

#### REF. E-MDR-MOD-01 - E-LEARNING

Medical Device Regulation
Regulation (EU) 2017/745

**Overview of the European** 

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The Overview of the European Medical Device Regulation module presents the vocabulary and general principles of Regulation (EU) 2017/745.

#### **DURATION**

23 minutes, including a 15-minute lesson and 8 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

No prerequisite

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Learn the vocabulary and scope of the Regulation
- Become familiar with the major points of the Regulation
- Become aware of the parties governed by the Regulation

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. Definitions
- 2. Scope
- 3. Structure of the regulation
- 4. Parties involved

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







### REF. E-MDR-MOD-02 – <u>E-LEARNING</u>

**Operators** 

**Obligations of Economic** 

**Regulation (EU) 2017/745** 

EUROPEAN REGULATIONS

#### **COURSE DESCRIPTION**

The Obligations of Economic Operators module presents the parties who are governed by Regulation (EU) 2017/745 and the obligations incumbent on economic operators active on the Medical Device (MD) market

#### **DURATION**

**37 minutes**, including a 25-minute lesson and 12 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

Knowledge of the Regulation's main definitions and general principles (N.B. covered in the Overview module)

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Become familiar with the economic operators on the medical device market
- Identify the obligations incumbent on each of these operators

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. Parties involved
- 2. Manufacturer's obligations
- 3. Authorized representative's obligations
- 4. Importer's obligations
- 5. Distributor's obligations

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







### **EUDAMED & UDI**

**Regulation (EU) 2017/745** 

#### REF. E-MDR-MOD-03 - E-LEARNING

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The EUDAMED & UDI module presents the EUDAMED database and different types of device identifiers

#### **DURATION**

37 minutes, including a 25-minute lesson and 12 minutes of quizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

No prerequisite

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Understand EUDAMED and the main principles for entry into this database
- Become familiar with the different types of UDIs and their requirements

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. EUDAMED
- 2. UDI

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







### **Market Authorization**

**Regulation (EU) 2017/745** 

REF. E-MDR-MOD-04 - E-LEARNING

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The Market Authorization module presents the principles and steps to take to place Medical Devices (MD) on the market

#### **DURATION**

16 minutes, including a 10-minute lesson and 6 minutes of quizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

Knowledge of the Regulation's main definitions and general principles (N.B. covered in the Overview module)

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Learn the principles that govern market authorizations for Medical
- Become familiar with the market authorization conditions and steps for a Medical Device

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. Regulatory requirements
- 2. Market authorization steps
- 3. Parties involved

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







## Classification of Medical Devices

**Regulation (EU) 2017/745** 

REF. E-MDR-MOD-05 - E-LEARNING

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The Classification of Medical Devices (MD) module presents the different MD classes and the classification rules.

#### **DURATION**

23 minutes, including a 15-minute lesson and 8 minutes of guizzes



#### **E-LEARNING**

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

Familiarity with the Regulation's main definitions and general principles, more particularly regarding market authorization (N.B. covered in the Overview and Market Authorization modules)

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Understand what the different classes of medical devices refer to
- Know how to assign the right class to the different types of devices
- Become familiar with the principal special rules

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. Medical Device classes
- 2. Non-invasive devices
- 3. Invasive devices
- 4. Active devices
- 5. Special rules

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







### **Conformity Assessment**

**Regulation (EU) 2017/745** 

#### REF. E-MDR-MOD-06 - E-LEARNING

#### **EUROPEAN REGULATIONS**

#### **COURSE DESCRIPTION**

The Conformity Assessment module presents the procedures to assess conformity and the Technical Documentation's sampling principles in the framework of this assessment.

