

SCM Expert

Job ID REQ-10027859

Oct 30, 2024

USA

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.

At Advanced Accelerator Applications, a Novartis company, we are committed to leading innovation in nuclear medicine and delivering the next generation of targeted radioligand therapy to cancer patients. We are looking for experienced Manufacturing professionals to help us reach our ambitious goals.

Our SCM Expert is responsible for the site Master Data processes aimed to ensure accurate and timely creation and maintenance of the master data in the ERP System, in accordance with local and global requirements. The Material Planner drives the tactical purchasing process for all materials to ensure materials availability and execution plan feasibility in accordance with Site inventory policy.

About the Role

Major Accountabilities:

- Master Data -Set up and perform complex local And global master data set up And establish process, guidelines, business matrix and SLA time lines in in GxP environment in adherence to Novartis compliance.
- Be Subject Matter Expert in the assigned area of work -Support data collection and reporting
 of KPIs -Logistic, WAndD -Responsible that all logistic processes are proceed in a timely,
 high quality, efficient and effective manner and in full compliance with all laws and supply
 chain management policies and procedures.
- Identify and drive continuous improvement projects.
- Ensure material availability in line with the approved production plan.
- Ensure daily MRP oversight for all BOM material, analysis of requirements and Purchase Orders management.
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- Own, in ERP System, MRP relevant data and materials technical specifications and ensure no Purchase Orders are past due in the ERP System.
- Management of purchased item Complaints/Returns to supplier.
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- Provide a load-balanced dispatch list for incoming materials to the warehouse and Quality department that ensures these activities are completed in line with the production needs.
- Control and follow-up of supplier's service level indicators: Quality OTIF, lead-time violation.
- Lead action plans to achieve supplier delivery performance targets and drive for continuous improvement.
- Reporting of technical complaints / adverse events / special case scenarios related to
 Novartis products within 24 hours of receipt -Distribution of marketing samples (where
 applicable) -Provide guidance and system support for returns, rebates and credit/debit notes
 -Define and decide on priorities for physical deliveries in collaboration with other functions
 (e.g. LEX) for 3rd party customers and cross divisional supplies -Act as SPOC for escalation
 issues for assigned customers -Influencing and connecting relevant stakeholders to speed up
 (system) issue resolution -Material Planner -Ensure management of daily MRP exception
 messages And appropriate follow up.
- Provide a load balanced dispatch list for incoming materials to the warehouse and Quality department that ensures these activities are completed in line with the production needs.

Minimum Requirements:

- Education: Bachelor's degree required.
- 3+ years of pharmaceutical experience.
- 3+ years of project management or governed change control process experience.
- 5+ years of experience working with ERP system. Preferred: dynamics or SAP.

Commitment to Diversity & Inclusion: Novartis is committed to building an outstanding, inclusive work environment and diverse teams' representative of the patients and communities we serve.

Why Novartis: Helping people with disease and their families takes more than innovative science. It takes a community of smart, passionate people like you. Collaborating, supporting and inspiring each other. Combining to achieve breakthroughs that change patients' lives. Ready to create a brighter future together? https://www.novartis.com/about/strategy/people-and-culture

Join our Novartis Network: Not the right Novartis role for you? Sign up to our talent community to stay connected and learn about suitable career opportunities as soon as they come up: https://talentnetwork.novartis.com/network

Benefits and Rewards: Read our handbook to learn about all the ways we'll help you thrive personally and professionally: https://www.novartis.com/careers/benefits-rewards

EEO Statement:

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Accessibility & Reasonable Accommodations

The Novartis Group of Companies are committed to working with and providing reasonable accommodation to individuals with disabilities. If, because of a medical condition or disability, you need a reasonable accommodation for any part of the application process, or to perform the essential functions of a position, please send an e-mail to <u>us.reasonableaccommodations@novartis.com</u> or call +1(877)395-2339 and let us know the nature of your request and your contact information. Please include the job requisition number in your message.

部門 Operations 部門 Innovative Medicines



State Indiana

勤務地 Indianapolis

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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