QC Compliance Manager
Focus on Small-Molecule APIs and Drug Products!

SPEAKERS:

Dr Thomas Backensfeld
Bayer AG, Germany

Dr Raphael Bar
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Dr Kerstin Hartisch
Bayer AG, Germany

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27 – 29 March 2019, Vienna, Austria

PROGRAMME:

- Regulatory Requirements for Analytical Labs (EU and U.S.)
- Analytical Instrument Qualification According to USP <1058>
- Computerised Systems in Analytical Labs
- Sampling of APIs and Excipients
- Documentation
- Specifications, SOPs, Test Procedures
- Laboratory Data Integrity
- Analytical Methods
  - Lifecycle Approach
  - Method Transfer and Equivalence Testing
- Managing OOS/OOT Results
- Stability Data: Presenting and Evaluating
- QA Aspects Applicable for QC Compliance Managers

Participate in 4 Workshops

This education course is recognised for the ECA GMP Certification Programme „Certified Quality Control Manager“. Please find details at www.gmp-certification.eu
Objectives

This Education Course will give a comprehensive overview of the main GMP requirements for quality control laboratories, from a European as well as from the U.S. (FDA) perspective. It is the aim of the course to address the challenges that QC Compliance Managers face today regarding the relevant regulatory requirements and how to successfully implement these requirements in the analytical lab.

Background

Due to changing regulatory requirements pharmaceutical Quality Control Compliance Managers are continuously facing new challenges. There are many regulatory requirements relevant for the pharmaceutical quality control, both in EU and in the US, for instance:

- EU GMP Guide (Part 1 / Part 2 / Annexes)
- 21 CFR Part 210/211 (USA)
- Guidances (EMA and FDA)
- ICH Guidelines
- WHO and PIC/S Recommendations
- Pharmacopoeias (Ph.Eur., USP)

QC Compliance Managers must be familiar with all these GMP-related topics and must be aware of the latest updates and the current interpretation of all these guidance documents.

In addition, analytical QC laboratories are increasingly in the focus of GMP inspections, both in Europe and in the US. For instance after FDA inspections, many laboratory-specific citations can be found in 483s and Warning Letters. And many findings related to the laboratory can also be found after inspections of European GMP supervisory authorities. Key compliance requirements include:

- Change control systems
- Calibration and qualification of analytical instrument
- Reference standards
- GMP compliant documentation
- Validation of analytical methods
- Stability program
- Validation of computerised systems
- Procedures for handling OOS results

All these key compliance issues will be addressed in this course and the main topics also deepened in workshops.

This year’s programme has been further updated to include the current challenges of Laboratory Data Integrity issues in analytical labs about reporting and invalidating laboratory results.

Please note that the emphasis of this course is on small-molecule pharmaceuticals. The course will not focus on biotech products.

Target Audience

This Education Course will be of significant value to
- Laboratory managers,
- Quality control managers,
- Analytical scientists,
- Senior laboratory staff
from quality control units in the pharmaceutical industry who are responsible for GMP Compliance in the analytical laboratory.

Programme

Regulatory Requirements for Analytical Labs and QC (EU and US)
- EU GMP Guide Part 1
- EU GMP Guide Part 2
- US 21 CFR Part 210/211
- FDA Guidelines for Industry with relevance for labs
- Inspection of analytical labs (EMA, FDA, etc.)
- FDA Warning Letters relating to QC

Analytical Instrument Qualification
- USP General Chapter <1058> Analytical Instrument Qualification
- Risk Analysis
- Qualification steps: DQ/IQ/OQ/PQ
- Practical Qualification of typical instruments such as
  - Balances
  - HPLC
  - UV spectrophotometer
  - Dissolution apparatus

Sampling
- EU GMP Part 1, Chapters 4, 5, 6
- EU GMP Part 2, Chapter 7 (7.1 – 7.5)
- EU GMP Chapter 4
- Statistical sampling – requirements and interpretation
- EU GMP Annex 8
- EU GMP Annex 19
- US / FDA Requirements
- WHO
- PIC/S
- ISO (former Military Standard)

Documentation in QC Laboratories
- Regulatory requirements (EU / US)
- Specifications, Test Procedures, SOPs, etc.
- Handling of data (paper, electronic, hybrid)
- Laboratory Data Integrity
- Analytical results (Raw data, Raw data check, averaging, rounding of results, …)
- Case Studies
- Laboratory Data Integrity issues related to documentation - issues to be aware of
The Analytical Aspects of Laboratory Data Integrity
- Overview of deficiencies in laboratory data integrity
- Structure of a typical assay in a QC laboratory
- When system suitability requirements are not met
- Is there a system suitability test based on samples?
- When sample variability criteria are not met
- QC samples for method validity check
- Integration of chromatographic peaks: Automated versus Manual
- Reprocessing of raw data and re-integration
- Accepting or invalidating test results?
- Review of audit trail of an analytical run

