

Symeres is the leading mid-sized transatlantic contract research organization for drug discovery and development needs. With over 700 highly educated scientists and professionals in six locations in Europe and two in the USA, we offer best-in-class solutions for drug discovery and drug development, from small- to medium-sized molecule hits. We are large enough to matter, and small enough to care. Our services span across early stage hit finding all the way to the delivery of your early clinical drug substance API. Making molecules matter. Together.

Due to rapid international growth and increasing demand for our services at our location in Weert, we are currently looking for a skilled and enthusiastic:

Analyst **(M/F) Full-time**

The Quality Control Department is responsible for analytical method validation, the testing of raw materials, in-process controls, intermediates, active pharmaceutical ingredients (APIs), and stability testing. The Quality Control Department is equipped with state-of-the-art equipment for analytical techniques such as GC (MS), (u)HPLC (MS-MS), ICP (MS), NMR, IC, wet chemistry, and FT-IR.

Key job responsibilities:

- Carry out analytical testing intended for the release of (critical and project) raw materials, intermediates, and final products, based on approved methods of analysis.
- Compare results against established specifications.
- Report and examine aberrant and OOS and OOT results. Where possible, suggest solutions and/or improvements.
- Follow analytical protocols for stability studies, retesting of reference standards, and validation of analytical methods.
- Write analytical reports on stability studies and validation of methods.
- Calculate, interpret, and report on the obtained test results.
- Write/review procedures and working instructions.
- Collaborate closely with project teams and manufacturing to discuss timelines, flag and discuss deviations from procedures, and share results.

Qualifications

- Minimum MBO+/HBO level.
- Knowledge of analytical chemistry acquired through education and/or working experience in a pharmaceutical environment.
- Broad (practical) analytical knowledge of analytical techniques such as (U)HPLC (MS), GC (MS), titration, FT-IR, NMR, and ICP-MS.
- Knowledge of Good Manufacturing Practices and EHS directives.
- Strong communication skills, both verbally and in writing. Fluent in English and Dutch.
- A good working and quality attitude, accurate, with an eye for detail.
- Team player.

Job offer

An exciting position in a dynamic fast-growing organization with an attractive remuneration package and opportunities for learning and development.

Application

Please send an email with your CV and motivation letter to our corporate recruiter, Bhumika Chawla, via: Bhumika.Chawla@Symeres.com. For questions about the vacancy, please contact Chantal Rademaekers-Litjens (Director – Quality) by phone: +31 6 30 99 29 63.

More information about the Symeres organization can be found on our website: www.symeres.com.

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