#### **DURATION**

**16 minutes**, including a 10-minute lesson and 6 minutes of quizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

Familiarity with the Regulation's main definitions and general principles, more particularly regarding market authorization, as well as classification and UDI rules (N.B. covered in the Overview, EUDAMED and UDI, Market Authorization and MD Classification modules)

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Understand the conformity assessment procedures for medical devices according to Regulation (EU) 2017/745
- Understand the Technical Documentation's sampling principles in the framework of Medical Device conformity assessments

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. Preamble
- 2. Conformity assessment procedures
- 3. Technical Documentation assessment via sampling
- 4. Quality Management System evaluation

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







REF. E-MDR-MOD-07 - E-LEARNING

**EUROPEAN REGULATIONS** 

### **General Safety and Performance Requirements**

**Regulation (EU) 2017/745** 

#### **COURSE DESCRIPTION**

The General Safety and Performance Requirements (GSPR) module presents the safety and performance, design and manufacturing requirements and the information that must be provided with a Medical Device (MD).

#### **DURATION**

37 minutes, including a 25-minute lesson and 12 minutes of guizzes



#### **E-LEARNING**

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

Knowledge of the Regulation's main definitions and general principles (N.B. covered in the Overview module)

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Become familiar with the safety and performance requirements set out in Annex I of the Regulation
- Differentiate the general requirements from those regarding performance, design and manufacturing
- Be aware of the information to be provided with an MD

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. General requirements
- 2. Performance, design and manufacturing requirements
- 3. Requirements relative to information provided with the device

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation guiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







### **Technical Documentation**

**Regulation (EU) 2017/745** 

#### REF. E-MDR-MOD-08 - E-LEARNING

#### **EUROPEAN REGULATIONS**

#### **COURSE DESCRIPTION**

The Technical Documentation module presents the content of the Technical Documentation and the information in the documentation regarding post-market surveillance.

#### **DURATION**

23 minutes, including a 15-minute lesson and 8 minutes of guizzes



#### **E-LEARNING**

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

Familiarity with the Regulation's main definitions and general principles, and more particularly the General Safety and Performance Requirements (N.B. covered in the Overview and GSPR modules)

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Familiarity with the Regulation's Annexes that set out the content of the Technical Documentation (TD)
- Familiarity with the requirements in terms of the TD content for the EU conformity declaration
- Familiarity with the documentation's elements on post-market surveillance

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. Prerequisites for the conformity declaration (Annex II)
- 2. Post-market surveillance (Appendix III)

#### **ADDITIONAL BENEFITS**

**Summary** presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







### **Clinical Evaluation**

**Regulation (EU) 2017/745** 

#### REF. E-MDR-MOD-09 - E-LEARNING

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The Clinical Evaluation module presents the clinical evaluation methodology and the data to be used for this evaluation.

#### **DURATION**

37 minutes, including a 25-minute lesson and 12 minutes of quizzes



#### **E-LEARNING**

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

Knowledge of the Regulation's main definitions and general principles (N.B. covered in the Overview module)

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Learn the methodology for a clinical evaluation
- · Know which data to use and how to obtain it
- Know how to analyze and organize clinical data to comply with regulatory requirements

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. Clinical evaluation objectives and methodology
- 2. Nature, source and equivalence of the data
- 3. Clinical evaluation report

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







## Vigilance

**Regulation (EU) 2017/745** 

Post-Market Surveillance &

REF. E-MDR-MOD-10 – E-LEARNING

#### **EUROPEAN REGULATIONS**

#### **COURSE DESCRIPTION**

The Post-Market Surveillance & Vigilance module describes the purpose of post-market surveillance and the procedures to implement based on the type of data collected.

#### **DURATION**

30 minutes, including a 20-minute lesson and 10 minutes of quizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

Knowledge of the Regulation's main definitions and general principles, more particularly those regarding product classification and the principles to demonstrate and monitor clinical benefits (N.B. covered in the Overview, MD Classification and Clinical Evaluation modules)

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EDUCATIONAL GOALS**

- Understand the need for post-market surveillance
- Become familiar with the procedures to apply based on the type of data collected

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. Post-market surveillance (PMS)
- 2. Claim processing
- 3. Vigilance
- 4. Processing other relevant information
- 5. Post-market clinical follow-up (PMCF)

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







### **Quality Management System**

**Regulation (EU) 2017/745** 

#### REF. E-MDR-MOD-10 - E-LEARNING

#### **EUROPEAN REGULATIONS**

#### **COURSE DESCRIPTION**

The Quality Management System module presents the manufacturer's obligations regarding the Quality Management System and how it is assessed.

