

# Essentials of European Medical Device Regulatory Affairs

22 September 2026



## Programme

Please note all times are in BST

| Time  | Presentation  | Speaker   |
|-------|---|---|
| 09:00 | Introduction from TOPRA   | TOPRA   |
| 09:05 | Introduction  | <b>Theresa Jeary</b><br>BSI Group                           |
| 09:15 | Scope and definitions   | <b>Angela Stokes</b><br>Sharp Regulatory Consulting Limited |
| 10:05 | Making available on the market and putting into service of device, obligations of economic operators, reprocessing, CE marking and free movement                                  | <b>Theresa Jeary</b>  |
| 10:40 | Break   |   |
| 10:55 | Identification and traceability of devices, registration of devices and of economic operators, summary of safety and clinical performance, European data base on medical devices. | <b>Theresa Jeary</b>  |
| 11:10 | Notified bodies   | <b>Theresa Jeary</b>  |
| 11:35 | Classification  | <b>Angela Stokes</b>  |
| 12:15 | Lunch Break   |   |
| 13:00 | Conformity Assessment – Annex IX – XI   | <b>Theresa Jeary</b>  |
| 13:45 | Clinical evaluation and clinical investigation  | <b>Angela Stokes</b>  |
| 14:15 | Break   |   |
| 14:30 | Post-market surveillance vigilance and market surveillance  | <b>Theresa Jeary</b>  |
| 15:05 | Cooperation between member states, medical device coordination group, expert laboratories, expert panels and device registers   | <b>Theresa Jeary</b>  |
| 15:15 | Confidentiality, data protection, funding and penalties   | <b>Theresa Jeary</b>  |
| 15:20 | Final provisions  | <b>Theresa Jeary</b>  |
| 15:30 | Annex 1 and labelling   | <b>Angela Stokes</b>  |
| 15.45 | Technical documentation   | <b>Theresa Jeary</b>  |
| 16:00 | Close   |   |