

Production Technician I (Weekend PM)

Job ID REQ-10048268

USA

May 08, 2025

Summary

This role is located on-site in Indianapolis, IN. Novartis is unable to offer relocation support for this role: please only apply if this location is accessible for you.

Production Technicians play an active role in daily production of Radioligand Therapies (RLT) as well as setup and preparation of instruments and equipment. The Production Technician adheres to regulatory requirements while performing job functions, executing production as per batch records and SOPs. Responsibilities are performed within a team and according to an assigned production shift schedule. The Production Technician works closely with the Manager and Lead to ensure production is executed in a safe and timely manner.

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Key Responsibilities:

- Executes all activities related to the manufacturing of RLT products. Responsibilities include operating and maintaining grade A isolators, focusing on KPI goals as well as ensuring all state, federal and Novartis radiation safety guidelines are adhered to.
- Responsible for successful on-time completion of required training curriculum comprising of the necessary Standard Operating Procedures (SOPs) and Aseptic Techniques, Gowning Qualifications and other relevant training including HSE for the specific role.
- Supports all technical aspects related to production readiness including manually cleaning the cell and performing sterilization of the isolators. Conducts routine and dynamic environmental monitoring as required.
- Prepares all materials while maintaining material identity in accordance with the batch monitoring system as defined by procedure.
- Ensures all cGMP compliance activities are followed.
- Participation in assigned qualification/validation activities, and assist on deviation investigations and inspections, as necessary.
- Prepares applicable documents and records such as batch records, shipping documents, and training materials.

Shift: Week 1Thurs-Sun, 6:00pm-6:00am, Week 2 Fri-Sun 6:00PM-6:00am. This position may involve mandatory overtime as needed.

Essential Requirements:

- Bachelor's degree in relevant Engineering or Scientific discipline is highly preferred; If the
 applicant does not have a degree, a minimum of 1+ year' of experience in cGMP or aseptic
 environment is required.
- Knowledge of cGMP regulations and FDA guidance applicable to aseptic manufacturing is highly preferred.
- Ability to gown aseptically and work in a clean room environment (Grade C) area for extended periods of time is required.
- Near vision performance should be the equivalent of 20/20 with no impairment of color vision.
 The use of corrective lenses to achieve the desired visual acuity is permitted.
- Ability to lift or carry up to 35 pounds
- Radio Pharma experience preferred.

Languages:

• English.

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部門 Operations
部門 Innovative Medicines
国 USA
State

Indiana

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Company / Legal Entity U469 (FCRS = US469) AAA USA Inc.

Functional Area Technical Operations

職種

Full time

雇用形態 Regular

Shift Work No

Apply to Job.



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