#### **DURATION**

23 minutes, including a 15-minute lesson and 8 minutes of quizzes



#### **E-LEARNING**

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The module is composed of a lesson in the form of an interactive video with voice-off and learning activities. The module can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector

#### **PREREQUISITES**

Familiarity with the Regulation's main definitions and general principles, more particularly market authorization procedures, conformity assessments and post-market surveillance (N.B. covered in the Overview, Market Authorization, Conformity Assessment and Post-Market Surveillance and Vigilance modules)

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of two months within two business days at most of approval of your order.

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EDUCATIONAL GOALS**

- Become familiar with the manufacturer's obligations regarding the quality management system
- Understand how this QMS is assessed by a notified body

#### **COURSE CONTENT**

Assistance with navigation/Module objectives/Presentation of the course content

- 1. Requirements applicable to the Quality Management System
- 2. Quality Management System evaluation
- 3. Quality Management System evaluation request

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This training is assessed based on each individual's acquisition of knowledge via a quiz at the end of the program (success criteria >= 80%). If trainees pass the quiz, they receive their training certificate, as well as the corrected quiz, by email.







#### REF. E-MDR-F-01 – E-LEARNING

EUROPEAN REGULATIONS

# Package for MANUFACTURERS

**Regulation (EU) 2017/745** 

#### **COURSE DESCRIPTION**

The MANUFACTURERS' package is composed of 11 modules and has been designed specifically to present, in an interactive format, the main obligations of Medical Device (MD) manufacturers to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

310 minutes, including 200 minutes of lessons and 110 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly MD manufacturers

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of six months within two business days at most of approval of your order.

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on manufacturing activities

#### **COURSE CONTENT**

- 1. Overview of Regulation (EU) 2017/745
- 2. Obligations of economic operators
- 3. EUDAMED & UDI
- 4. Market authorization
- 5. Classification of Medical Devices
- 6. Conformity assessment
- 7. General Safety and Performance Requirements
- 8. Technical Documentation
- 9. Clinical Evaluation
- 10. Post-Market Surveillance & Vigilance
- 11. Quality Management System

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 11 modules (final quiz/success criteria >= 80%). If trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.







#### REF. E-MDR-F-02 - E-LEARNING

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**EUROPEAN REGULATIONS** 

## Package for DISTRIBUTORS

**Regulation (EU) 2017/745** 

#### **COURSE DESCRIPTION**

The DISTRIBUTORS' package is composed of 7 modules and has been designed specifically to present, in an interactive format, the main obligations of Medical Device distributors to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

**195** minutes, including 125 minutes of lessons and 70 minutes of quizzes



#### **E-LEARNING**

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly MD distributors

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on distribution activities

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of four months within two business days at most of approval of your order.

#### **COURSE CONTENT**

- 1. Overview of Regulation (EU) 2017/745
- 2. Obligations of economic operators
- 3. EUDAMED & UDI
- 4. Market authorization
- 5. Classification of Medical Devices
- 6. Post-Market Surveillance & Vigilance
- 7. Quality Management System

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 7 modules (final quiz/success criteria >= 80%). If trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.







#### REF. E-MDR-F-03 - E-LEARNING

EUROPEAN REGULATIONS

## Package for IMPORTERS

Regulation (EU) 2017/745

#### **COURSE DESCRIPTION**

The IMPORTERS' package is composed of 7 modules and has been designed specifically to present, in an interactive format, the main obligations of Medical Device importers to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

195 minutes, including 125 minutes of lessons and 70 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly MD importers

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on importing activities

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of four months within two business days at most of approval of your order.

