

# QC Analyst

Job ID REQ-10051724

5月 12, 2025

Switzerland

### 摘要

We are seeking a motivated QC-Analyst for Biopharmaceutical Production within the framework of raw materials, drug substance, and in-process controls. The candidate will perform analytical testing and ensure compliance with SOPs, analytical methods, and compendial standards.

#### About the Role

Major accountabilities:

- Conduct and coordinate quality control tests on biologics drug substances (Physicochemical testing,
   e.g. HPLC, Capillary Electrophoresis, UV) ensuring compliance with regulatory requirements
- · Independent planning, implementation, and evaluation of routine and special analyses in a GMP-regulated environment

- · Interpret test data, prepare detailed reports, and maintain accurate record of test results.
- · Troubleshoot testing procedures and make recommendations for improvements, with a focus on HPLC and Capillary Electrophoresis
- Conducting microbiological tests such as total germ count determinations (MET) and bacterial endotoxins (BET)
- · Participate in the validation of analytical procedures
- Collaborate closely with the internal teams to optimize quality control processes
- · Instrument responsibilities, including qualification, maintenance, and calibration documentation
- Support in ensuring that the laboratory is maintained in a ready state of inspection.

#### Key performance indicators:

- · Timely test record completion and accurate processing without delays
- Prompt reporting of missed deadlines and aim for shortest possible lead times
- Continuous readiness for inspection
- · Consistently follow the GMP and GSU guidelines, and SOPs, ensuring no critical irregularities
- Proactively identify and implement cost-reducing optimizations
- Complete all assigned training as per the provided plan

#### Minimum Requirements:

- · Completed scientific education (e.g., Laboratory Technician, Bachelor or Master)
- Practical experience in a GMP-regulated lab and document creation
- · Knowledge in common analysis techniques, especially HPLC and photometry; microbiological knowledge is an advantage
- Working experience in laboratory environment in the pharmaceutical industry
- · Good IT skills (MS Office) and laboratory software like LIMS, Chromeleon, Empower are an advantage
- Ability to work precisely, independently, and proactively
- · Reliability, flexibility, resilience, and strong teamwork skills
- Shift work with normal working times (one shift) including weekends

#### Skills:

- · Continuous learning
- Dealing with ambiguity
- Decision making
- · GMP
- Industry standards
- Laboratory equipment
- Laboratory excellence
- Quality Control (QC) testing
- Quality Control sampling
- Self awareness
- Technological expertise

#### Languages:

· Fluent in German (spoken and written) and proficient in English

#### Skills Desired

Continued Learning, Dealing With Ambiguity, Decision Making Skills, Gxp, Industry Standards, Laboratory Equipment, Laboratory Excellence, Quality Control (Qc) Testing, Quality Control Sampling, Self-Awareness, Technological Expertise

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Operations	
Business Unit Innovative Medicines	
地点 Switzerland	
站点 Basel (City)	
Company / Legal Entity C028 (FCRS = CH028) Novartis Pharma AG	
Functional Area Quality	
Job Type Full time	
Employment Type Regular	
Shift Work No	
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