From theory to practice: Implementing API Process Knowledge

A Pre-Conference Session of the 27th APIC/CEFIC on 22 October 2024 On-site in Vienna, Austria or live online

Send us your questions and real-life scenarios/challenges!

Highlights

Practical assistance and experiences how to implement API Process Knowledge from a quality and regulatory perspective of:

- Control Strategy
- Change Control
 - Reprocessing and Rework





a sector group of







Objectives

This Pre-Conference Session provides an overview of the regulatory and quality GMP requirements and approaches for implementing API Process Knowledge in the pharmaceutical industry. During interactive sessions you will get to know:

- Which GMP and regulatory aspects have to be considered for establishing an adequate Change Control Management?
- What do the guidelines tell us for reprocessing and reworking? And what are the practical consequences during the production of APIs?
- What has to be taken into account when implementing control strategies?

Furthermore, you will have the opportunity to reach clarification on ambiguous issues by bringing your questions up for discussion.

This Pre-Conference Session ideally complements the following 27th APIC/CEFIC Global GMP & Regulatory API Conference.

Background

The API world is changing rapidly. Nowadays, Process Knowledge is one of the most important topics in the API industry. Companies are struggling with the challenge how to implement and how to obtain Process Knowledge and ensure the quality of their products during their life cycles. On the one hand, for that matter the theoretical obligations need to be considered and well known, while on the other hand the practical implementation needs to be valuable and manageable in the daily API work.

Three hot topics out of the long list of options how to obtain and implement Process Knowledge in your company are selected and will be considered and explained during the Pre-Conference Session. Besides explaining the requirements of the guidelines of e.g. the ICH Q7 and ICH Q12 guideline for the respective themes, the speakers will share their approaches, experiences and best practices on these API related quality and regulatory topics.

Target Audience

This Pre-Conference Session is designed for all persons involved in the manufacture of APIs especially for persons from production, quality control, quality assurance, technical and regulatory affairs departments. We are also addressing interested parties from the pharmaceutical industry and GMP inspectorates.



APIC Industry Best Practice documents

APIC developed a number of guidances on GMP and Regulatory Affairs, e.g. "Guidance on aspects of cleaning validation in active pharmaceutical ingredient plants" or "How to do – Interpretation of ICH Q7 document & « Review form »".

All APIC guidance documents are available for free download on the APIC/CEFIC website: https://apic.cefic.org/publications/

Important Information

The presentations will be made available to you prior to the Pre-Conference Session as PDF files. After the event, you will automatically receive your certificate of participation.

Programme

Control Strategy from a Quality perspective

- Sound science & risk management all along the life cycle
- Quality trends
- Elements of a control strategy
- QRM as integral part of quality by design
- Areas of controls (hierarchy and practical examples)
- Impact on cost of quality

Control Strategy from a Regulatory perspective

- The control strategy in the regulatory files
- A complete story
- What to share with the customers

Take advantage of the experiences of our speakers and send us your questions and real-life scenarios/challenges prior to the Pre-Conference Session. Your questions and examples are welcome and will be answered as comprehensively as possible by the experts during the Q&A sessions and exchange sessions.

Changes from a Quality perspective

- Different changes during product life cycle
- Scope and importance of a change control
- General change control requirements
- Detailed change control requirements for specific systems (i.e. materials, process, equipment, utilities, specs & methods) with practical examples
- Implementation requirements

Reprocessing and rework: options and obligations

- Definitions
- ICH Q7 section 14.2 and ICH Q7 Q&A section 14
- Q&A's on practical real-life examples

Changes from a Regulatory perspective

- Change procedures in different regions
- Supply chain issues related to changes
- The impact of ICH Q12

Speakers



Marieke van Dalen

Aspen Oss B.V., The Netherlands

Ms van Dalen is the global regulatory specialist in the regulatory group dedicated to APIs, with over 35 years of experience in the regulatory field. She is a board member of APIC, the European API Industry organization, and she actively participates and/or (co-)chairs in a number of task forces. She frequently represents APIC in meetings and conferences organized by EMA, EDQM, CPhI etc.