Lifecycle Approach to Analytical Procedures
- Developing robust, stability indicating methods
- Analytical Target Profile
- Implementation of QbD in development of analytical methods
- Life-cycle of an analytical method
- Are we estimating the real method precision?
- The concept of Assay Format

Handling and Qualification of Primary and Secondary Chromatographic Reference Standards
- Procedure for qualification of a primary reference standard
- Procedure for qualification of a secondary reference standard
- Pharmacopeial standards: handling and re-use
- Will the certified reference standards (CRM) come to the QC lab?
- Assigning purity values to reference standards
- Calculation examples of assigning purity

Managing Out of Specification and Out of Trend Results
- OOS / OOE / OOT
- FDA and MHRA Guidance
- Reportable Value
- Case Study: Practical approach for handling OOS results
- Issues with OOT results and how to manage these

Computer Validation and Integrity of Electronic Data in Analytical Labs
- Regulatory requirements (EU Annex II / US 21 CFR Part II)
- GAMP / GAMP Laboratory Guide
- PIC/S Guide Computerised Systems in GXP Environments
- Validation of Excel spreadsheets
- UCB Case Study: Implementation of a computerised system at UCB from IT infrastructure components to final VSR and periodic review
- Facilitating and ensuring data integrity through validated paperless processes

Presenting and Evaluating Stability Data
- Overview of ICH storage programs for new drugs
- Generic drugs
- Presenting stability data
- Derivation of shelf life according to ICH Q1E

Transfer of Analytical Methods
- Definition and regulatory requirements
- How to perform a method transfer
- Case studies
- Typical and critical issues

Method Comparison – Equivalence Testing of Two Methods
- Is a newly issued pharmacopeial method equivalent to an in-house validated and practiced method?
- Traditional way of comparison
- Equivalence testing with two one-sided t-test (TOST)
- Examples

QA Aspects in QC (relevant for QC Compliance Managers)
- Defining responsibilities for analysts, head of analytical lab, QPs (EU and US)
- Release of APIs, excipients, packaging materials, finished products, etc.
- Contract Labs
- CAPA (Corrective Actions and Preventive Actions)
- Change Control (regulatory framework)
- PQR
- Training (GMP training / training on the job, training records)

Four Workshops

Some of the most important laboratory compliance topics will be further discussed in interactive workshops:

**Topic I: Analytical Instrument Qualification**
Moderator: Dr Thomas Backensfeld

**Topic II: Validation of Analytical Test Procedures**
Moderator: Dr Raphael Bar

**Topic III: Method Transfer**
Moderator: Dr Kerstin Hartisch

**Topic IV: Sampling of Raw Materials, Packaging Components, Devices and Finished Products – Practicing with the Sampling Standard ISO-2859-1**
Moderator: Dr Raphael Bar
Speakers

**Dr Thomas Backensfeld**
Bayer AG, Berlin, Germany
Thomas Backensfeld studied Pharmacy at the University of Muenster and received his PhD in Pharmaceutical Technology at the University of Kiel. He joined Schering AG in 1990 and held different positions in Analytical and Pharmaceutical Development. He has more than 10 year experience in formulation development of solid dosage forms including GMP manufacturing of clinical trial materials. After leading a production plant he returned back to Analytical Development. In his current position as head of an Analytical Development T. Backensfeld is responsible for several analytical project groups developing pharmaceutical products of small molecules. The work covers the development and validation of analytical methods, stability testing, quality control, tech transfer, dossier filing and support of life cycle management.

**Dr Raphael Bar**
BR Consulting, Israel
Raphael Bar is presently a pharmaceutical consultant for the Pharma and bio-Pharma industries. He is consulting various companies and provides development and analytical support to investigational, generic, new drugs as well as combination device-drug products and CMC project management. With a doctorate in Chemistry (1984), Dr Bar joined Teva Pharmaceuticals (1995) and headed for three years the Analytical R&D Laboratory. He was involved in preparation of ANDA files. He then joined Pharmos where he managed the quality control and R&D laboratory till 2007. As Senior Director of Analytical Development he was actively involved in preparation of CMC packages for clinical trial studies. From 2009 until June 2015, he was a member of the scientific advisory board of global PDA (USA).

**Dr Kerstin Hartisch**
Bayer AG, Berlin, Germany
Kerstin Hartisch studied Food Chemistry and Pharmacy at the University of Bonn and received her PhD in Pharmaceutical Analytics. In her position as head of Analytical Development within Bayer, Global Chemical and Pharmaceutical Development she is responsible for all aspects regarding special analytical techniques in product development, i.e. dissolution testing, particle identification, packaging material and excipient testing and also sample and data management (LIMS). She is specialized in the area of dissolution testing including all aspects of automation (Robot Technology).

**Mathieu Materne**
UCB BioPharma sprl, Belgium
IT Compliance Lead, Global TO, NewMeds & Patient Solutions (Medical Device), IT Strategy & Technology Solutions,
Quality, Compliance and IT Professional with over 15 years of experience in BioPharma industry for international Technical and SupplyChain Operations. Strong experiences in GMP/GDP (Engineering, Manufacturing, Laboratory and Supply Chain) and Medical Device projects and data management.

