


Design Quality Engineer New Product Development

Solothurn, Switzerland



**Are you looking for a company that truly has a
care for quality?**

**Do you want to work in one of the top employers
in the industry?**

**Are you excited by new product development
projects?**

We may have a great opportunity for you!






The Company

We are recruiting a Senior Design Quality Engineer focused on Design and Maintenance for New Product Development Projects of medical implants and instruments.

This is the chance to make your own mark in a brand new role, in a growing team in one of the best performing divisions in the company! Constantly investing in R&D and its employees, you will have the opportunity to join in social events, team building activities and much more!

This company has been voted as one of the top employers in Medtech over and over again and is known for creating high performing and fun teams.



In this new and exciting Senior Design Quality Engineer role you will:

- Ensure compliance throughout the design and development process by reviewing risk management files, design transfer protocols and development documents
- Work on exciting development projects with partners both internally and externally on cross company collaboration projects
- Represent quality interests in product development projects of implantable devices
- Collaborate in multi functional teams alongside R&D, Quality, Regulatory Affairs, Process Management and Biomechanics
- Supporting global regulatory submissions
- Be responsible for risk management moderation in development projects
- Review and validate reports and take CAPA where needed
- Continuously improve quality processes in compliance with relevant legal guidelines

Contact us if you have...

- A technical/engineering background, with a good eye for details
- 3+ years experience in a quality function in the medtech industry
- Experience with ISO 13485 and MDD/ EU MDR
- Hands-on experience in risk management and design controls/quality processes



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