



## Speakers



**Frank Behnisch**  
CSL Behring GmbH



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Körber Pharma Consulting



**Yves Samson**  
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**Dr Robert Stephenson**  
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## Computerised System Validation:

- Leveraging Suppliers
- Computerised System Validation Master Class



Live Online Training on 4 & 5 - 7 May 2021



*Qualify yourself as an expert for the validation of computerised systems*

## Highlights

- Regulatory Update
- Leveraging Suppliers
  - Managing Quality
  - Leveraging Test Activities
  - Supplier Assessment
- Good Validation Practices
- System Classification
- Scalability of Validation
- Advanced Risk Management
- IT Governance
- Data Integrity
- Alternative & Agile Approaches
- Today / Future IT Compliance Challenges
- Learning by doing: up to 9 Case Studies
- Interactive sessions



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## Objective

- Learn what activities and deliverables you should expect to see from your IS/IT supplier to demonstrate Supplier Good Practice
- Learn how to verify your supplier’s capabilities so that there are “no surprises”.
- Learn how to plan verification and validation activities, leveraging the expertise of your supplier
- Learn how to minimise duplication of effort between the supplier and your regulated company in order to achieve lean and effective processes throughout the system life cycle
- Learn how to work with your supplier to build a strong and lasting client-supplier relationship

## Background

Recognising the potential savings and flexibility available, regulated companies are increasingly withdrawing from ‘in-house’ developed solutions and looking to their external suppliers to provide them with innovative and compliant products and services which fulfil their operational and business needs.

The EU-GMP Annex 11 on Computerised Systems states that ‘the competence and reliability of a supplier are key factors when selecting a product or service provider’; ‘Leveraging Supplier Involvement’ is also one of the 5 key concepts of the GAMP®5 guidance ‘A Risk-Based Approach to Compliant GxP Computerized Systems’.

This Live Online Trainings aims to provide attendees with the knowledge, and opportunities to practice the skills required, to achieve successful partnerships with their IS/IT suppliers and to improve the efficiency and effectiveness of their validation processes.

## Target Audience

This ECA Live Online Training is directed at employees from Production, Quality Control/Quality Assurance, Engineering and IS/IT, who have to assess, manage or work with computerised system or service providers.

The Live Online Training will also be of value to representatives from supplier organisations that are working or seeking to work with Regulated Companies in the Life Sciences Sector.

## Programme

### Introduction – What the Participants Expect

An open session capturing the expectations of the delegates.

### Leveraging Supplier Expertise: An Overview of Good Practice

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- What is current Good Practice?
- Optimising Supplier involvement
- Integrating the supplier’s expertise and deliverables into your validation process
- How to do more with less

### Performing a Supplier Assessment

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- Why Assess the Supplier?
- The Overall Process
- Assessment Topics
- Types of Assessment
- Corrective Actions & Follow Up Audits



#### Case Study: Selecting a Supplier

- What factors to consider?
- How to focus the assessment?
- How to engage with the supplier?
- How to report and manage the findings?
- Regulatory expectations



#### Case Study: Quality Planning within a Supplier’s QMS - Developing a Quality Plan that Delivers

- Quality Management System
- Establishing Requirements
- Producing Specifications
- Testing and Release
- Support and Maintenance

### Identifying Leveraging Opportunities

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- Defining the role of the supplier
- What must the supplier do?
- What must the regulated company do?



#### Case Study: Leveraging Supplier Testing

- Test script development
- Test script execution
- Test script review and approval

### Managing Quality within an Outsourced IS/IT Environment

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- Making a Business Case
- Outsourced Supplier Specification and Selection
- Implementation
- Monitoring
- Contract Change and Exit

## Objectives

As a specialist in the validation of computerised systems, this event will provide you with

- Suggestions on how current regulatory guidance on computerised systems relating to data integrity, critical thinking and CSA (Computer Software Assurance) can be put into practice
- Real-life examples of how validation effort can be scaled according to risk-based approaches
- Answers to specific questions, e.g. on source code review or on creating specification documents
- The opportunity to bring questions from your own practice up for discussion

## Background

The V-model has become a standard worldwide methodology for the validation of computerised systems. Regulatory requirements, as well as industry guidelines, like GAMP®5, are orientated towards this model. In practice, you as a validation specialist will want to know how to apply this model to current and increasingly complex validation projects.

## Target Audience

The Master Class is directed at employees from

- IT
- Production
- Engineering
- Quality Assurance
- Quality Control

Participants should already have gained experience in the validation of computerised systems, and preferably will have also attended a basic CSV Course.

