

*Webinar on*

# **Best Practices for Deviation Training**

*Date : August 18, 2021*

# Areas Covered

- *Review of FDA and Regulatory Requirements for Investigations*
- *What is the definition of a Deviation?*
- *Types of Deviations/Identification of Deviations. Conducting the Investigation*
- *Interviews – dos and don'ts, Source Documents/Evidence*
- *Determining Root Cause Effective/Sustainable CAPA to observations*
- *Key Elements of the Investigation Report*

# Learning Objectives

- Discuss what to do when problems occur*
- Outline the requirements of the Deviation and procedure including the deviation report*
- Choose the most appropriate Root Cause Analysis methods for the situation*
- Discuss how to conduct the deviation and the tools to be used in the process*

In this webinar you will learn how to identify and avoid potential pitfalls during deviation investigations.

**PRESENTED BY:**

*Danielle DeLucy, MS, is the owner of ASA Training and Consulting, LLC which provides Pharmaceutical and Biologics based companies with training and quality systems assistance in order to meet regulatory compliance. Prior to this role, Danielle has been in the industry for 15 years serving in numerous Quality Management Roles, such as the Director of Product Quality.*



Date : August 18, 2021

Time : 01 : 00 PM EST

Duration : 60 Minutes

Price: \$179

# Webinar Description

Deviation investigations are a very common topic to be reviewed during FDA investigations. Ensuring the firm has robust and complete write-ups of these situations is key to a successful audit. Having inadequate investigations could potentially cause some severe issues for the company and may sometimes result in warning letters, consent decrees or recalls.



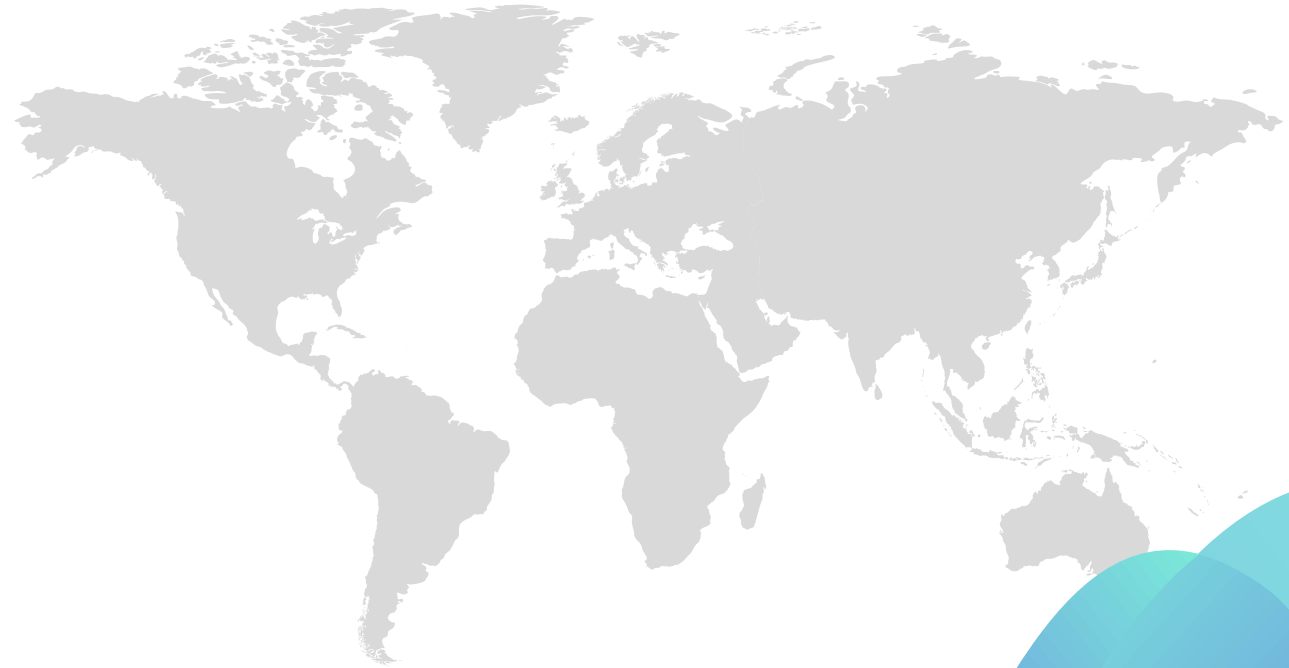
# Who Should Attend ?

- *Site Quality Operations Managers*
- *Quality Assurance Personnel*
- *Plant Managers and Supervisors*
- *Manufacturing Superintendents and Managers*
- *Regulatory Affairs Managers*
- *QC Lab staff*



# Why Should You Attend ?

*This webinar will help attendees understand the fundamental steps of a deviation investigation with a focus on using facts and objective evidence to arrive at root cause and CAPA. This webinar will focus on how to avoid the pitfalls that may occur during FDA inspections and help eliminate 483 observations. Learn how to identify and avoid potential pitfalls during deviation investigations.*



To register please visit:

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