Bioanalysis, LC-MS/MS
Study Director

Job offer

**AnaPath Research**

AnaPath Research is a CRO located in Barcelona, with extensive experience in carrying out preclinical trials for pharmaceutical laboratories, the chemical industry and other research organizations. In our more than 30 years of activity we have worked with the main pharmaceutical industries being part of different multinationals (RCC, Harlan and Envigo). In November 2019, AnaPath Services acquired the company and re-founded it as AnaPath Research, thus undertaking together a new project of scientific quality and close contact with new and old sponsors.

With a multidisciplinary team of scientific experts, AnaPath Research covers most fields of preclinical pharmaceutical development and chemical and food safety.

**Analytics & Bioanalytics Unit**

The AnaPath bioanalytical team supports the industry R&D projects all the way from drug discovery, through preclinical development and up to the clinical phase. Whether it is method development or routine sample analysis, we are dedicated to providing accurate, reliable and streamlined bioanalytical services for the most challenging programs to help our clients make confident research decisions.

The Analytics & Bioanalytics Unit provides services to regulated bioanalytics by LC-MS/MS and ImmunoAssay platforms. Additionally, we also provide support to the preclinical drug development by performing ADA assessment, flow cytometry and qPCR analysis. Finally, we also assist our toxicology studies by carrying out dose formulation analyses.

**Position**

We are looking for a Study Director specialized in the LC-MS/MS field to work in our Analytics & Bioanalytics Unit and to also give support to dose formulation analysis by HPLC-UV. The job holder will be responsible for the scientific direction of the studies assigned to them ensuring that all work is conducted in accordance with the study plan, standard operating procedures (SOPs) and appropriate regulatory standards (such as GLP & GCP) and guidelines (BMV, FDA, EMA, SANCO etc.).

**Responsibilities**

Technically, the Study Director will have the scientific responsibility for study plan design and approval; oversight of data collection and data interpretation, analysis and documentation; providing of study progress updates to Sponsors, reporting of results and drawing of corresponding study conclusions.

Additionally, from a business perspective, the study director will oversee that the study is cost-effective and meets budgetary constraints.

Finally, the Study Director will have the GLP responsibilities detailed in the OECD Principles on Good Laboratory Practice guidance.

**Requirements**

- BSc, MSc , PhD in Chemistry/Biochemistry/Pharmacy
- Experience in liquid chromatographic technique and UV and MS/MS detections
- Proven experience in the pharmaceutical industry or similar
- Experience working under/following GLP principles
- English level: Advanced/Proficiency
- Good communication skills
- Immediate start
- Own vehicle

**Contact**

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**Terms of employment**

- Indefinite contract
- Full-time
- Morning work shift