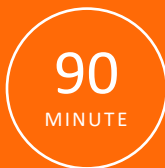


# Data Integrity Practices for the Laboratory and Beyond

GMP Laboratory Control Professional Certification Program

DIRECTED BY

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ACCREDITED  
COURSE

#### Course Topics Include:

- Improving Data Integrity in laboratory Environment
- Learning from Real World FDA Citations
- Mitigating Common Data Integrity Issues

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## about the course

Data integrity is the assurance that data records are accurate, complete, intact and maintained within their original context, including their relationship to other data records. This definition applies to data recorded in electronic and paper formats or a hybrid of both. To assure the quality of raw materials, in process materials and finished goods, laboratory data integrity is assuming greater importance in current Good Manufacturing Practices (CGMP) for US Food and Drug Administration (FDA)-regulated industry. Data integrity and security infractions are not only 21 Code of Federal Regulations (CFR) Part 11 issues but also severe CGMP violations. The reasoning behind this complex issue is quite simple: if the integrity of laboratory data is compromised, batches of finished goods may not comply with regulatory authorization terms and, consequently, will not be released for sale.

In this 90-minute accredited training, we will discuss effective batch record review and assurance to Data Integrity compliance.

**This training is one part of the 10-course series required for the GMP Laboratory Control Professional Certification Program.**

Attend this as a step in the certification process or as a stand-alone course for personal career advancement and training.

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## who should attend

This course is intended for professionals in the pharmaceutical industry who are charged with all aspects of maintaining compliant data integrity practices.

Personnel working in Quality Control, Manufacturing, Regulatory Affairs and Quality Assurance along with their management and supervisory teams will benefit greatly from this training.

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## learning objectives

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

- Describe the current regulatory position on data integrity
- Discover the criteria for data integrity
- Recognize what needs to be addressed to ensure data integrity within a regulated GXP laboratory
- Implement approaches to improve data integrity in a laboratory environment
- Discuss Part 11 compliance
- Review FDA citations related to data integrity issues

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## course outline

### Review of Learning Objectives

#### Module 1: Background and Regulations related to Data Integrity

- Regulatory requirements for Data Integrity
- Definition of ALCOA

#### Module 2: Data Integrity Basics

- Identifying Common Issues relating to Data Integrity
- How to mitigate issues

#### Module 3: Examples of Proper Data Integrity

- What Regulators look For
- Real World Citations about Data Integrity

### Question and Answer Session

### Assessment Opportunity

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## course instructor

**Danielle DeLucy, MS**, is currently the owner of ASA Training & Quality Consulting, LLC, an Independent Consultant Agency to the Biologics and Pharmaceutical Industries specializing in the areas of Quality Assurance and Quality Systems. Prior to this role, Danielle has been in the industry for 18 years serving in numerous Quality Management Roles, such as the Director of Product Quality, the oversight of Sterility Assurance practices, and provided QA oversight of numerous filling and packaging operations. Danielle began her QA career as a Quality Control Pharmaceutical Microbiologist at a contract laboratory where she performed various tests for their clients. In the years after, she has held positions in the Quality management arena while increasing her responsibility. She has helped to lead many Regulatory Health Inspections and was instrumental in the coaching process of her peers prior to any inspection. Currently, Danielle assists companies who are faced with warning letters and consent decrees establish more robust quality systems so that the company can succeed.

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## Accreditations



### **International Accreditors for Continuing Education and Training (IACET )**

Cobblestone has been approved as a CEU Accreditor by IACET and awards CEUs for participation in qualified courses. Cobblestone has demonstrated that it complies with the ANSI/IACET Standards and is authorized to offer IACET CEUs for its programs. CEUs will be awarded for participation in Cobblestone’s courses at the rate of .1 CEU per contact hour upon successful completion of the entire course and 70% accuracy in the required Learners’ Assessment. A minimum score of 80% is required for all courses within a Cobblestone Certification Program. This course offers a total of 1.5 contact hours, or .2 CEUs. For further information, visit [www.iacet.org](http://www.iacet.org)



### **Regulatory Affairs Professional Society (RAPS)**

Cobblestone is committed to enhancing the ongoing professional development of regulatory affairs professionals and other stakeholders through appropriate regulatory affairs learning activities and programs. Cobblestone has agreed to follow RAPS- established operational and educational criteria. This course may be eligible for up to 12 credits towards a participant’s RAC recertification upon full completion. The requirements and standards for recertification are developed and administered by the Regulatory Affairs Certification Board (RACB), which manages all areas of the RAC program. Additional information about RAC is available on the RAPS website at [RAPS.org/rac](http://RAPS.org/rac).