

Regional Quality Assurance Specialist

Eindhoven, Netherlands

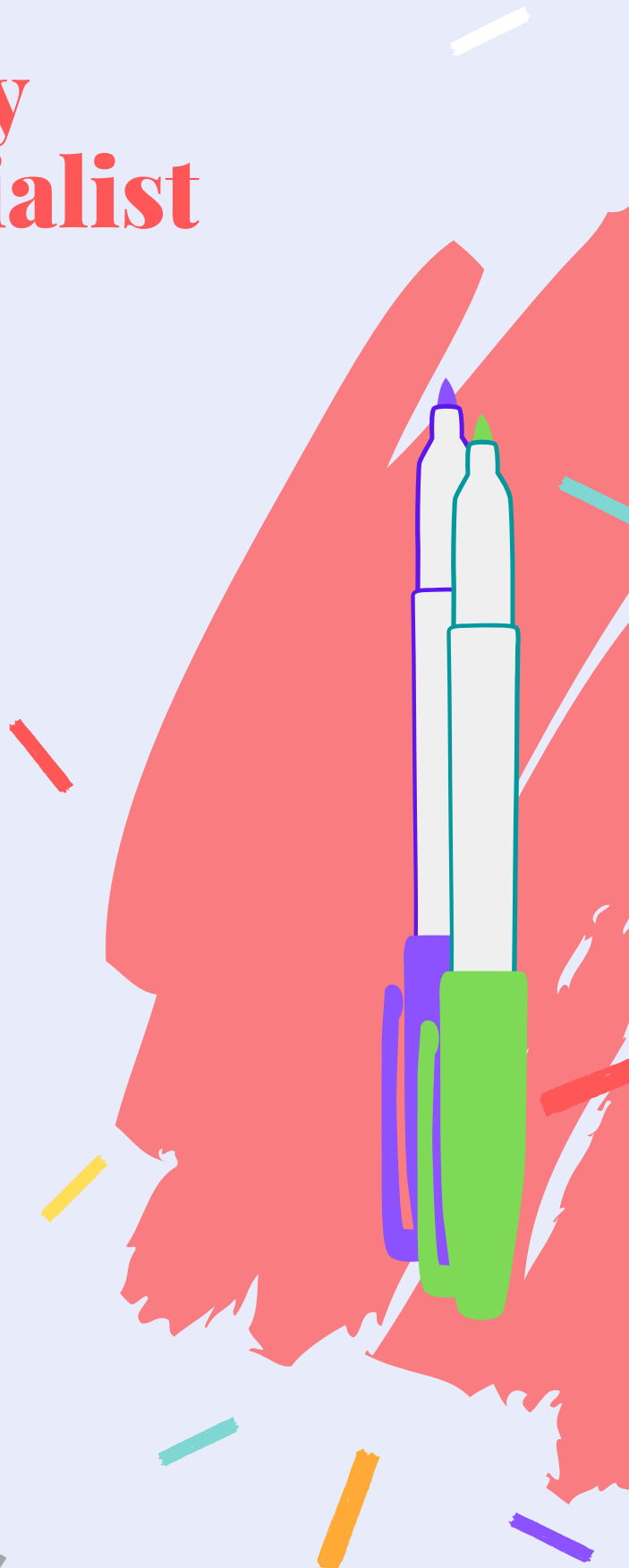
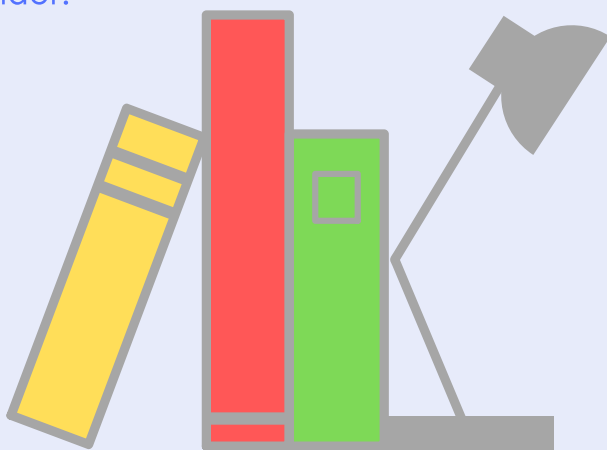
Come and join a global leader in healthcare with locations across the world and around 8,000 employees. The devices (manufactured and sold globally) are groundbreaking AND life changing. Thanks to a full pipeline of R&D and new products, this company is really leading the way and innovating within their space.

At this company you'll enjoy a flat hierarchy (no top down management here!) where communication is open and you are valued not just as an employee but as a person. This company has very high ethics and a unique culture where the work-life balance is important!

The Opportunity

You will have the opportunity to be a part of the Regional Quality Assurance team for EMEA.

As Regional Quality Assurance you will be based at the European distribution warehouse for global markets. Not only will you support the warehouse operation, you will also interface with the core company's operation and 3rd party logistics provider.





Your responsibilities:

- Drive improvement efforts and ensure compliance of the Quality Management Systems
- Create a balanced perspective for improvements and changes by interfacing with the 3rd party logistics provider
- Support the Regional Quality Assurance operations locally and globally by designing and developing processes
- Be the representative for Quality for reboxing, boxing and labelling operations on site under the internal QMS
- Host and lead internal audits as well as support external audits
- Travel up to 10% in EU and internationally

Your qualifications:

- 3+ years experience in Quality Assurance or other Quality function for the medical technology industry
- Knowledge and experience with ISO 13485
- Fluent English speaking, reading and writing

**We'd love to
hear from you!**

**Please send your CV to
hello@elemed.eu for 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.

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