct of clinical trials that generate high quality data and ensure patient safety
- Collaborative and accessible
- Preference for working in a fast-paced, team-oriented, environment
- Strong commitment to goals and timelines
- Ability to absorb new information quickly and gain command of relevant literature
- Desire for new professional challenges

- Possessing excellent problem-solving & decision-making skills; strives for appropriate balance between taking initiative and seeking input from others
- Ability to analyze and summarize complex data and information concisely
- Excellent written and verbal communication skills, including fluency in English

About Aimmune Therapeutics, Inc.

Aimmune Therapeutics is a clinical-stage biopharmaceutical company founded to address the unmet medical need in food allergy, which currently has no approved treatments. Our mission is to improve the lives of people with food allergies, based on our proprietary desensitization treatments in development. Aimmune's recently announced positive top-line data from its pivotal Phase 3 PALISADE trial for AR101, the company's lead investigational drug for peanut allergy. Headquartered in the heart of San Francisco's biotechnology hub, Brisbane, California, Aimmune has additional offices in the Kings Cross area of London, and in Raleigh, North Carolina.

Qualified candidates should forward a resume and cover letter, including a statement of interest, availability, and experience to Human Resources (careers@aimmune.com) with the job title in the subject line.

Aimmune Therapeutics is an Equal Opportunity Employer.
Principals only; no recruiters please.