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ST. GALLEN REGION

Head Quality Management System

The opportunity

A unique opportunity to lead a team of managers and engineers, while also remaining in the exciting technical aspects of technical; from working with the QMS, CAPAs and dealing with conformity issues.

A great role to report directly into the Site Manager and work alongside them to develop key ideas and strategy for the growing quality department.

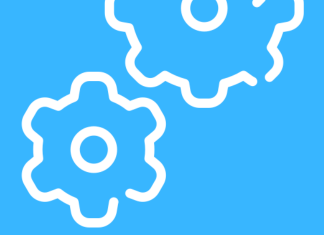
Be the next manager of a growing and global medical device company who specialise in innovative medical devices.

The role

Because this role is located in the company HQ, where the senior management team sit, this role will have significant interaction and exposure to the company's Leadership Team. We are looking for dynamic, seasoned Quality professionals with experience in operations and stakeholder management who are able to influence both up and down reporting lines.

- Lead the quality organization in Switzerland in line with the Company's Vision and Values of Ambition, High standards and Trust.
- Drive compliance of both products and processes to applicable standards and regulations globally (e.g. QSR, ISO 13485)
- Lead a team across quality assurance, validation and compliance.
- Have a commercially minded approach to quality to support the business achieve its goals and objectives
- Drive projects to improve the reliability and quality of products by establishing effective control mechanism in operations and supplier managementDrive accountability towards effectiveness of the QMS







Expectations

We are looking for seasoned professionals with a proven background in medical device quality assurance and leadership:

• 5+ years experience working in a quality assurance role within the medical device industry and mastery of the relevant standards: ISO 14971, ISO 13485, EU regulations, quality and regulatory compliance processes.

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- Expertise within MDR implementation.
- Fluency in English AND German
- Proven experience with manufacturing quality processes, operational quality,
 Quality Systems Management, Audits, Quality Engineering

Interested to explore this further?

Please send your CV to tamanna@elemed.eu to arrange a confidential career discussion.



Only applicants who meet the requirements above will be considered for the role. Unless otherwise stated we are not able to consider applicants without EU work authorization.

Elemed is an executive search firm, specialized in finding and representing exceptional talent in medtech. To find out more about our Candidate Services click here:https://www.elemed.eu/candidates/