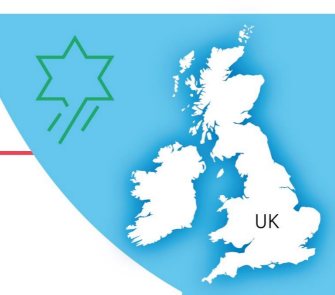
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Lab Manager

Department: QC

Hours per week: 37.5

Basis: Permanent



Job Purpose

As Laboratory Manager — Biochemistry, you will be responsible for managing and leading a team of skilled and experienced QC Scientists. You will oversee all Biochemistry tests (Elisa, SDS-PAGE, Western Blotting and other immune assays) performed on the allergen product portfolio and ensure that the laboratory is running effectively and developing relationships across ATL.

Key Responsibilities

- Line management of the Biochemistry team, ensuring you are a role model for the wider QC function and working to develop relationships with peers, stakeholders and the wider ATL group
- Taking a lead in both strategic and operational improvements across QC in line with the QC transformation strategy
- Taking overall responsibility for Health and Safety in the QC labs, ensuring that processes are reviewed on a regular basis to ensure compliance and ensure continuous improvement
- Overseeing the planning, coordination and daily testing activities in collaboration with the team
- Acting as main point of contact for all requests coming into the Biochemistry team and managing workflows
- Participating in Problem Solving / Root Cause Analysis
- Optimisation and streamline of processes using LEAN approach
- Setting direction by clarifying mission and purpose and translate department goals into team activities
- Obtaining and increasing the knowledge and experience within the team, including development of team
 members and creating a working environment that fosters enthusiasm, initiative and empowerment
- Deputise for QC Manager where appropriate

Knowledge, Experience & Skills Required

- Previous experience in a similar role you will be comfortable championing and promoting the team
- Strong background within a leadership role managing a GMP regulated team in a lab environment
- In depth knowledge of compliance, method validation, GMP and ICH guidelines
- Able to write GMP protocols/reports and other key GMP documents
- Experience of using visual planning tools.

If you are interested in this role, a job description is available on request.

Please send all applications to: careers@allergytherapeutics.com

