KiOmed Pharma hires:
“Design & Development Project Leader”
Implantable Medical Devices - innovative biomaterial products.
Available immediately / permanent contract / Herstal (Belgium).

KiOmed Pharma (http://kiomedpharma.com) provides safer and efficient chitosan-based therapeutic implants to patients who suffer from high impact pathologies.

We are specialized in the development, manufacturing and commercialization of innovative pipeline of implantable medical devices in compliance with requirements of European regulations and ISO 13485 quality standards. Our products aim at addressing unmet medical needs in regenerative medicine, mainly focused on joint health and dermatology markets. A clear example is our revolutionary clinical-stage injectable chitosan gel, which has been designed to combat the progression of osteoarthritis.

To grasp a new growth opportunity, we are actively seeking a “Design & Development Project Leader”:

REQUIRED QUALIFICATIONS:

- Scientific background, minimum master in chemistry, life sciences or engineer, preferentially a PhD.
- At least 5-year experience in the development of implantable devices.
- Ability to conduct all stages of the value chain from design and development, pre-clinical activities till process transfer / industrialization.
- Knowledge of Design and Development activities required for CE marking of implantable devices.
- Knowledge of the European regulations applied to medical devices for CE marking.
- Background in biopolymers is an asset.
- Effective project management and leadership skills.
- Strong analytical and problem-solving skills, result- and customer-oriented.
- Ability to work in a dynamic, fast-moving environment.
- Rigorous, you dispose of a good sense of organization.
- Excellent communication and reporting skills, French and English both written and oral.
- Computer skills (Microsoft Office suite) and knowledge of project management tools.
MISSIONS:

The D&D Project Leader will take a leading role in Design & Development (D&D) programs in relation to the strategic development roadmap of the company. He/she will ensure the effective management and leadership of the program. He/she will provide scientific advices with respect to design controls required for CE marking of implantable devices and will consequently provide a clear definition of the scope of work of the program. He/she will establish the project plan into appropriate work packages and will contribute actively to the progress of the project plan. He/she will ensure that the right product is developed and delivered to meet Company’s strategy and quality standards. Strongly result- and customer-oriented, he/she will lead the program with a focus on meeting established projects milestones in compliance with the budget and timelines. He/she will report on progress and issues to all stakeholders of the project and will provide thorough control of subcontracted activities.

He/she will report to the CEO of the company and will interact transversally with the business development and compliance teams.

TASKS AND RESPONSIBILITIES:

1. Lead the development programs in relation to the strategic development roadmap of the company,
2. Develop a project plan and identify project timelines and milestones,
3. Translate the project plan into well-defined task orders and communicate effectively to both internal (project team, D&D lab, production) and external (subcontractors, customer) partners involved on the project,
4. Manage the execution of the project with a focus on meeting established milestones in compliance with the budget and timelines,
5. Support the management in decision making in line with product development,
6. Prepare periodic management and/or customer reports and presentations,
7. Organize and archive the documentation in compliance with Quality standards for Design & Development,
8. Contribute to the preparation of documents for regulatory purposes,

If you are interested in this challenge, please send your full application (CV and motivation letter) to jobs@kiomedpharma.com , until June 30th, 2016.
Your application will be processed with full confidentiality.