



QUALITY ASSURANCE SPECIALIST

Greater Zurich Area

CONTACT

MONIA@ELEMED.EU



THE COMPANY

Join the company's headquarters, based in a great location within the Greater Zurich area. This is a varied and responsible position in an innovative company that opens up long-term perspectives.

A fantastic new and exclusive role has come in, with the opportunity to develop your career and knowledge in Quality. You will work with experts willing to develop your skills.

As you learn and develop your skills, you will be provided with the perfect platform to develop a strategy for the department according to your own ideas and expertise; thus allowing you to leave a legacy within a company at their exciting point of growth.

The company is developing novel devices on a global scale.

There are strong opportunities in the future for interesting evolution.

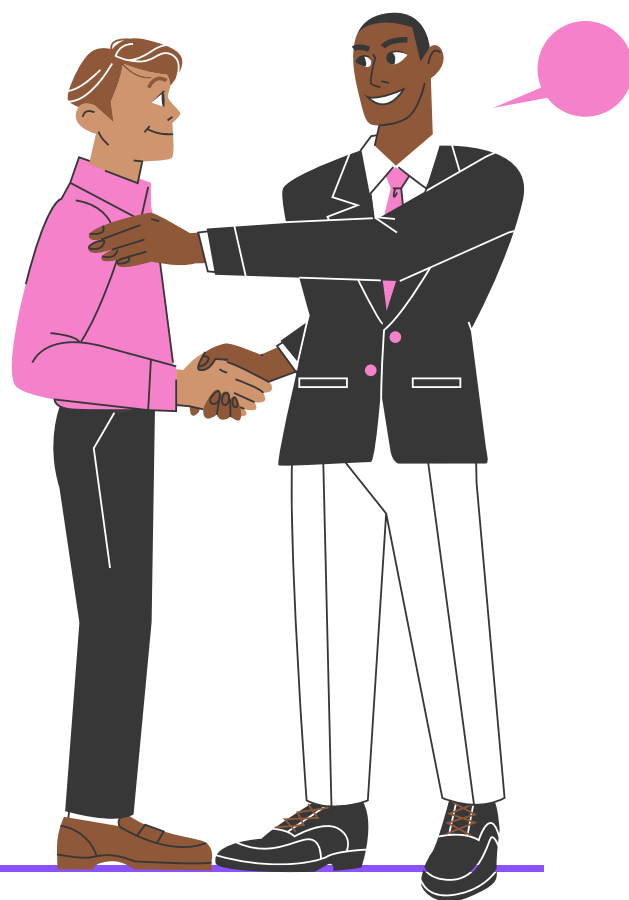


THE OPPORTUNITY

In this position you will be responsible for ensuring quality in terms of function, safety and compliance with regulatory requirements in their development and manufacturing phase.

YOUR RESPONSIBILITIES (NON EXHAUSTIVE LIST):

- Implementation and support of validation activities internally and with external partners (measuring equipment, software, production machines and processes)
- Monitoring of the production processes
- Quality Assurance and batch releases
- Supervision and implementation of internal and external audits
- Contribution towards the creation of an efficient quality management system
- Cooperation with risk management and post market surveillance team
- CAPA evaluation and processing
- Supporting R&D department with the interpretation of regulatory requirements and, if necessary during product development
- Responsible for regulatory content of product - related information
- Supervision and coordination of internal and external stakeholders in regards to complaint handling
- Ensuring all specified quality and documentation requirements



YOUR QUALIFICATIONS:

- Scientific or technical studies or equivalent
- Experience in quality management in the medical device sector
- Business fluent in German and English





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**ARE YOU
INTERESTED
IN THIS
EXCITING
ROLE?**

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