

## Senior Sterility Assurance Expert (m/f/d) / Vi š ji strokovnjak za zagotavljanje sterilnosti (m/ ž /d)

Job ID  
REQ-10034729

3月 28, 2025

Slovenia

### 摘要

#LI-Hybrid

We are thrilled to announce the establishing of new clinical manufacturing facility in Slovenia, dedicated to accelerating the creation of innovative medicines for patients around the globe. Don ' t miss the chance to join our international team.

As Senior Sterility Assurance Expert Drug Supply you will be part of our Drug Product Clinical Manufacturing Team at our Technical Research and Development site in Menges, Slovenia and be primarily responsible for ensuring sterility assurance and contamination control at the Clinical Manufacturing Plant.

About the Role

## JOB DESCRIPTION:

### Key Responsibilities:

- Sterility assurance and contamination control within the Clinical Manufacturing Plant.
- Investigation of microbiological related deviations / OOX, conducting root cause analysis, and implementation of CAPAs and corresponding Risk Assessment.
- Contribution in the preparation and execution of validations (clean room validation, aseptic process validation), including process and environmental monitoring.
- Providing technical expertise during regulatory inspections and ensuring compliance with regulatory requirements.
- Developing and improving cleaning, sanitation, and environmental monitoring programs.
- Cross-Functional Collaboration to align activities with organizational goals.
- Mentoring and providing technical guidance to junior team members.
- Utilizing data analytics, machine learning, and artificial intelligence to optimize performance parameters.
- Proactive review and improvement of aseptic programs and contamination control strategies.

### Essential Requirements:

- Accountability: responsibility for assigned tasks and reliability.
- Decision Making: correct interpretation of analyses and evaluations and identifying appropriate measures to be taken.
- Ability to work in a team (constructive and reliable contribution in a group setting) and in a matrix environment. Influencing without authority.
- Results driven self-motivation and motivation of others to achieve outstanding results while ensuring adherence to ethical and legal principles, with a continuous drive for improvement.
- Customer focus as the highest priority.
- Quality focus: providing the highest quality products and services that meet the needs and requirements of internal and external customers.
- Significant experience in CMC development and/or production.
- 5 years of experience in Pharmaceutical Industry and 3 years of Microbiological Experience; thorough knowledge of cGMP requirements

### Desirable Requirements:

- 2 years of experience within Manufacturing QA

We offer permanent employment with 6 months of probation period. Submit your application with the CV in English language.

You ' ll receive:

Competitive salary, Annual bonus, Flexible working schedule, tailored to your needs, possibility to work from home, Pension scheme, Employee Recognition Scheme, Expanded program for the promotion of health in the field of physical, mental and social well-being (Wellbeing), employment at Top SI Employer, Unlimited learning and development opportunities.

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.

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  
Development

Business Unit  
Innovative Medicines

地点  
Slovenia

站点  
Mengeš

Company / Legal Entity

SI19 (FCRS = SI019) Novartis farmacevtska proizvodnja d.o.o.

Functional Area

Research & Development

Job Type

Full time

Employment Type

Regular

Shift Work

No

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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 [diversity.inclusionslo@novartis.com](mailto:diversity.inclusionslo@novartis.com) and let us know the nature of your request and your contact information. Please include the job requisition number in your message.

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



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