## Programme

Introduction – Gain Understanding of Delegate Experience and Background



### Case Study: What the Delegates expect

- Capturing delegates expectations
- Sharing and reducing to key points
- Facilitated discussion

### Current Challenges and Evolution for CSV Activities

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- What does compliance really mean?
- Data Integrity
- Securing operation: cybersecurity
- Project agility
- Cost efficiency vs effective risk management
- Applying critical thinking

### Roles, Responsibilities and Governance

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- PQS – Pharmaceutical Quality System according to ICH Q10
- Responsibilities
  - Operational ownerships
  - Supporting roles
- QA oversight



### Case Study: Governance Benchmark

- Polling Exercise plus facilitated discussion
- IT and System Governance
- CSV Roles and Responsibilities
- Role of Quality Unit

### Practical Use of Scaleability

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- What do we mean by Scaleability?
- How does it work in practice?
- How can we combine documents successfully?
- How much is enough?



### Case Studies: Scaleability of Validation Activities

- LIMS – Laboratory Information Management System
- Laboratory computerised equipment
- Process control system: PLC – Programmable Logic Controller

### Writing Requirements Documents

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- Requirements gathering
- Writing good requirements
- Use of templates / boilerplates
- Requirement Quality

### Requirements: The Good, the Bad and the Ugly

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- Good, bad, and could-be-better requirements
- Testability
- Use of templates / boilerplates
- What you should never write

### Ideal Content of a CSV SOP

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- Embedding the CSV SOP into the PQS
- Topics to address

### Data Integrity and Record Management: A Necessary Long-Term Approach

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- Regulatory context
- Document life cycle
- Retention requirements and constraints
- Supporting processes
- Areas of concern

## System Classification – A Record-based Approach

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- Needs for record-based system classification
- Classification criteria
- Class A, B, C, D

## Design Review – How to Apply Critical Thinking?

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- CSA – Computer Software Assurance
- Scaleable Risk Management
- Document Review



### Case Study: Design Review Scaleability

- Combining Risk Management & Design Review

## Bringing Legacy Systems into Compliance

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- How to approach legacy system remediation
- Examples
  - Learning management system
  - Laboratory Instruments



### Interactive Session: Good Validation Practices

*Open session in which delegates score their CSV approach against 12 good validation practices*

- Each good practice introduced
- Delegates score themselves
- Results consolidated and fed back
- Allows delegates to compare their CSV system against best practice and other practitioners

## Alternative & Agile Approaches

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- Alternative software development models
  - Unified Process, Scrum
- Agility objectives
  - Need for flexible engineering methodologies
- What Agile engineering is not
  - What Agile engineering needs
- Practical approaches and recommendation
  - Conditions for success

## Validating Spreadsheets

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- Why are spreadsheets high risks?
- Design considerations
- What is important (risk again!)
- How to document spreadsheet validation



### Case Studies: Complex Projects

- Global projects
  - Roles & Responsibilities
  - Data-related requirements
- Large systems
  - Phase-based implementation and deployment
- Interface projects
  - Roles & Responsibilities
  - Testing

## Today / Future IT Compliance Challenges

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- Open Source Software validation
- Challenge demands Infrastructure platforms for applications
- Global systems validation vs local defence
- Paperless recipes based production – ISA 95 / S 88
- Cloud Computing – Data Integrity
- Validating Artificial Intelligence (AI)
- Challenges for data integrity on Lab-Systems

## Elaboration of a Data Integrity Programme

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- Data Integrity Programme: What to do?
  - Topics to address
  - Action planning
- Embedding the Data Integrity Programme into the PQS
- Progress Reporting

## Code Review

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- Principles of code review
- Regulatory expectations
- Performing code reviews
- How to document code reviews

## Real Life Qualification / Validation Projects

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- Real Life Qualification / Validation Projects
- Requirements
- Constraints
- Issues identified
- Lessons learned

## This could be of interest for you as well

**Would you like to train a larger group of participants in your company?** We offer practice-oriented GMP/GDP training courses on:

- Basic GMP:  
APIs (ICH Q7), Medicinal Products, Biopharmaceuticals
- Quality Assurance
- Quality Control
- Validation/Qualification
- Regulatory Affairs
- Sterile Manufacturing
- IT / Computer Validation
- Good Distribution Practice (GDP)
- Data Integrity
- Packaging
- Medical Devices
- Technical Operations

You will find a time schedule for each training course at <https://www.gmp-compliance.org/training/gmp-gdp-in-house-trainings>.

### Why not online? GMP/GDP seminars, webinars and e-learning

Take advantage of the wide range of „on demand“ training opportunities offered by the ECA Academy. You can use various online offers at any time without software installation. There is an extensive selection of courses available. Simply book online - with a certificate of completion, of course. Find out more at <https://www.gmp-elearning.com> and <https://www.gmp-compliance.org/gmp-webinars/recorded-gmp-webinars>

## Ordering a Recording

Independent from the Live Online Training, you can also order a recording of this training at the same conditions. This recording will be provided on our media server. All you need to watch it is an Internet browser – no additional software. You can order the recording of the Live Online Training at the earliest 10 days after the live performance at <https://www.gmp-compliance.org/on-demand-online-training/recorded-online-training-webinars>.

