

## Speakers



Dr Hiltrud Horn  
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# How to provide Process Validation data in a regulatory submission

04 - 05 November 2020 | Hamburg, Germany



## Highlights

- Dossier requirements for description of the manufacturing process validation
- How to provide manufacturing process data in an NDA – FDA requirements
- Key aspects of traditional process validation and continuous process verification with regard to regulatory submissions
- Providing stability data in regulatory submissions
- Process validation and GMP issues
- How to handle post-approval changes
- Process validation of biotech-derived APIs

## Objective

This education course focuses on how to compile and provide information and data from Process Validations for Drug Substances and Drug Products both of chemical and biotechnological origin. You will learn

- How to prepare and process the data derived from validation runs of drug product manufacturing processes
- What needs to be documented about drug substance manufacturing processes
- How to manage and document post approval changes in manufacturing processes
- What to consider for compiling stability data for the dossier
- How to provide validation data of biotech manufacturing processes

## Background

Process Validation can be defined as documented evidence that the process, operated within established parameters, can perform effectively and reproducibly to produce a medicinal product meeting its predetermined specifications and quality attributes. In general there are two options to validate a manufacturing process: a traditional approach or an enhanced approach where continuous process verification is applied. Irrespective of which approach is used the manufacturing process should be validated before the product is placed on the market. Therefore complete data have to be provided in the dossier at the time of regulatory submission. These data should cover the validation for all manufactured strengths, batch sizes, pack sizes and proposed manufacturing sites.

Guidance on process validation information to be provided in regulatory submissions is given in 2 EMA Guidelines: “*Guideline on process validation for finished products – information and data to be provided in regulatory submissions*” and “*Guideline on process validation for the manufacture of biotechnology-derived active substances and data to be provided in the regulatory submission*”.

Provisions of both GMP compliant manufacture and dossier requirements are laid down in FDA’s Guidance for Industry entitled “*Process Validation: General Principles and Practices*”.

## Target Audience

This education course is designed for all persons involved in the compilation of dossiers for regulatory submissions who want to become familiar with the requirements for the documentation of Process Validation data. The course will be of interest in particular for personnel from Regulatory Affairs as well as for personnel from Quality Assurance, Production and Quality Control.

## Programme

### Process validation of manufacturing processes – Dossier requirements in the EU

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- Relevant guidance documents
- Finished product process validation
- Traditional and enhanced approaches
- Process validation schemes
- Standard vs. non-standard processes

### Traditional Process Validation and Continuous Process Verification

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- What are the opportunities and challenges?
- What are the key-aspects for the CTD?
- What should you consider for selecting the right validation strategy?
- What should you know about design space?
- What are typical validation questions to be addressed?

### How to provide stability data in regulatory submissions

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- Stability data from drug substances and drug products in the CTD
- Long-term and accelerated conditions, in-use stability
- Requirements for the different climatic zones
- Stability summary and conclusion
- Process parameters with potential impact on drug substance/drug product stability
- Changes in the process: what has to be considered regarding stability?

### Manufacture of active substances – Process validation and GMP issues

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- API manufacture – What needs to be documented in the dossier?
- Process validation for APIs – Key aspects
- GMP for APIs

### How to provide data from API manufacturing process validation – case study

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### Case Studies on typical Validation Projects

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- Standard and non-standard processes
- Validation approach for drug substances and drug products
- Validation strategy and planning from development to registration
- Specific points to be considered for EU and US

## Basic requirements and expectations of the FDA regarding process validation

- Approach to and considerations for process validation
- Process Design
- Process qualification and process performance qualification (PPQ)
- The PPQ protocol – execution and report

## Handling Post-Approval Changes in manufacturing processes

- Which GMP and regulatory aspects need to be considered (e.g. site /process changes)?
- How to define the validation strategy?
- What are the challenges?
- How to be successful?

## Process validation for the manufacture of biotech-derived APIs – process evaluation and verification

- Process evaluation
- Critical quality attributes (CQAs) of the active substance
- Small scale models
- Process verification studies and data
- Number of batches to be presented
- Design space option
- Evaluation of the upstream process
- Criticality assignment of process parameters
- Potential impact of raw materials
- Verification of upstream process
- Single use equipment
- Evaluation and verification of downstream process
- Comparability of products manufactured in different sites



### GMP/GDP Certification Scheme

Building on your education, the ECA GMP/GDP certification programmes provide you with the appropriate supplement to acquire this qualification. 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. More details at:

[www.gmp-compliance.org/gmp-gdp-certification-programme](http://www.gmp-compliance.org/gmp-gdp-certification-programme)

## Speakers



**Dr Hiltrud Horn**  
Horn Pharmaceutical Consulting, Germany

Dr Horn is managing director of HORN PHARMACEUTICAL CONSULTING with focus on CMC, GMP and Regulatory Affairs. She started in pharma industry in 1990 and held several managerial positions within Hoffmann-La Roche in Basel and Knoll (Abbott) with global responsibility within QC / QA / Regulatory Affairs / Project Management / Medical Writing.



**Dr Nils Jost**  
Gründau, Germany

Dr Jost is an expert for the assessment of CMC dossiers for clinical trial applications, EMA centralized marketing authorizations and national marketing authorizations for biological medical products. He studied biology at the Ruhr-University Bochum and the University of Essen-Duisburg.



**Dr Wilhelm Schlumbohm**  
Berlin, Germany

Dr Schlumbohm worked 30 years with German drug licensing authorities. He was an expert for the Certification Procedure of the European Pharmacopoeia and a member of the TAB for several years. He was also a member of the ASMF working group, and the CVMP co-opted member for quality. He is a pharmacist, holds a Ph D in biochemistry, and is further qualified as pharmacist for drug information and for public health.



**Dr Norbert Skuballa**  
Biologische Arzneimittel Heel, Germany

Dr Skuballa is head of the Pharmaceutical Compliance Management function at Heel and responsible for development and coordination of all compliance related GxP and regulatory affairs processes. He has been working in the pharmaceutical industry since 1991, mainly for Schering (now Bayer Pharmaceuticals) in Research, Production and Quality Management.



### Social Event

In the evening 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.

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Reservation Form (Please complete in full)

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Title, first name, surname

Department

Company

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Purchase Order Number, if applicable

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GERMANY

### 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 100 %.

- Cancellation until 1 week prior to the conference 50 %.

- Cancellation within 1 week prior to the conference 100 %.

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to pay the full registration fee, even if you have not made the payment yet. Only

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ference (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.

## Date

Wednesday, 04 November 2020, 9.00 – 18.00 h  
(Registration and coffee 8.30 – 9.00 h)

Thursday, 05 November 2020, 9.00 – 15.00 h

## Venue

Hotel

Barceló Hotel Hamburg

Ferdinandstrasse 15

20095 Hamburg

Phone: +49 (0) 40 22 63 62 0

Fax: +49 (0) 40 22 63 62 999

## Fees (per delegate, plus VAT)

ECA Members € 1,590

APIC Members € 1,690

Non-ECA Members € 1,790

EU GMP Inspectorates € 895

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 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](http://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.

CONCEPT HEIDELBERG

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### For questions regarding content:

Ms Anne Günster (Operations Director) at

+49 (0) 62 21/84 44 50, or per e-mail at

[guenster@concept-heidelberg.de](mailto:guenster@concept-heidelberg.de).

### For questions regarding reservation, hotel, organisation etc. please contact:

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