



John Cuspilich, Sr. Auditor
The Auditing Group

John is the CEO and Senior Auditor for The Auditing Group, Inc and GMP Boot Camps. In addition, John also serves as the Senior Editor at **GMP Publications, Inc.** and the CEO of **The Validation Group.**

www.johncuspilich.com

John has conducted more than 1200 GxP audits, Gap Analysis and remediation projects world-wide, with over 35 years, hands-on technical and management level experience within the Food, Supplements, Pharmaceutical, Biotechnology, Medical Device, Cosmetics, Petrochemical, Electronic Systems, Validation, and regulated industries.

John has conducted hundreds of speaking engagements, seminars and bootcamp style training seminars world-wide. Published, co-sponsored and conducted audit reviews of thousands of technical and professional papers, journals and books for hundreds of Companies in the regulated industry.

John has assisted hundreds of companies in meeting and exceeding regulatory compliance, pertaining to 'for-cause' or 'due-diligence' initiatives. Assisting companies to achieve, resolve, remediation and exceed regulated industry requirements, mandates, 'for-cause' and 'due-diligence' priorities with the technique of promoting GxP standards and practices through interactive hands-on training.

John has extensive knowledge in industry standards; FDA (CDER, CBER, CDRH, CVM, CFSAN), cGMP, GLP, ICH, OECD, GAMP, ISO, OECD, OSHA, HACCP, HIPPA, EPA and GCP regulations with thorough knowledge in implementation of these standards.

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federal regulations

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21 CFR Part 11
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GMP Boot Camps
Divisions of GMP Publications and
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856-437-588033

info@auditing.com



Course Agenda

GMP QMS 101 The Basics - The required GMP Training for all employees who work in regulated industry.

- GMP and the GMP Focus and the GMP Lifestyle
- The Predicate Rules
- The FDA Agency, Inspections, Warning letters and 483s
- Quality Terminology

The QMS

- The Basics of QMS
 - CAPA - The basics about Corrective and Preventive Actions;
 - Non-Conformance - Materials Supply, to End of Use Complaints;
 - Change Control - Documentation, Engineering, Production and Distribution;
 - Deviations - Deviation management essentials;
 - Out of Specifications / Out of Trend - Management of OOS and OOT;
 - Complaints - Receipt, Qualifications, Risk Assessments and Remediation/Resolution;
 - Recalls - The process, and indications of failures;
 - Product Traceability;
 - Audit - Internal, Agency, Customers and External Audit;
 - Vendors, Suppliers and Contractors - Quality Agreements;
 - The Meetings - Management and Quality Meetings;
- The 11 General Orders / Principals of GMP
 1. Writing Procedures (with template examples);
 2. Following Procedures (and the failures that occur);
 3. Documentation (Requirements, and the Regulations);
 4. Validation (Concepts, and basic process requirements);
 5. Design/Build Facilities and Equipment (In-depth look into a GMP facility requirement);
 6. Maintain Facilities and Equipment (Maintenance, Calibration, Use and Cleaning of Facilities/Equipment);
 7. Competency (The Training Requirements);
 8. Sanitation and Good Housekeeping Practices (General requirements for all facilities);
 9. Control for Quality (from Materials Receipt, Production, Packaging and Distribution);
 10. Audit Requirements
 11. Prepare for Battle! The Traceability Process;

21 CFR Part 11 Electronic Records

- 21 CFR Part 11 Basic Overview
- Definitions, System Types and Classifications
- 21 CFR Part 11 – Predicate Rule
- Part 11.10 Sections a) - k)
- Microsoft Excel



856-596-2333

info@gmpbootcamps.com

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- 1 Day GMP Training – Good Laboratory Practice (GLP)
- 1 Day and 2 Day Good Clinical Practice (GCP) Training
- 1 Day and 2 Day Compounding Facilities 503b
- 1 Day – DEA GMP Training
- 1 Day – Good Engineering Practice Training – Design/Build Requirements for Drug Manufacturing
- 1 Day Good Validation Practice – Process
- 1 Day and 2 Day – Good Validation Practice – Software and Systems – 21 CFR Part 11
- 1 Day and 2 Day – Good Validation Practice – Equipment (Process and Utilities)

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