



BlueReg Group is a consultancy company specialized in Development, Regulatory Affairs, quality management and Pharmacovigilance for the Pharmaceutical Industry.

Based in Paris, Sophia Antipolis and London, BlueReg Group has over 100 clients ranging from large multinationals to small start-up companies, from innovators to generic companies.

We are recruiting:

Associate Director, CMC Regulatory Affairs (CDI)

Your key duties and responsibilities:

- Is a source of CMC regulatory expertise in the development, registration and post-licensing activities of pharmaceutical products (CMC advice, pharmaceutical development strategy, writing of CMC documents)
- Project management activities
- People management / Review of work of junior personnel and BlueReg partners
- Ensures high quality and on time delivery of consulting services to clients.
- Contributes to the effective functioning and to the business growth of BlueReg EU

Your profile :

- Pharmacist, Life Sciences Graduate, ideally with post-graduate qualification.
- Sound knowledge of pharmaceutical regulations and guidelines.
- Significant regulatory affairs experience including a successful track record in the registration of medicinal products
- A least 15 year experience in the pharmaceutical industry, broadly based Regulatory Affairs experience and CMC, including a successful track record in the registration and maintenance of pharmaceutical products within Europe .

You have excellent written and verbal communication skills, good organisational and analytical skills. You are fluent in written and spoken English.

This position is based in Sophia Antipolis (South of France).

If you are interested in, please send a copy of your CV and a cover letter to contact@blue-reg.com.