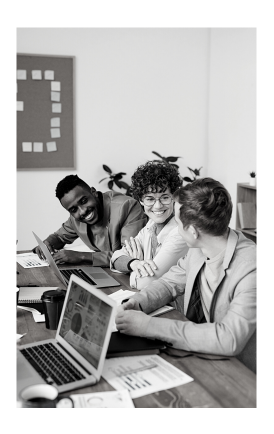


The Company



Are you looking to join a company with open communication, positivity and an excellent social environment? Look no further! This is an outstanding opportunity to join a world leading medtech company that has been recognised to be one of the top employers in medical devices. This is a brand new position focussed on implementing the new MDR for a wide range of devices (class I-III), focusing on trauma and extremities

Your Mission

As Associate Regulatory Affairs: You will be able to develop and apply all your knowledge and understanding of the RA frameworks, legislative requirements, processes and procedures in the EMEA distribution organisation.



Your Responsibilities

- Executes RA activities in line with defined procedures and processes for the UK and BeNeNord region.
- Provides support to EMEA / Country RAQA teams as appropriate.
- Work with the Benelux leadership teams and acts as a business partner for the local region
- Identifies the need for new regulatory procedures
- Identifies opportunities for continuous improvement and supports those activities across RAOA.
- Collects, organises and maintains files on local, regional, and global RAQA intelligence.
- Lead teams to support regulatory processes for market access of products



Essential requirement



- 1.3rd level Degree in Life Sciences. RAC certified preferred
- 2. Minimum 3 years experience in regulatory affairs for medical devices
- 3. Fluent in Dutch and English

