Learn How to Plan, Implement and Document Effectively Computer Validation Activities

**SPEAKERS:**

- **Frank Behnisch**  
  CSL Behring GmbH, Germany

- **Dr David Selby**  
  Selby Hope International, UK

- **Dr Robert Stephenson**  
  Rob Stephenson Consultancy, UK

**LEARNING OBJECTIVES:**

- The New EU GMP Guide Annex 11
- 21 CFR Part 11
- The GAMP® 5 Lifecycle
- Practical Risk Management – ICH Q9 and FMEA Methodology
- Validation Planning
- Change Control
- Validation Documentation
- Presentation to Inspectors
- **Learning by doing:** up to 10 Workshops

**GAMP®** is a trademark of ISPE - http://www.ispe.org/gamp5

This education course is recognised for the ECA GMP Certification Programme „Certified Computer Validation Manager“. Please find details at www.gmp-certification.eu
**Learning Goals**

- You get to know the current risk management approaches of ICH Q9 and GAMP®5
- You become familiar with the latest methods and tools for risk analysis and can assess their relevance to practice in the validation of computerised systems
- You learn how the activities involved in the validation of computerised systems can be controlled efficiently by means of risk management
- In 4 workshops you can apply the procedures and discuss them

**Background**

The current GMP regulations and guidelines (ICH Q9, GAMP®5, EU GMP Guide Annex 11 "Computerised Systems") focus more and more on the topic of risk management. However, the regulations do not offer much concrete advice on how its principles should be translated into practice during the validation and operation of computerised systems. Therefore, it is the aim of this course to provide you with practice-oriented guidance in performing this task.

**Target Group**

This Education Course is directed at employees from Production, Quality Control / Quality Assurance, Engineering, IT who have to deal with risk assessment and risk management in the field of computer validation.

**Programme**

**Introduction - What do you want from this day?**
- Capturing delegates expectations
- Sharing and reducing to key points in groups
- Sharing with all delegates and tutors

**An Introduction to Risk Management (including ICH Q9)**
- Definition of "Quality Risk Management"
- Principles of Quality Risk Management
- Application of the principles in validation
- Methods of assessing and controlling risk
- Regulatory expectations for risk management

**Risk Management the GAMP® 5 Way**
- The GAMP methodology for risk management
- Where to apply risk management in validation
- Methods of assessing risk

**Workshop 1: Risk Assessment in Validation**
- Risk management applied to a computer system
  - Evaluating identified risks
  - Classification of risks into H, M, L
  - Controls to mitigate unacceptable risks
  - Links to the validation plan and protocols

**Workshop 2: Risk Management in Validation**
- Risk management applied to a control system
  - What are the conclusions from the risk assessment?
  - What options do you have to mitigate (reduce) the higher risks?
  - How will the output affect the protocol?

**Workshop 3: Applying Risk Ranking to determine periodic review priorities**
- What factors influence supplier assessment?
- What risks are associated with supplier selection?

**Workshop 4: Assessing and Selecting a supplier**
- What factors influence supplier assessment?
- What risks are associated with supplier selection?

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Learning Goals

This is why you should attend this course:

- You will systematically be introduced to the principles and methods of the validation of computerised systems (according to GAMP®)
- You will learn the skills to plan, implement and document effectively validation activities for computerised systems and to assess them with respect to their GxP compliance
- You have the opportunity to practically apply the theoretical foundations in 6 workshops

Background

Computerised systems are a central factor determining work sequences in the pharmaceutical industry. Their use increases product safety and saves time and costs of manual intervention. This creates the requirement and necessity, however, to validate all computerised systems which can influence the quality of pharmaceutical products. The basis of the education course will be the current requirements for the validation of computerised systems like GAMP® and their GxP-oriented application in practice. Experts from the pharmaceutical industry and from the GAMP® Committee will show you efficient ways to validate your computerised systems.

Target Group

This course is directed towards specialists and executives in the pharmaceutical industry entrusted with the planning, implementation and evaluation of the validation of computerised systems.