#### **COURSE CONTENT**

- 1. Overview of Regulation (EU) 2017/745
- 2. Obligations of economic operators
- 3. EUDAMED & UDI
- 4. Market authorization
- 5. Classification of Medical Devices
- 6. Post-Market Surveillance & Vigilance
- 7. Quality Management System

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 7 modules (final quiz/success criteria >= 80%). If trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.







# Package for AUTHORIZED REPRESENTATIVES

**Regulation (EU) 2017/745** 

REF. E-MDR-F-04 - E-LEARNING

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The AUTHORIZED REPRESENTATIVES' package is composed of 9 modules and has been designed specifically to present, in an interactive format, the main obligations of authorized representatives of Medical Devices to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

235 minutes, including 150 minutes of lessons and 85 minutes of guizzes



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#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly authorized representatives of MD

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on authorized representatives' activities

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of six months within two business days at most of approval of your order.

#### **COURSE CONTENT**

- 1. Overview of Regulation (EU) 2017/745
- 2. Obligations of economic operators
- 3. EUDAMED & UDI
- 4. Market authorization
- 5. Classification of Medical Devices
- 6. Conformity assessment
- 7. Technical Documentation
- 8. Post-Market Surveillance & Vigilance
- 9. Quality Management System

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 9 modules (final quiz/success criteria >= 80%). If trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.







REF. E-MDR-F-05 – E-LEARNING

**EUROPEAN REGULATIONS** 

### **Package for REGULATORY AFFAIRS**

Regulation (EU) 2017/745

#### **COURSE DESCRIPTION**

The REGULATORY AFFAIRS package is composed of 11 modules and has been designed specifically to present, in an interactive format, the main Regulatory Affairs obligations to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

310 minutes, including 200 minutes of lessons and 110 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly MD Regulatory Affairs Departments

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on the Regulatory Affairs Department

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of six months within two business days at most of approval of your order.

#### **COURSE CONTENT**

- Overview of Regulation (EU) 2017/745
- 2. Obligations of economic operators
- 3. **EUDAMED & UDI**
- 4. Market authorization
- 5. Classification of Medical Devices
- Conformity assessment 6.
- 7. General Safety and Performance Requirements
- 8. **Technical Documentation**
- 9 Clinical evaluation
- Post-Market Surveillance & Vigilance 10
- Quality Management System 11

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 11 modules (final quiz/success criteria >= 80%). If trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.







# Package for RESEARCH & DEVELOPMENT

Regulation (EU) 2017/745

REF. E-MDR-F-06 – E-LEARNING

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The RESEARCH & DEVELOPMENT package is composed of 6 modules and has been designed specifically to present, in an interactive format, the main obligations of the R&D Department to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

185 minutes, including 120 minutes of lessons and 65 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly the R&D Department

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on research and development activities.

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of four months within two business days at most of approval of your order.

#### **COURSE CONTENT**

- 1. Overview of Regulation (EU) 2017/745
- 2. Obligations of economic operators
- 3. EUDAMED & UDI
- 4. Classification of Medical Devices
- 5. General Safety and Performance Requirements
- 6. Technical Documentation

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 6 modules (final quiz/success criteria >= 80%). When trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.







# Package for QUALITY ASSURANCE

**Regulation (EU) 2017/745** 

REF. E-MDR-F-07 - E-LEARNING

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The QUALITY ASSURANCE package is composed of 7 modules and has been designed specifically to present, in an interactive format, the main obligations of the Quality Assurance Department to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

175 minutes, including 110 minutes of lessons and 65 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly the Quality Assurance Department

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on the Quality Assurance Department's activities

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of four months within two business days at most of approval of your order.

#### **COURSE CONTENT**

- 1. Overview of Regulation (EU) 2017/745
- 2. Obligations of economic operators
- 3. Market authorization
- 4. Classification of Medical Devices
- 5. Conformity assessment
- 6. Post-Market Surveillance & Vigilance
- 7. Quality Management System

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 7 modules (final quiz/success criteria >= 80%). When trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.







