

COURSE ID 3091

Scope of the FDA

History and FDA regulations

DIRECTED BY

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Course Topics Include:

- What are GMPs
- Legal Issues
- History of GMPs
- Purpose of GMPs

about the course

Most industries in the US are regulated by local, state, and federal regulators and pharmaceutical manufacturing companies are some of the most highly regulated, particularly at the federal level. Pharmaceutical Quality affects every American hence the FDA regulates the quality of pharmaceuticals very carefully. The main regulatory standard for ensuring pharmaceutical quality are the Current Good Manufacturing Practice (cGMPs) regulations.

This 90-minute, accredited course will discuss the background of the cGMPs and their applicability to the pharmaceutical industry. The course will also cover how the cGMPs are enforced by FDA and how companies should work to comply with them.

This training is one part of the 10-course series required for the GMP Professional Certification Program. Attend this as a step in the certification process or as a stand-alone course for personal career advancement and training.



who should attend

This course has been developed to meet the training needs of professionals in the Pharmaceutical, Medical Device and Cosmetic Industries. It will be especially valuable to those engaged in compliance with FDA regulations, Compliance, QA/QC, Formulation, Production, Manufacturing and Regulatory areas.

Engineers, chemists, QA/QC and manufacturing personnel will benefit greatly by attending this course by understanding the extent of the FDA regulations and how they affect daily job responsibilities.

learning objectives

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

- Outline the regulations to properly comply
- Contrast and summarize current regulations and guidelines
- Outline how the regulations affect you and your job function
- Comprehend the meaning of "c" in cGMP
- Explain why and how regulations came about
- Show what pharmaceutical product types are affected
- Explain to Management who have the Oversight of GMP and GDP Compliance

course outline

Review of Learning Objectives Scope of the FDA

- Background to the GMPs
- How they evolved
- Purpose of the regulations
- Industry references

Roles and Responsibilities

- Manufacturing
- Quality Assurance/Quality Control
- Management

Question and Answer Session Assessment Opportunity

course instructor

Mr. Angelucci is a pharmaceutical professional with over 30 years' experience in Quality Assurance, Quality Control, Regulatory Affairs, Validation, consent decree remediation as well as cGMP Compliance in the Medical Device and Pharmaceutical industries

He has worked for several well-known companies including Johnson & Johnson, Bristol-Meyer Squibb, Pfizer, Schering Plough and Merck. His experience has been either as a direct employee or as a contracting consultant while employed at Foster Wheeler and Aker Kvaerner. As a consultant Mr. Angelucci has worked at various project locations domestically and in Europe and Asia. With these firms he has been involved with auditing, GMP training, Bio-tech consulting as well as FDA audit and PAI readiness training and participation.



Mr. Angelucci is a degreed engineer with two master's degrees in engineering, holds industry certifications with ASQ as a CQE, CQA and CPGP. Previously, he was the ASQ Philadelphia chapter Education Chair. In addition, he holds a PMP certification through the Project Management Institute. He has published numerous articles on the subjects of Validation and compliance and has been a speaker to industry groups such as ISPE, IVT, DIA, PDA, ASQ and CFPA.

Accreditations

International Accreditors for Continuing Education and Training (IACET)



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