Programme 1st Day

Introduction – What the Participants Expect
An open session capturing the expectations of the delegates

Laws, Regulations and Guidelines for Computer Validation
- The historical perspective
- Current regulations and regulatory guidelines from US
- New regulatory guidance (GAMP® 5, GAMP® Good Practice Guides, ASTM)
- New industry guidance
- Regulatory training
- Harmonisation

Electronic Records and Signatures
- What Part 11 means – Now!
- Identify e-records in predicate rules
- Identify risks to records
- Identify appropriate controls for records

The EU Annex 11 “Computerised Systems”
- What is new?
- What are the important points?
- How can you implement it?

Workshop 1: Self Evaluation of Compliance with Annex 11

The GAMP® 5 Approach to Computer Validation
- Validation needs structure
- The GAMP® approach
- What is new in GAMP® 5
- General validation activities
- The GAMP® Categorisation System
- Life Cycle cost reduction

Workshop 2: Review of User Requirements Specifications
A short review of the URS and how to write specifications, as a prelude to a workshop in which delegates will evaluate a real requirements specification.
- What is a URS?
- Why is it important?
- Contents of a URS
- Characteristics of good specifications
- Testable specifications

Risk Management – A Key Point Review
- How to use the FMEA tool to reach better risk-based decisions?
- Risk registers
- Documenting risk
- How to make use of risk information
Programme 2nd Day

Validation Planning
- Why is a validation plan important?
- Definitions and regulatory expectations
- Building risk management into planning phase
- Structure and contents of validation plans
- Discussion of best approach
- The impact of scaleability

Workshop 3:
Validation Planning
Based on considerations of the type of application, knowledge of the supplier and how it will be used, delegates will work out the best approach to delivering the benefits of a GxP system
- What are the risks associated with delivering the system?
- What options do you have to manage the most critical risks?
- How can they be best managed?
- What are the key issues to monitor to ensure delivery of the project benefits?

Specifications, Design Review and Traceability
- What sorts of specifications are needed?
- How are they constructed?
- Can they be combined?
- How to carry out a design review?
- How to construct a traceability matrix?

Protocols, Test Scripts and Deviation Management
- Principles of Risk-Based Qualification
- Leveraging the Supplier
- Commissioning vs Qualification
- Test Script Design
- Deviation Management

Workshop 4:
Risk Management in Protocol Planning
Based on a real case study, delegates will use the same risk assessment techniques as in Workshop 2 to determine where to focus the qualification of a packaging line.
- Risk management applied to a control system
- Using FMEA to assess risks to be managed and controlled in validation
- Identifying options to mitigate (reduce) the higher risks
- Using the output in creating the testing protocol

Change Control
- Regulatory requirements
- Configuration management
- Responsibilities
- Planned/unplanned changes
- Classification
- Sources of changes

Workshop 5:
Change Control
The participants will work on a number of case studies and define the change control activities needed.
- Change Control forms
- Approval process
- Standard Changes
- Committees

Workshop 6:
Managing Deviations
In this workshop examples of deviations will be examined and methods of resolution discussed. The examples are based on real-life protocols.
- Test failures found during IQ/OQ
- Manage the deviations
- Suggest solutions

Programme 3rd Day

Automation Aspects
- System Overview
- GAMP® and risk analysis
- Specifications
- Qualification / Validation

Validation Reporting & Presentation to Inspectors
- The link between the plan and the report
- Key documents
- Validation summary reports
- Style and emphasis
- Managing the inspection

Introduction to IT-Infrastructure Qualification
- The qualification lifecycle
- How to deal with user requirements
- Qualification documentation
- Critical issues
- Qualification summary report

Regulatory Comments
- Recent general trends
- Highlights from Warning Letters and 483s
- Lessons we must learn
Accommodation

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

Registration

Via the attached reservation form, by e-mail or by fax message. Or you register online at www.gmp-compliance.org.

Conference Language

The official conference language will be English.

Organisation and Contact

ECA has entrusted Concept Heidelberg with the organisation of this event.

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For questions regarding reservation, hotel, organisation etc.: Ms Marion Grimm (Organisation Manager) at +49-(0)62 21 / 84 44 18 or per e-mail at grimm@concept-heidelberg.de.

