

About the Role

Your Responsibilities:

Your responsibilities include, but not limited to:

Job Description

Manufacturing Systems Expert

Define User requirement specifications (Voice of customer)

- Design and create electronic batch files (EBR) with respect for quality, costs and deadlines
- MBR / BOM/ Recipe creation in production IT systems
- · Participate in the qualification and risk assessment processes
- · Propose and technically validate the choice of solutions proposed by the IT teams and Automation in collaboration with the Quality team
- · Participate to the deployment of MES across the manufacturing units, runs performance qualification and validation batches (commissioning of the line) with the manufacturing units
- Responsible for the manufacturing documentation update following the implementation of electronic batch files
 Provide training for MES users in partnership with the training team
- Ensure follow-up and processing of deviations and Change Control in compliance with deadlines and applicable regulations
 Real time shop floor troubleshooting with the implementation of appropriate immediate corrective actions
- Ensure the transfer of information to production teams following issues or modifications having a technical, quality or HSE impact Responsible for MES / MIS technical knowledge transfer to the shop floor Ensure the preparation of audits and inspections for related topics
- Collaborate with process experts in the context of deviations related to MES / MIS and continuous process improvement
- Participate to change management in close collaboration with change champion and P&O partner
 Ensure the application of local rules, procedures and policies

What you'll bring to the role:

- * Minimum 5 year experience in GMP manufacturing process support role · Proven experience in MES expert role
- * University degree in Science is required, Pharmacy or Chemical Engineering, Pharmaceutical Technology or equivalent job experience
- *Competencies Good scientific and technical (automation) understanding Deep process understanding
- *Quality and compliance skills
- *Team player with strong team spirit
- * Change management, adaptability, ability to work under pressure
- * Mastering of automation and computer skills

- *Good understanding of regulatory requirements across multiple health authorities
- *Mastering of manufacturing execution systems (MES, SAP, or other as applicable)
- * Good office software applications

Why Novartis?

Our purpose is to reimagine medicine to improve and extend people's lives and our vision is to become the most valued and trusted medicines company in the world. How can we achieve this? With our people. It is our associates that drive us each day to reach our ambitions. Be a part of this mission and join us!

Learn more here: https://www.novartis.com/about/strategy/people-and-culture

Commitment to Diversity and Inclusion:

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

Join our Novartis Network: If this role is not suitable to your experience or career goals but you wish to stay connected to hear more about Novartis and our career opportunities, join the Novartis Network here: https://talentnetwork.novartis.com/network

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

部门 Operations
Business Unit Innovative Medicines
地点 India
站点 Hyderabad (Office)
Company / Legal Entity IN10 (FCRS = IN010) Novartis Healthcare Private Limited
Functional Area Technical Operations
Job Type Full time
Employment Type Regular
Shift Work No
Apply to Job

Accessibility and accommodation

Novartis is 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 recruitment process, or in order to perform the essential functions of a position, please send an e-mail to diversityandincl.india@novartis.com and let us know the nature of your request and your contact information. Please include the job requisition number in your message.

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Job ID REQ-10021438

Manufacturing Systems Expert

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