Highlights

- GMP: Where do we come from – where do we go?

- Basic principles of GMP
  - Personnel
  - Hygiene
  - Premises / Production
  - Documentation
  - Risk management
  - Qualification / Validation
  - Communication with clients/authorities

- Elements of a QA System
  - Change Control
  - Deviations
  - CAPA (Corrective Actions – Preventive Actions)
  - Failure Investigations
  - OOS (Out of Specification)
  - Audits – Inspections
  - Falsified products
Objective

The course is designed for people who have no or little knowledge of GMP.
- You get to know the most important pharmaceutical regulations and their importance,
- you get a basic overview of GMP requirements in pharmaceutical production, and
- you become familiar with technical terms from the field of GMP and their meaning

Background

In the manufacture and quality control of medicinal products, compliance with the GMP rules is the decisive aspect for manufacturing high-quality products. For this reason, every staff member in the pharmaceutical industry has to be familiar with the basic GMP requirements. The relevant European GMP regulations define the following prerequisites:

- **Commission directive 2003/94/EC**
  The personnel shall receive initial and ongoing training, the effectiveness of which shall be verified, covering in particular the theory and application of the concept of quality assurance and good manufacturing practice.

- **EudraLex Vol. 4 Good manufacturing practice (GMP) guidelines**
  2.9 Besides the basic training on the theory and practice of Good Manufacturing Practice, newly recruited personnel should receive training appropriate to the duties assigned to them.

In practice, many members of staff are often unaware of the contents and meaning of the different GMP requirements from Europe and US and their consequences for product quality. During this course, speakers with long-standing experience in the training of employees introduce and explain the most important elements of a pharmaceutical GMP system in an easy-to-understand way.

Target Audience

The course is directed to staff from the pharmaceutical industry having no or little experience with the current GMP requirements. This includes both employees who, after their vocational training, enter the pharmaceutical industry and experienced colleagues who work for the first time in a GMP-regulated environment. Participation is also recommended for personnel from suppliers who have to understand the quality requirements of their customers.

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.
Risk Management
- Main topics of ICH Q 9 / Part 3 EU GMP Guideline
- Risk management tools
- Use of risk analysis during qualification
- How to handle FMEA?

Premises / Production
- Requirements for room and equipment
- Classification of rooms
- Sterile production/isolator
- Maintenance of hygiene
- How to behave during production

Qualification/Calibration/Maintenance
- Definitions: Qualification, validation, calibration, maintenance, risk analysis
- Organizing qualification and validation: the validation master plan (VMP)
- Steps in Qualification studies: DQ, IQ, OQ, PQ
- Qualification parameters of typical types of equipment: Clean rooms, water systems, production equipment, analytical equipment
- Performing risk analysis: tools and practical tips
- Calibration: critical types of equipment
- How to build up a calibration system
- Maintenance: Requirements and system
- Validation of computerised systems

Process Validation, Computer Validation and Validation of Analytical Methods
- General aspects and requirements
- Process Validation
- Documentation of process validation
- Validation of analytical methods
- Documentation of analytical methods validation

Cleaning Validation
- Regulators requirements
- The cleaning procedure
- Building up a cleaning validation
- Sampling
- Analytical tests

Audits and Inspections
- Types of audits
- Requirements
- Dos and don’ts for the auditee - How to survive audits?
- Performing audits and self-inspections
- Good audit practices

Packaging/Storage/Transportation
- Packaging/Storage/Transportation in the regulations
- Managing of packaging process
- What is necessary to regulate in a pharmaceutical company
- WHO good storage practice – elements and requirements
- Transportation as part of storage
- How to maintain the quality during transportation

Falsified Products
- Regulatory requirements
- Responsibilities of QP and the pharmaceutical industry
- What measurements can be taken
- Strategies against falsified products

Speakers

Dr Bettina Pahlen, Quality x Pharma Consulting GmbH, Germany
Bettina Pahlen, PhD, studied pharmacy at the University of Muenster, Germany, graduated in pharmaceutical chemistry and performed post-docs in USA and Germany. During the last 15 years she worked at university, authority and in different areas of the pharmaceutical industry (R&D, manufacturing, quality control, quality assurance). Since July 2007, she has been working as a consultant in the pharmaceutical industry focussing on GxP Quality Assurance aspects.

