Job Posting

Regulatory Affairs Associate

OSSIO is seeking a talented and dedicated individual who is interested in applying his/her scientific knowledge in the technical field of Regulatory Affairs. This diverse and challenging role will evolve as his/her technical knowledge and experience increases. This individual will be responsible for assisting the Regulatory Affairs department by reviewing and updating documentation, writing/updating Technical Files and Essential Requirements, compiling data, and providing information necessary to register OSSIO’s products in the global market through collaboration with all parties involved.

OSSIO Ltd. is an orthopedic fixation company committed to transforming the orthopedic experience. The company is focused on developing and commercializing bio-integrative fixation implants and is based in the industrial park in Caesarea.

The Regulatory Affairs Associate will be required to gain intimate knowledge of the Company's product line, including the general scientific technology used to manufacture and test the product, how the product is used, and how the product is marketed. The Associate should possess a driving need to understand the how or why, should not be afraid to ask questions, and should seek and provide candid feedback. In addition, the Associate should be able to form beneficial professional relationships with individuals at OSSIO and with regulatory agencies/companies.

Requirements

• B.Sc. Degree in Life Science or related field. (M.Sc. preferred)
• Excellent written and verbal communication skills in the English language.
• Strong organizational skills with attention to detail and accuracy as well as good analytical skills.
• Ability to conduct thorough research for global regulatory submissions.
• Ability to work well under pressure.
• Ability to multi-task to meet strict deadlines.
• Computer literate with proficiency of Microsoft Office package including Word, Outlook, Excel, and Adobe PDF.
• Mental flexibility to solve complex problems.
• Ability to work independently and as part of a team.
• Ability to maintain high ethical standards of integrity and quality.
• Knowledgeable about current Good Manufacturing Practices (cGMPs), Good Clinical Practices (GCPs), and Good Laboratory Practices (GLPs) preferred.
• Familiar with regulations/guidelines governing development of medical devices preferred.
• Enthusiasm and willingness to learn are more important than work experience. More experienced candidates may be considered.

We look forward to you joining our team!

Qualified candidates can submit a cover letter along with their resume or CV in English to taly@ossio.io.