

# LEGACY DEVICES

## HOW TO PREPARE FOR THE MDD/MDR TRANSITION BY USING POST-MARKET DATA IN THE TRANSITION PERIOD



### MAIN TOPICS

The course will outline how to apply the requirements of MDR **Art. 120** to devices that are already MDD-certified, seeking to understand how regulatory strategies can be optimized to produce adequate clinical evidence.

- The Guideline MDCG 2021-25 "Regulation (EU) 2017/745 application of MDR requirements to 'legacy devices' and to devices placed on the market prior to 26 May 2021 in accordance with Directives 90/385/EEC or 93/42/EEC"
- The Guideline MDCG 2020-6 "Regulation (EU) 2017/745: Clinical evidence needed for medical devices previously CE marked under Directives 93/42/EEC or 90/385/EEC - A guide for manufacturers and notified bodies"
- How to efficiently develop a PMS Plan according to standard ISO/TR 20416:2020 "Medical devices - Post-market surveillance for manufacturers"
- The tools for the manufacturer.

# PROGRAM

The course will be divided into two 4-hours modules, which can be attended separately:

- **MODULE 1 TECHNICAL/NORMATIVE:** technical/knowledge in-depth study on the Regulation (EU) 2017/745 and the MDCG guidelines
- **MODULE 2 EXPERIMENTAL:** regulatory overview + practical training on post-market surveillance activities

The course will be held in **e-learning** mode: it will be possible to participate online remotely.

#### SPEAKER

Dr. Marianna Mastroroberto - Clinical Manager Ente Certificazione Macchine

#### FOR INFORMATION:

Diego Stevanella - Sales Manager Medical Devices ph. (+39) 393 2471040 - diego.s@entecerma.it