Dr Heinrich Prinz
PDM Consulting, Germany
Dr Prinz worked with Boehringer Mannheim before he joined Biotope where he was Head of Quality Assurance, responsible for both the pharmaceutical and the medical device division. Since 2003 he works as a freelance consultant.

Dr Wolfgang Schumacher, formerly F. Hoffmann-La Roche Ltd., Switzerland
Dr Schumacher studied chemistry and pharmacy. After entering Asta Medica, he headed different positions. From 2001 to 2016 he was Head of the department of Quality Computer Systems at F. Hoffmann-La Roche, Baar. He is a member of the ECA Advisory Board.

Participants’ comments of March 2018 course:
“In total very helpful and well presented/explained lectures. Thank you.” Katharina Bubb, Janssen Vaccines AG, Switzerland

“Highly informative. Worthy & completed answers received for questions – highly open discussion atmosphere. Highly informative for getting the overview of GMP Basics.” Laia Camprubi Gallet, Bayer Bitterfeld GmbH
If you cannot attend the conference you have two options:
1. We are happy to welcome a substitute colleague at any time.
2. If you have to cancel entirely we must charge the following processing fees:
   - Cancellation until 2 weeks prior to the conference 10%,
   - Cancellation until 1 week prior to the conference 50%,
   - Cancellation within 1 week prior to the conference 100%.

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Terms of payment: Payable without deductions within 10 days after receipt of invoice.

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In case you do not appear at the event without having informed us, you will have to pay the full registration fee, even if you have not made the payment yet. Only after we have received your payment, you are entitled to participate in the conference (receipt of payment will not be confirmed)! (As of January 2012).

German law shall apply. Court of jurisdiction is Heidelberg.

Privacy Policy: By registering for this event, I accept the processing of my Personal Data. Concept Heidelberg will use my data for the processing of this order, for which I hereby declare to agree that my personal data is stored and processed. Concept Heidelberg will only send me information in relation with this order or similar ones. My personal data will not be disclosed to third parties (see also the privacy policy at http://www.gmp-compliance.org/eca_privacy.html). I note that I can ask for the modification, correction or deletion of my data at any time via the contact form on this website.

CONCEPT HEIDELBERG
P.O. Box 101764
Fax +49 (0) 62 21/84 44 34
D-69007 Heidelberg
GERMANY

Reserve Form (Please complete in full)

GMP for Beginners
☐ 24/25 March 2020, Hamburg, Germany
☐ 20/21 October 2020, Berlin, Germany

Title, first name, surname

Department

Company

Important: Please indicate your company's VAT ID Number

Purchase Order Number, if applicable

City

ZIP Code

Country

Phone / Fax

E-Mail (Please fill in)

Fees (per delegate, plus VAT)

ECA Members € 1,490
APIC Members € 1,590
Non-ECA Members € 1,690
EU GMP Inspectors € 845

The conference fee is payable in advance after receipt of invoice and includes conference documentation, dinner on the first day, lunch on both days, and all refreshments. VAT is reclaimable. Accommodation

CONCEPT HEIDELBERG has reserved a limited number of rooms in the conference hotel. You will receive a room reservation form/POG when you have registered for the course. Reservation/POG should be made directly with the hotel. Early reservation is recommended.

Date & Venue March 2020

Tuesday, 24 March 2020, 09.00 h – 18.00 h
(Wednesday, 25 March 2020, 08.30 h – 17.00 h)
Barcelo Hamburg
Ferdinandstr. 15
20095 Hamburg, Germany
Phone +49(0)40 22 63 62 0
berlin@barcelo.com

Date & Venue October 2020

Tuesday, 20 October 2020, 09.00 h – 18.00 h
(Wednesday, 21 October 2020, 08.30 h – 17.00 h)
InterCityHotel Berlin Hauptbahnhof
Katharina-Paulus-Straße 5
10557 Berlin, Germany
Phone +49(0)30 288 755 0
berlin-hauptbahnhof@intercityhotel.de

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