



DIRECTOR REGULATORY AFFAIRS, QUALITY & CLINICAL

Lyon, France

DO YOU HAVE THE FOLLOWING EXPERIENCE?

Essential requirements:

- 10+ years of experience from the perspective of a legal manufacturer:
 - a) Regulatory affairs with class IIb medical devices
- OR
- b) Regulatory & Clinical experience
- Leadership experience of managing a team
- Fluent English and French

Desirable (but not a requirement).

- Quality
- Auditing experience / Lead auditor certification

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THE COMPANY

Tired of working for the same big slow moving corporation where the key decisions are made outside of Europe? Are you looking for a role that will allow you significantly more autonomy in taking QA/RA/CL decisions, where you will sit at the table of the company's leadership team?

Located near the beautiful cultural city of Lyon, is Elemed's newest client; a small medical device manufacturer. Small in terms of people, but not small in terms of presence. This is not an opportunity in a mono-product, mono-franco environment. This company has a significant product portfolio and designs, develops, manufactures and sells their products for the global market: EMEA, N.AMERICA, LATAM & APAC.

In this role you'll be at the heart of the company's operations as legal manufacturer, overseeing a broad range of activities covering Regulatory, Quality and Clinical globally, as well as leading a young and dynamic team.

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Responsibility

THE ROLE

In this role as Director Regulatory, Quality and Clinical; your responsibilities will centre on the following goals:

Leadership: Manage the QA/RA/CL team, organising their work and priorities, supporting them to achieve their objectives and ensuring the continuous skills-development of the group.

CL/RA: Oversee regulatory and clinical compliance of the products manufactured by the company, defining clinical and regulatory strategy and clinical studies are executed in compliance with regulatory requirements.

Quality: Be the PRRC, and the quality management representative. You'll be responsible for ensuring the quality system is compliant and implemented correctly, as well as making sure that products comply with qualified and approved standards/processes.

Want to learn more? Read on for a detailed breakdown of the role's responsibilities.

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LEADERSHIP

- Define your team's goals and objectives, and ensure they are achieved by driving, supporting, mentoring and motivating them. Define and implement key KPIs and ways to measure your team's performance
- Manage the team and allocated resources and budget, in order to ensure the leadership's vision and objectives are achieved
- Review and approve the team's solutions and proposals, as well as be responsible for managing the calendar and pipeline of projects; providing progress reports and alerting the leadership team in case of problems or delays
- Maintaining an open channel of communication by sharing any changes or advancements ongoing within the company with your team
- Unlock talent and potential by ensuring each team member is performing at their best ability, providing them support and guidance to help them advance in the development of their skills and competences.

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REGULATORY

REGULATORY AFFAIRS

- In this role you'll also assume responsibility as PRRC; named person responsible for regulatory compliance
- You'll oversee regulatory monitoring and ensure compliance with regulatory requirements specific to Medical devices (21CFR/820, 93/42/EEC) including the continual work on the transition to the new MDR (2017/745)
- Drafting and updating of CE technical files in accordance with MDD/MDR as well as providing input to R&D teams on new product development projects/changes to existing products.
- Overseeing international registrations through connections with local partners, subsidiaries and distributors and ensure compliance with local country regulations
- Leading the relationship with competent authorities and Notified Body as the main point of contact
- Carrying out Regulatory Intelligence: to make sure you are updated on the changing regulatory requirements around the world; in particular in markets of interest.

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QUALITY ASSURANCE

- Be the Company's quality representative in accordance with ISO 13485
- Driving the Quality agenda and building an internal culture of quality
- Advise the leadership team on Company quality policy: supporting with the definition of key objectives and the relevant strategies on how to achieve them
- Overall management of the QMS, ensuring it is implemented correctly, and ensuring compliance with the standards and regulations applicable to Medical Devices such as ISO 13485, and 21 CFR/820, MDD 93/42/EEC, as well as the new medical device regulation 2017/745
- Monitor and look for opportunities to continually improve the Quality Management System
- Oversee the management of operational quality activities
- Audit: Conducting Internal and supplier audits, Being responsible for third-party audit as the main point of contact and for follow-up

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CLINICAL

- Lead and define the clinical strategy for the company in relation to clinical evidence and clinical investigations
- Oversee the carrying out of clinical studies through all the phases: from setup to close-out, with CRO or internal resources
- Sign the declaration linked to annex XV, chapter II, section 4.1 of the MDR 2017/745

DESIRED EXPERIENCE

Essential requirements:

- 10+ years of experience from the perspective of a legal manufacturer:
 - a) Regulatory affairs with class IIb medical devices
- OR
- b) Regulatory & Clinical experience with minimum class IIb devices
 - Leadership experience of managing a team
 - Fluent English and French

Desirable (but not a requirement).

- Quality
- Auditing experience / Lead auditor certification

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Interested to explore this further?

Please send your CV to Elena at elena@elemed.eu to arrange a confidential career discussion.

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