

THE COMPANY

Are you an expert in process verification and validation for medical devices? Are you tired of working in large, slow-moving corporations but worried about joining an unstable start-up? Why not join a well-established, commercial stage company that continues to grow each year?

This company is designing, developing and producing an innovative Class III active-implantable medical device that is already CE marked and changing the lives of patients. If you enjoy working in a fast-paced and highly dynamic environment, this could be the place for you!

To support their continuous growth, this company is expanding their QA/RA team and creating brand new roles such as this verification and validation engineer.



THE OPPORTUNITY

This is an exciting time to be joining the company as they grow in all areas including people, product indications and markets. It is an opportunity to step into a brand new position and gain more skills and knowledge working on a complex portfolio with 1300 components. You will work collaboratively across all departments of the business and dive deep into your V&V activities.

WANT TO FIND OUT MORE ABOUT THE ROLE? GO TO THE NEXT PAGE!



AS VERIFICATION AND VALIDATION ENGINEER YOU WILL:

- Execute verification and validation activities including validation of software, processes, products, equipment etc. and all other associated revalidation activities
- Generate, review, edit and approve qualification and validation / revalidation protocols, process specifications, FMEAs, test plans, summary reports and other associated documents to allow conformance to regulations and standards
- Analyse test results using statistical techniques and determine acceptability
- Investigate and troubleshoot problems identified during and associated to verification and validation activities
- Determine solutions or recommendations for changes and/or improvements
- Support the validation of software used within the quality management system (e.g. software controlled test equipment used during in-process inspection)
- Coordinate with cross-functional teams to plan and facilitate Qualification/Validation activities within the organization

AS VERIFICATION AND VALIDATION ENGINEER YOU WILL HAVE:

- A minimum of 3 years demonstrated Verification and Validation experience
- Experience working in the medical device industry
- Fluent reading, writing and speaking in English
- A hands-on approach



Interested to explore this further?

Please send your CV to Elena at <u>kristina@elemed.eu</u> to arrange a confidential career discussion.

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