## Organisation and Contact

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

CONCEPT HEIDELBERG  
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69007 Heidelberg, Germany  
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Ms Marion Grimm (Organisation Manager) at  
+49(0)62 21 /84 44 18, or per e-mail at  
grimm@concept-heidelberg.de.

## Your Benefits

### Internationally Acknowledged Certificate from ECA Academy

The EU GMP Guide requires: „... All personnel should be aware of the principles of Good Manufacturing Practice that affect them and receive initial and continuing training...“ This is why you receive an acknowledged participant certificate, which lists the contents of the Live Online Training in detail and with which you document your training.



### Computerised System Validation Master Class is recognized for the GMP/GDP Certification Scheme

Building on your education the ECA GMP/GDP certification programmes provide you with the appropriate supplement to acquire this qualification. The the Live Online Training

Computerised System Validation Master Class is the first element for your additional certification. Simply choose any three courses within the programme according to your professional interest. Your certificate is then valid for two years. To renew it, you can pick any training from the ECA courses and conferences list within that two-years period – allowing you to broaden your knowledge in GMP and GDP compliance. Please find more information at [www.gmp-certification.org](http://www.gmp-certification.org)

## 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 (SIG) for “Small Systems”.



Stefan Münch  
Körber Pharma Consulting / Systec Services GmbH, Karlsruhe, Germany

Stefan Münch, Business Director Validation, is responsible for all validation services of Körber Pharma Consulting. He has more than 20 years of experience in software development for the pharmaceutical industry (MES). Furthermore, Mr. Münch is actively engaged in GAMP D-A-CH for many years and member of the steering committee.



Yves Samson, Kereon AG  
Basel, Switzerland

Yves joined the industry where he served as project and site engineer automation. In 2002, he founded Kereon AG. He is member of GAMP® Europe Steering Committee, co-founder and chairman of GAMP® Francophone and edited the French version of GAMP® 4 / 5. Membership: Active member of the GAMP working group ‘IT Infrastructure Compliance and Control’ / ECA ‘DI & IT Compliance Group’.



Dr Robert Stephenson  
Rob Stephenson Consultancy, UK

Rob has had more than 30 year experience in Pharmaceutical and Personal products industries (Boots, Lilly, Unilever, Pfizer). As a long-standing member of the GAMP Europe Steering Committee 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.



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Reservation Form (Please complete in full)



## Live Online Training

- Computerised System Validation: Leveraging Suppliers, 4 May 2021
- Computerised System Validation Master Class, 5-7 May 2021

Title, first name, surname

Department

Company

Important: Please indicate your company's VAT ID Number

Purchase Order Number, if applicable

City

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Country

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E-Mail (Please fill in)

### 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:
  - 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 %

CONCEPT HEIDELBERG reserves the right to change the materials, instructors,

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If the event must be cancelled, registrants will be notified as soon as possible and will receive a full refund of fees paid. CONCEPT HEIDELBERG will not be responsible for discount, airfare penalties or other costs incurred due to a cancellation.

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 can-

cellation 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).

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](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.



## Dates of the Live Online Training

### Computerised System Validation: Leveraging Suppliers

Tuesday, 4 May 2021, 09.00 h – 18.00 h

### Computerised System Validation Master Class

Wednesday, 5 May 2021, 09.00 h – 17.30 h

Thursday, 6 May 2021, 09.00 h – 17.30 h

Friday, 7 May 2021, 09.00 h – 13.30 h

All times mentioned are CEST.

## Technical Requirements

For our webinars, we use Cisco WebEx, one of the leading suppliers of online meetings. At <https://www.webex.com/test-meeting.html> you can check if your system meets the necessary requirements for the participation at a WebEx meeting and at the same time install the necessary plug-in. Please just enter your name and email address for the test. If the installation is not possible because of your rights for the computer system, please contact your IT department. WebEx is a standard nowadays and the necessary installation is fast and easy.

## Fees (per delegate plus VAT)

### Computerised System Validation: Leveraging Suppliers

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.

### Computerised System Validation Master Class

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.



## Save money and book both courses:

ECA Members € 2,390

APIC Members € 2,490

Non-ECA Members € 2,590

EU GMP Inspectorates € 1,440

The conference fee is payable in advance after receipt of invoice.

## Registration

Via the attached reservation form, by e-mail or by fax message. Or you register online at [www.gmp-compliance.org](http://www.gmp-compliance.org)

## Presentations/Certificate

The presentations will be made available to you prior to the Live Online Training as PDF files. After the event, you will automatically receive your certificate of participation.

## Conference language

The official conference language will be English.