# Package for CLINICAL DEPARTMENTS

**Regulation (EU) 2017/745** 

#### REF. E-MDR-F-08 - E-LEARNING

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The CLINICAL DEPARTMENTS' package is composed of 5 modules and has been designed specifically to present, in an interactive format, the main obligations of the Clinical Department to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

155 minutes, including 100 minutes of lessons and 55 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly the Clinical Department

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on the Clinical Department's activities

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of four months within two business days at most of approval of your order.

#### **COURSE CONTENT**

- 1. Overview of Regulation (EU) 2017/745
- 2. Classification of Medical Devices
- 3. General Safety and Performance Requirements
- 4. Clinical Evaluation
- 5. Post-Market Surveillance & Vigilance

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 5 modules (final quiz/success criteria >= 80%). If trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.







# Package for PRODUCTION/PURCHASING

**Regulation (EU) 2017/745** 

REF. E-MDR-F-09 - E-LEARNING

**EUROPEAN REGULATIONS** 

#### **COURSE DESCRIPTION**

The Production/Purchasing package is composed of 5 modules and has been designed specifically to present, in an interactive format, the main obligations of the Production and Purchasing Departments to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

150 minutes, including 95 minutes of lessons and 55 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly the Production/Purchasing Departments

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on the Production/Purchasing Departments' activities

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of four months within two business days at most of approval of your order.

#### **COURSE CONTENT**

- 1. Overview of Regulation (EU) 2017/745
- 2. EUDAMED & UDI
- 3. General Safety and Performance Requirements
- 4. Technical Documentation
- 5. Quality Management System

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 5 modules (final quiz/success criteria >= 80%). When trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.







#### REF. E-MDR-F-10 - E-LEARNING

KLI. L-MDK-I-IO - L-LLAKINING

# Package for SALES DEPARTMENTS

**Regulation (EU) 2017/745** 

### EUROPEAN REGULATIONS

#### **COURSE DESCRIPTION**

The SALES DEPARTMENTS' package is composed of 6 modules and has been designed specifically to present, in an interactive format, the main obligations of the Sales Department to ensure compliance with Regulation (EU) 2017/745.

#### **DURATION**

170 minutes, including 110 minutes of lessons and 60 minutes of guizzes



**E-LEARNING** 

#### **TECHNICAL PREREQUISITES**

Computer and internet connection

#### **METHODS USED**

The modules are composed of lessons in the form of interactive videos with voice-off and learning activities. They can be accessed 24/7. Reminders are sent to monitor knowledge acquisition and avoid a break in the learning curve.

#### **TARGET TRAINEES**

All individuals involved in the MD sector, and more particularly the Sales Department

#### **EDUCATIONAL GOALS**

- Understand the regulatory framework for medical devices
- Identify the requirements that apply and their impact on the Sales Department's activities

#### **PREREQUISITES**

Familiarity with the MD sector

#### **ACCESS**

The CEISO ACADEMY platform can be accessed for a period of four months within two business days at most of approval of your order.

#### **COURSE CONTENT**

- 1. Overview of Regulation (EU) 2017/745
- 2. Obligations of economic operators
- EUDAMED & UDI
- 4. Market authorization
- 5. Post-market surveillance & vigilance
- 6. Quality Management System

#### **ADDITIONAL BENEFITS**

Summary presentation of the regulation

Support materials designed by experienced consultants-trainers

Interactive lesson

Short modules that can be accessed as often as required and autonomously

As an option, you may select Q&A sessions with one of our consultants-trainers

#### **EVALUATION METHODS**

Course evaluation quiz Satisfaction survey Attendance certificate

This package is assessed based on each individual's acquisition of knowledge via a quiz at the end of each module (practice quiz) and one at the end of the 6 modules (final quiz/success criteria >= 80%). When trainees pass the final quiz, they receive their training certificate, as well as the corrected quiz, by email.



