Introduction To The EU Medical Device Regulation (MDR) Training Course



COURSE CONTENT

- An overview of the new regulation
- The key changes you need to be aware of
- What EU MDR means in practice
- Where you can find further information
- A guiz to test your knowledge





AN OVERVIEW OF EU MEDICAL **DEVICE REGULATION**

LEARNING OBJECTIVES

Upon completion of this course, learners will be able to:

- Give an overview of EU Medical Device Regulation
- Understand implementation timescales

AUDIENCE

This course is intended for, but not limited to, regulatory personnel, managers, CEO, CFO, medical device industry's employees and medical device users who want to learn about the new regulation.

COURSE CERTIFICATION

On completion of this course:



- The SGS certificate is available to download once you have achieved a pass mark of 80% or more in the final assessment which completes this course.
- You have 12 months access to this course, effective from the date of purchase.









