

Bringing your project to success while mastering the challenges of MDR (medical device & software)

Goal:

The goal of this workshop is to give an overview of the regulatory landscape for medical devices in Europe (MDR), the product development process, as well as the requirements regarding technical documentation. Within the MDR the workshop will focus on the development of medical devices and software.

Contents:

- The regulatory landscape for medical devices and software in Europe (MDR)
- The product development process and its milestones
- Technical documentation according to MDR Experience from previous projects
- Q & A Session

Who should attend:

Start-up Founder, Start-Up Team seeking advice on regulatory affairs (medical devices and medical software)

Time:

This two hours workshop will be held on

Thursday September 17th, 2020 at 14:00 - 16:00

Location

Online Webinar

Price

As we like to accelerate and support innovation in Switzerland this workshop is **free of charge for start-ups**. The fee for all other companies is 140 CHF.

Register

Please register by sending an email to: **workshops@effectummedical.com**Please note that space is limited, priority is given to start-ups.

Speakers:



Nila-Pia Rähle

COO and Co-Founder of Effectum Medical, a legal manufacturer for medical devices offering an outsourced quality management system (QMS) solution. Nila has been working in medical devices for nearly 20 years, for both global players and start-ups. She has broad experience along the whole value chain with a key focus on project management, regulatory affairs and quality management.