



Process Change Assessment

Course description

The objective of this course is to provide students with an understanding of the change control process, interaction with different elements of the quality system and considerations when creating a process change assessment in the medical devices industry.

Throughout the course, students will engage in hands-on learning experiences, applying their knowledge of change control in the medical device industry, ensuring regulatory compliance, maintaining quality standards, and addressing real-world challenges.

At the end of the course you will be able to:

- Understand the concepts of change control in the medical device industry.
- Understand an overview of the medical device change control and regulatory and quality management system (QMS) compliance.
- Understand the types of changes according to different criteria.
- Learn the elements of the impact assessment in medical device change control process.
 - Change Description
 - Validation Assessment
 - Scope of Chance
 - Production and Process Assessment
 - Risk Assessment
 - Regulatory Assessment
- Experience in a practical way, real-world situations, and examples on change control.

Main topics

- 1 Regulatory Requirements.
- 2 Definition of Change.
- 3 Change Workflow.
- 4 Change Types.
- 5 The Impact Assessment:
 - Change Description
 - Validation Assessment
 - Scope of Chance
 - Production and Process Assessment
 - Risk Assessment
 - Regulatory Assessment

Contact

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Course features

- Instructor Led 
- Duration: 4 hours 
- Tools and templates 
- Simulated learning 
- Course certification 