Social Event

On 18 November 2015 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.

Speakers

Frank Behnisch
CSL Behring GmbH, Germany
Frank is Senior Manager Project Engineering at CSL Behring GmbH in Marburg, Germany. He is member of the GAMP® D-A-CH „steering committee“ and chairman of a GAMP® Special Interest Group (SIP) for “Small Systems”.

Dr David Selby
Selby Hope International, UK
David Selby, BSc., PhD., was with Glaxo for many years in different positions. He occupied the role of Site Quality Assurance Manager there and latterly, he was the Site Manager. He is a founder member and Chairman of the GAMP® Forum and 2004 Chairman on the International Board of ISPE. He has established his own consultancy, Selby Hope International, specialising in the compliance of computerised systems and automated equipment used in pharmaceutical manufacturing.

Dr Robert Stephenson
Rob Stephenson Consultancy, UK
Rob has had extensive experience with the implementation and operational control of a wide range of applications within the Pharmaceutical and Personal Products sector. He joined Pfizer Sandwich UK in 2000 as member of their Quality Unit operating within the IT group where his responsibilities included coordinating the manufacturing site’s initiative to achieve 21 CFR Part 11 compliance and authoring their IT Quality Management System. As a long-standing member of the GAMP® Forum Rob has contributed material to GAMP®5 and the ISPE GAMP Good Practice Guide on “A Risk-Based Approach to Operation of GxP Computerized Systems” for which he was co-leader. Rob now works as an independent IT Systems Validation Consultant.

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.
Reservation Form (Please complete in full)

☐ Computer Validation: Introduction to Risk Management,
17 November 2015, Copenhagen, Denmark
☐ Computer Validation – The GAMP® 5 Approach,
18-20 November 2015, Copenhagen, Denmark
☐ Mr. ☐ Ms.

Title, first name, surname

Company
Department

Important: Please indicate your company's VAT ID Number
P.O. Number if applicable

Street/P.O. Box
City Zip Code Country

Phone/Fax
E-Mail (please fill in)

Date and Venue
Computer Validation:
Introduction to Risk Management
Tuesday, 17 November 2015,
09.00 h – 18.15 h
(Registration and coffee
08.30 h - 09.00 h)

Computer Validation –
The GAMP® 5 Approach
Wednesday, 18 November 2015,
09.00 h – 17.30 h
(Registration and coffee
08.30 h - 09.00 h)
Thursday, 19 November 2015,
09.00 h – 17.30 h
Friday, 20 November 2015,
08.30 h – 13.00 h

Radisson Blu Scandinavia Hotel
Amager Boulevard 70
2300 Copenhagen S, Denmark
Phone +45 33 96 50 00
Fax +45 33 96 55 00

Fees (per delegate plus VAT)

Computer Validation: Introduction to Risk Management
ECA Members € 790
APIC Members € 840
Non-ECA Members € 890
EU GMP Inspectorates € 445

The conference fee is payable in advance after receipt of invoice and includes conference documentation, lunch and all refreshments. VAT is reclaimable.

Computer Validation – The GAMP® 5 Approach
ECA Members € 1,790
APIC Members € 1,890
Non-ECA Members € 1,990
EU GMP Inspectorates € 995

The conference fee is payable in advance after receipt of invoice and includes conference documentation, lunch, two lunches and all refreshments. VAT is reclaimable.

Save Money and book both courses
ECA Members € 2,190.
APIC Members € 2,290.
Non-ECA Members € 2,390.

General terms and conditions
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:
   - until 2 weeks prior to the conference 10 %,
   - until 1 weeks prior to the conference 50 %,
   - within 1 week prior to the conference 100 %.
   CONCEPT HEIDELBERG reserves the right to change the materials, instructors, or speakers without notice or to cancel an event. If the event must be cancelled, registrants will be notified as soon as possible and will receive a full refund of fees paid.

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

Important: This is a binding registration and above fees are due in case of cancellation or non-appearance. If you cannot take part, you have to inform us in writing. The cancellation fee will then be calculated according to the point of time at which we receive your message. 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)

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.