R&D Medical Device Specialist

Location: IL- Caesarea

A Day in the Life

As an R&D Medical device specialist, you will be leading the Materials of Concern compliance within the site and support the R&D by being an SME (subject matter expert) in Biocompatibility and Surgical instrument reprocessing, while working with some of the best consultants and knowledge centers in the field in the world. The role involves the application of critical thinking, technical skills, teamwork, and creative problem solving to support the design and delivery of high-quality innovative medical devices intended to improve patient outcomes and the quality of life for millions of people.

As part of your day to day you will:

Be a part of a mechanical design group within the R&D that is responsible for developing surgical platforms.

- Lead the Materials of Concern compliance including a deep understanding of relevant internal procedures and different laws and regulations governing MoC (for example: RoHS, REACH).
- Work with different functions in order to demonstrate compliance with laws and regulations governing Materials of Concern.
- Provide guidance, technical evaluations, and training regarding MoC laws, regulations, and internal procedures.
- Be the site SME (subject matter expert) for Surgical instruments Biocompatibility, Cleaning, and Sterilization requirements and testing, working with knowledge experts in the various fields in Israel and abroad.
- Define and lead the implementation of procedures required as part of the R&D development process, according to company guidelines.
- Write technical assessments and reports.
- Preform and support evaluations for the impact of changes in laws and regulations.
- Participate in internal and external audits as needed.

Must Haves

- Bachelor’s Degree with at least 2 years of relevant experience from a Medical device company or a highly regulated industry.
- Strong written and verbal communication skills in Hebrew and English.
- Ability to define and implement complex processes and guidelines within an R&D matrix organization.
- Highly organized.
- Strong initiative.
- Problem-solving skills.
• Team player with the ability to work on your own.
• Able to effectively work with cross-functional teams in a multinational matrix organization.

**Nice to Haves**

• Advanced knowledge of MoC laws and regulations (for example: RoHS and REACH).
• Knowledge in medical devices Biocompatibility, Cleaning and Sterilization processes, requirements and regulations.

**About Medtronic**

Together, we can change healthcare worldwide. At Medtronic, we push the limits of what technology, therapies and services can do to help alleviate pain, restore health and extend life. We challenge ourselves and each other to make tomorrow better than yesterday. It is what makes this an exciting and rewarding place to be.

We want to accelerate and advance our ability to create meaningful innovations - but we will only succeed with the right people on our team. Let’s work together to address universal healthcare needs and improve patients’ lives. Help us shape the future.

**Physical Job Requirements**

The physical demands described within the Responsibilities section of this job description are representative of those that must be met by an employee to successfully perform the essential functions of this job. Reasonable accommodations may be made to enable individuals with disabilities to perform the essential functions. For Office Roles: While performing the duties of this job, the employee is regularly required to be independently mobile. The employee is also required to interact with a computer, and communicate with peers and co-workers. Contact your manager or local HR to understand the Work Conditions and Physical requirements that may be specific to each role. (ADA-United States of America)

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