

SPARK EUROPE WEBINAR SERIES 2023

Wednesdays
at 4 pm CET

SPARK Europe
Education

Regulatory landscape for medical devices - how to approve your solution quickly in Europe or should you focus on the US market Dr. Jani Virtanen

8 November 2023 | 4 – 5 pm (CET) | Online Webinar



You have a solution (e.g. surgical tool, vascular implant, equipment or material for in vitro diagnostics, software etc.) which may become a medical device?

In this online **SPARK Europe Webinar Series**, Dr. Jani Virtanen is giving insightful information on getting regulatory approval needed to launch your medical device to the market. **You may implement this knowledge in constructing your research plan and planning budget for your next grant application.**

After this talk, you will:

- understand the right order of main steps to get CE approval needed for your medical device to launch on the EU market
- get an overview on timelines and the budget needed for the approval process
- know the requirements for selecting a suitable Notified Body
- understand when to choose US market as your first regulatory approval territory

Dr. Jani Virtanen has been working in several international medical device companies (e.g. Nexstim, Sooma Oy, GE Healthcare) for 17 years and has brought more than 15 medical devices to the market. He holds a PhD in Medical Engineering from University of Oulu.

Online via Zoom | Please register [here!](#)

Registration to the webinar is required in advance. Please register no later than **6 November 2023**. Please note that you will receive the Zoom link and access code the evening before the lecture and that access to last minute registrations, cannot be guaranteed.

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In unforeseen cases, the organizers may change and update topics and speakers.

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