

# MDR – THE NEW EU MEDICAL DEVICE REGULATION



**DIRECTIVE 93/42/EEC –**  
Medical Devices



**DIRECTIVE 90/385/EEC –**  
Active Implantable  
Medical Devices



**Medical Devices Regulation**  
**MDR**

## THE MOST IMPORTANT CHANGES

Extended area of application (includes non-medical devices)

UDI: Unique product number for every medical device

More stringent requirements for technical documentation (TD)

More stringent requirements for responsible persons: Expert knowledge of medical devices

New scrutiny procedure for high risk medical devices

More stringent requirements for clinical assessments and testing: Data collection to continue even after market launch

Notified bodies more strictly regulated: New bodies to be chosen and inspected

EUDAMED: Europe-wide database for more transparency and cooperation

## TRANSITIONAL PROVISIONS

