

T. Harris, MS, CQA

Senior Consultant 2300 Bethelview Road, Suite 110-189, Cumming, GA 30040 P: (470) 423-0185

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SKILL SET

Regulations: 21 CFR 210, 211, 11, 50, 111, 117 and 820, FD&C Act, ICH, ISO standards, EudraLex Parts I and II & Annexes; Annex 1, PIC/S, Good Distribution Practices (GDP), Good Clinical Practices (GCP), Good Documentation Practices, Data Integrity Principles and worldwide GMPs

Quality Assurance: GMP audits, Inspection Readiness Activities, Sterile Manufacturing, GMP and Auditor Training, Quality Management Systems (QMS), GMP Compliance, Gap Assessments, Risk Evaluation, Minister of Health Authority Responses, Investigations, Change Control, Process Validation, Consent Decree Remediation, Corrective and Preventive Actions (CAPA), Quality Metrics Assessment, Artificial Intelligence, Medical Device Design Controls and 510k

Quality Control: Data Collection, Data Integrity & Security, Data Analysis and Review, Microbiological Testing, UV/Vis, Chromatography and Spectrometry

Products/Services: Finished Drugs, Active Pharmaceutical Ingredients (API), Drug Substances, Biologics, Vaccines, Medical Devices, Combination Products, Raw Materials, Cosmetics & Cosmetic Ingredients, Packaging & Labeling and Distribution

Applications: Microsoft Office, Laboratory Information Management System (LIMS), Agilent, ChemStation, Empower, SharePoint, SAP, JD Edwards, TrackWise, and online learning platforms

Certifications: Former USFDA Certified Level I drug Investigator, Certified Quality Auditor (CQA), ASQ; Quality Instructor Certification, Train the Trainer, Baxter Healthcare; Pharmacovigilance Auditing Certification, RQA; Sterilization & Sterility Assurance Certification, Baxter Healthcare and FDA-certified training in GMP compliance and regulatory standards

Other Skills: Project Management, Computer System Validation, Regulatory Affairs and Physical & Analytical Chemistry

PROFESSIONAL EXPERIENCE

Polymath Regulatory Consultants, LLC GMP and Quality Consulting *Independent Contractor*

Sr. Quality ConsultantFebruary 2025 - present
Atlanta Metro Area



Baxter Healthcare Corp.

Senior Global Quality Technical Consultant

Regulatory Quality Assurance Corporate Compliance Division May 2017 – September 2024 Deerfield, IL

- Led pharmaceutical and medical device manufacturing audits driving compliance to applicable procedures, regulations and standards
- Performed global GMP compliance audits to assess quality management systems
- Evaluated and approved CAPAs, risk assessments, root cause analysis and remediation plans
- Conducted inspection readiness activities and regulatory compliance training
- Assessed manufacturing processes, test methods, validation and qualification and data integrity
- Provided subject matter expertise on regulatory intelligence and compliance
- Independently conducted pharmacovigilance audits
- Developed audit procedures and compliance initiatives
- Reviewed and evaluated quality control and manufacturing data for ensure compliance with requirements.
- Collaborated with contract facilities, internal sites, and regulatory authorities to facilitate audits, achieving 70% decrease in number of regulatory actions
- Trained and qualified lead auditors and evaluated performance
- Collaborated with all levels of internal management across functions and business units to foster culture of compliance quality, facilitating collaboration and promoting transparency across organization

US Food & Drug Administration

Consumer Safety Officer/Investigator

Office of Regulatory Affairs

January 2009 – May 2017 Cincinnati, OH

- Executed complex inspections and investigations of drug manufacturing facilities, ensuring compliance with safety and quality standards.
- Led numerous pre-approval inspections for drug applications including NDAs, INDs, ANDAs, NADAs, ANADAs and OTCs.
- Independently conducted CGMP inspections for pharmaceuticals (e.g., API, oral solid, liquid, sterile and nonsterile) and dietary supplements
- Assessed manufacturing processes, sterility assurance, microbiological controls, analytical testing, OOS investigations and data integrity
- Provided regulatory assessments on pre-market and post-market activities
- Authored comprehensive inspection reports (EIRs) using the FD&C Act, Compliance Programs, Guidance Manuals, and applicable 21 CFR regulations.
- Acted as a subject matter expert and expert witness

Sinclair Community College

Chemistry Instructor August 2001 – September 2008

Dayton, OH

Relevant Courses Taught: Intro to General Chemistry, Intro to Organic Chemistry, Laboratory

T. Harris Page 2 of 3



EDUCATION

Wright State University
Master of Science in Organic Chemistry

Dayton, OH (USA)

Wilberforce University
Bachelor of Science in Chemistry

Wilberforce, OH (USA)

T. Harris Page 3 of 3