

Susanne Thyssen Rasmussen

B.Sc. Clinical Chemistry

Senior Consultant Management and Compliance

ID1030

Direct (+45) 53 64 09 90 Susanne.Rasmussen@medicologic.com

medicologic°

Medical Device Development

Medicologic A/S Arne Jacobsens Alle 17 Ørestad City DK-2300 Copenhagen Denmark (+45) 48 24 51 13 contact@medicologic.dk www.medicologic.dk

Competences

- Project Management
- Quality Management
- Risk Management
- Medical Device
- People leadership and management
- Process mapping
- cGMP
- · Performance management
- Change Management

Standards

- MDR 2017/645
- MDD 93/42/EC
- ISO13485
- ISO14971
- ISO11607
- FDA regulation [CFR part 820]
- cGMP

Key results

Project Management as responsible person for implementation of large projects. E.g., construction and implementation of a pharmaceutical production site. Or as member of top management responsible for the PMO in Biogen

Change Management leading people towards major changes with success e.g., Implementation of LEAN leadership in several sites in Novo Nordisk

QMS and Technical File writing and updating QMS for LEO Pharma to fit a changed organization

Significant leadership skills leading teams and departments over the years in both quality control, aseptic production, and in-vitro production

Personal Characteristics

She is a competent leader, who takes responsibility for the people and the tasks from bottom to top.

She is a go-doer and has a no-nonsense attitude always finding a positive and constructive solution to a problem.

Susanne is an experienced consultant and is highly appreciated by her clients.

Privately She enjoys reading books, watching films and theatre, listening to concerts and records, and off course being a grandmother having fun with both dolls and LEGO.

References

Medicologic A/S (present)

Senior consultant Management and Compliance

Phillips Medisize

Project Manager for development and commercializing a medical device towards the US market. The project team is 25-50 people all involved in different roles during the project. My responsibility is to keep the project on track financially, and according to the overall milestones. Project management supporting engineering and product development in combination with labelling and documentation of a clinical trial variant for several markets.

Glostrup Apotek Magistrel Production

Overall Project Manager and responsible for coordination with the building project to ensure proper implementation of the production processes and installation of large machinery.

Price Invena

QA manager with overall responsibility for production, release of products and QMS. 13485:2016 certification has been received and MDR compliance in scope.

LEO Pharma:

Project Management in regulatory Affairs, Intermediate QA person writing and implementing an updated QMS. Senior Manager RA, Planning & Processes, LEO Pharma

Biogen

Associate Director Manufacturing, Biogen Site Leadership Team member, responsible for PMO

Novo Nordisk

Associate Manager and cLEAN® Partner, Novo Nordisk Senior cLEAN® consultant, cLEAN®Office, Novo Nordisk

ALK-Abello

Director of Production, IVBDD, Manager of Aseptic Production Manager Quality Control, Director Quality Control

Research assistant to Professor Klaus Øllgaard, RH Research assistant to Professor Sixtus Thorsen, KKHH Bio-analyst, Clinical Chemistry, Hvidovre Hospital