Francois Vandeweyer

VDWcGMP Consultancy, Belgium

Mr Vandeweyer joined Janssen Pharmaceutica (part of Johnson & Johnson) in 1981 in chemical development. Until 1995 increasing responsibilities within the organisation mainly in the Quality Control Unit. Starting from 1995 he joined the QA department. Several Senior Manager responsibilities. 2005 Sr Manager GMP Compliance Chemical Operations Belgium (sites Geel – Olen – Beerse). 2009 Director Global Compliance EMEA/AP for Johnson & Johnson. Since May 2019 he is a freelance consultant.

Date Pre-Conference Session

Tuesday, 22 October 2024, 09.30 - 17.15 h (Registration and coffee 09.00 - 09.30 h - for taking part onsite in Vienna only)

All times mentioned are CEST.

Venue - for taking part onsite in Vienna only

Austria Trend Parkhotel Schönbrunn Hietzinger Hauptstr. 10-14 1130 Vienna, Austria Phone: +43 (1) 878 04 0 parkhotel.schoenbrunn@austria-trend.at

Technical Requirements

We will stream the pre-conference and recommend using the latest version of Chrome, Firefox, Edge or Safari to participate. Technical instruction for the livestream will be provided shortly prior to the conference.

Fee

EUR 1090.- per delegate plus VAT.

A special fee of 990,- Euro is granted to participants who also register for the 27th APIC/CEFIC Global GMP & Regulatory API Conference.

The registration fee is payable in advance after receipt of invoice. VAT is reclaimable.

Accommodation - for taking part onsite in Vienna only

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.apiconference.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 P.O. Box 10 17 64 D-69007 Heidelberg, Germany Phone +49 (0) 62 21/84 44-0 Fax +49 (0) 62 21/84 44 34 info@concept-heidelberg.de www.concept-heidelberg.de

For questions regarding content:

Ms Anne Günster (Operations Director) at +49 (0) 6221/84 44 50, or at guenster@concept-heidelberg.de

For questions regarding organisation etc.:

Ms Sarah Schmidt (Organisation Manager) at + 49 (0)6221/84 44 16, or at s.schmidt@concept-heidelberg.de

If the bill-to-address deviates from the specification to the right, please fill out here:

> CONCEPT HEIDELBERG P.O. Box 10 17 64 Fax +49 (0) 6221/84 44 34

69007 Heidelberg Germany

General terms and conditions

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 4 weeks prior to the conference 10 %, - Cancellation until 3 weeks prior to the conference 25 %, - Cancellation until 3 weeks prior to the conference 25 %,

Cancellation until 2 weeks prior to the conference 50 %

- □ From theory to practice: Implementing API Process Knowledge, 22 October 2024 in Vienna or live online
- □ I also register for the 27th Global GMP & Regulatory API Conference, 23-24 October 2024 in Vienna or live online

Please choose 5 out of 10 parallel sessions (one choice in Session A, B, C, D and one in Session E): **Parallel Session A**

Session 1: Title to be announced

- \square Session 2: Differences in the focus of inspections by non-EU authorities
- **Parallel Session B**
- Session 3: Global DMF challenges
- Session 4: Prevention of (Cross-)Contamination in Non-Sterile API Manufacturing Processes Parallel Session C
 - Session 5: ICH M4Q(R2): designing the Common Technical Document for the future
- Session 6: New APIC Guidance on aspects of cleaning validation in API plants **Parallel Session D**
- Session 7: Title to be announced
- Session 8: Recycled raw materials in API manufacturing process for a greater sustainability Parallel Session E
- Session 9: API Registration in China: Recent Industry Experience Session 10: A new era in the assessment of Process-Equipment-Related Leachables (PERLs)

- l will participate on-site in Vienna.
- participate live online.
- □ decide later.

□Mr □Ms Title _

First name, surname

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