



## **Senior Clinical Research Associate / Clinical Trial Manager**

Do you want to be part of a dynamic team that drives clinical activities in an inspiring environment? Would you enjoy to work with different projects and with the ability to influence your job? Larix Clinical Operations is now seeking an experienced Clinical Research Associate / Clinical Trial Manager to join our Lund office.

**Larix A/S is a Nordic CRO** – we offer full-service solutions within the pharmaceutical and medical device areas. Our headquarters are located near Copenhagen in the middle of the Medicon Valley region, and we have strong ties to the thriving pharmaceutical and biotech activities in this region.

Clinical Operations consists of experienced and enthusiastic Clinical Trial Administrators, Clinical Research Associates and Clinical Trial Managers responsible for the planning and conduct of clinical activities in the Nordic countries.

**At Larix, we maintain a friendly atmosphere** – we think having fun while working is important. We are a relatively small company with approximately 70 employees, and we have all the benefits of being able to collaborate across functions, follow clinical study processes from start to finish, and learn from each other. We offer an exciting job in a dynamic company, which is developing extensively. You will have a great influence on the development and on future tasks.

**At Larix, you'll be part of our Lund office** and report to our Director Clinical Operations in Denmark. We are based at Ideon Gateway in Lund, an exciting and inspiring office environment. You will either work from our offices in Lund, but also be prepared to work at a client's office in the Lund area and in our Herlev office from time to time.

Your main tasks will include:

- Perform and handle all aspects of trial & site management of sponsor projects
- Perform monitoring (from Pre-study, Initiation, Monitoring and Close-out visits)
- Submission to regulatory authorities and ethic committees
- Perform and handle trial management of small or larger studies
- Prepare or contribute to the update of SOPs and procedures within Clinical Operations

**As an experienced Clinical Research Associate /Clinical Trial Manager** – we will get to know you as a proactive and competent colleague who approaches projects with a team players' spirit. At the same time, you are very good at working independently and are able to plan, structure and drive your own tasks. You are an open-minded, co-operative and service-minded person. Moreover:

- You have a relevant background from life science
- You have 5+ years of experience with on-site monitoring (site feasibility, selection, initiation, monitoring and closure) of clinical trials according to ICH GCP
- You have experience with submissions of clinical trials to Ethics Committees and Regulatory Authorities
- You have extensive knowledge of relevant national and European guidelines and regulations for clinical trials
- You have the flexibility to travel
- Experience with medical device studies is considered an asset
- You have experience from the CRO industry, the pharmaceutical industry and/or biotech companies

**Communication ♦ Proactivity ♦ Quality on time**



- You are a dedicated team player with a high-quality mind-set, meet your deadlines and know how to prioritise between different tasks in a dynamic environment which requires a high degree of flexibility
- You have good communication skills – and of course, you speak and write English and Swedish effortlessly

**At Larix, we look forward to welcoming you on board** - we offer an attractive salary package including pension scheme, life and medical cover, and the opportunity to work from home from time to time.

You are welcome to contact Iris Koenig, Director Clinical Operations at [iko@larixcro.com](mailto:iko@larixcro.com) or call for further information: +45 81 77 37 58. You can also visit our website at [www.larixcro.com](http://www.larixcro.com)

To apply for this position, please forward your application and CV to [info@larixcro.com](mailto:info@larixcro.com) no later than 28 February 2018. Applications will be handled in the order they arrive.