**Sue Mann**
Sue Mann Consultancy, UK
Sue Mann is a Pharmacist and a Qualified Person, and has spent over 35 years in the industry in various roles including technical support, clinical trial supplies and quality assurance/management. She has worked for many different types of company including multinational, contract manufacturing, Japanese and virtual. Sue has worked with both commercial and investigational medicinal products and most major dosage forms. Sue is presently a pharmaceutical consultant working for Pharmaceutical and Biopharmaceutical companies. She has a passion for developing simple, efficient, effective quality systems to support all operational activities including those performed in the analytical laboratory.

**Social Event**
At the end of the first course day you are cordially invited to a social event. This is an excellent opportunity to share your experiences with colleagues from other companies in a relaxed atmosphere.
Use the GMP App at no costs!

The ECA Foundation has developed a GMP App which offers a comprehensive GMP Guideline database with more than 1,400 GMP Guidelines and ten thousands of pages. Check relevant Guidelines (full text versions) during internal audits, regulatory inspections or GMP compliance meetings – simply on your smartphone or tablet PC. In addition to this wealth of guideline information the GMP App also comprises features like GMP News, a comprehensive GMP Search Engine. To open the app just enter app.gmp-compliance.org in your browser and the WebApp opens immediately.

What are The ECA Foundation and the ECA Academy?

The European Compliance Academy Foundation (ECA Foundation) is an independent professional organisation chaired by a Scientific Advisory Board with members from the pharmaceutical industry and regulatory authorities. The ECA Foundation’s goal is to support the Pharmaceutical Industry and Regulators to promote the move towards a harmonised set of GMP and regulatory guidelines by providing information and interpretation of new or updated guidances. The ECA Academy offers professional basic and advanced education (training) programmes. All services offered by the ECA Academy and with regard to ECA Academy Memberships are solely managed by Concept Heidelberg (a leading European training and information services provider). The ECA Foundation is conceptual sponsor of the ECA Academy.

How Do You Become a Member of ECA?

By participating in one of the ECA Academy Conferences or Courses you will automatically become a ECA Academy Individual Member for two years - free of charge. More information about ECA Academy can be obtained on the Website http://www.gmp-compliance.org

What Are the Benefits of ECA?

During the membership, you enjoy
- a EUR 200,- discount on the regular participation fee of any European Conference or Course presented by the ECA Academy.
- free access to the members’ area where you always find the latest update of the “GMP Guideline Manager” online version – allowing you to access a GMP tree with guidelines sorted by topics or by authority. It lets you find relevant guidelines quick and easy.

Lufthansa is Mobility Partner for all ECA Events

As an ECA course or conference attendee, you will receive up to 20% discounted travel fares (according to availability). And as Lufthansa German Airlines offers a comprehensive global route network linking major cities around the world you will most likely be able to benefit from these special prices and conditions.

And this is how it works: Once you registered for a course or conference you will receive a link together with your registration confirmation. Opening that link will take you to the Mobility Partner Program website where you can enter a code in the “Access to Event Booking” area you will also receive. This will take you into an online booking platform* that will automatically calculate the discount offered or provide you with an even better offer if another promotional fare is available.

We look forward to welcoming at one of our next events – and we already wish you a pleasant flight!

*Please note: You may have to enable pop-ups on the Mobility Partner Program website - otherwise the booking platform window will not open.

GMP Certification Programme

This seminar is recognised within the GMP Certification Programme. By attending selected seminars, the participant can acquire an additional certificate. We offer the following modules:

- ECA Certified Validation Manager
- ECA Certified QA Manager
- ECA Certified API Production Manager
- ECA Certified Quality Control Manager
- ECA Certified Technical Operations Manager
- ECA Certified Computer Validation Manager
- ECA Certified Regulatory Affairs Manager
- ECA Certified Microbiological Laboratory Manager
- ECA Certified Sterile Production Manager
- ECA Certified Biotech Manager
- ECA Certified Pharmaceutical Development Manager
- ECA Certified GMP Auditor
- ECA Certified GDP Compliance Manager
- ECA Certified Packaging Manager
- ECA Certified Data Integrity Manager

On the internet at www.gmp-compliance.org you will find a text explaining which seminars are recognised for which certificates.

Or you send an e-mail to info@gmp-compliance.org or a fax to +49-6221- 84 44 64 with the request for information about the GMP Certification Programme. We will then send you our brochure on the topic.
Reservation Form (Please complete in full)

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*Mr.*  
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Company Department

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For questions regarding reservation, hotel, organisation etc.:

Ms Nicole Bach (Organisation Manager) at +49-62 21 / 84 44 22, or per e-mail at bach@concept-heidelberg.de.

General terms and conditions

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1. We are happy to welcome a substitute colleague at any time.
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   - within 1 week prior to the conference 100 %.

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(As of January 2012)

German law shall apply. Court of jurisdiction is Heidelberg.

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