U NOVARTIS

QC Technician

Job ID REQ-10040017 Feb 07, 2025 Italien

Summary

The QC Technician performs the analysis on batches and incoming materials according to cGMP rules and SOPs.

About the Role

Major accountabilities:

- Perform the analysis of batches following specific training;
- Promptly report to the Quality Control Supervisor and the Qualified Person any deviation and/or out of specification detected during the analysis activities.
- Collaborate with the Quality Control Head/supervisor for the CAPA implementation in the quality control department.
- Control processes, equipment and area.
- Perform routine maintenance and cleaning, periodic microbiological verification of pharmaceutical areas quality control equipment and support external specialized personnel in carrying out extraordinary maintenance/qualification activities.
- Perform incoming and outgoing verification activities of raw materials and materials.
- Collaborate with the Quality Control Head to the management of material stocks and waste materials.
- Collaborate with the Quality Control Head to the training of new personnel.
- Execute the calibration, verification, qualification and validation activities of the Quality Control processes and equipment.

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- Execute the calibration, verification, qualification and validation activities of the Quality Control processes

and equipment.

Essential requirements:

- Degree in CTF, Pharmacy, Chemistry.
- Previous experience in a similar role within a GMPenvironment.
- Available to work in shifts, including night shifts.
- Fluent in Italian. Good knowledge of English.

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Division Operations Business Unit Innovative Medicines Standort Italien Site Ivrea Company / Legal Entity IT58 (FCRS = IT058) Advanced Accelerator Applications Italy Srl Functional Area Quality Job Type Full time Employment Type Regular Shift Work No <u>Apply to Job</u>

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