



Speakers



Dr Rainer Gnihl
GMP Inspector for EMA



Arno Hoekstra
Kite Pharma



Dr Andreas König
Fidelio Healthcare



Dr Jens-Uwe Rengers
JeRo Consulting

Improve your Quality Reviews

PQR, APR, Management Review, Quality Metrics



Live Online Training on 03/04 September 2020



Highlights

- Quality Reviews in the Context of FDA, EU and ICH
- Expectations of the Agencies
- How to set up efficient
 - PQRs and APRs
 - Management Reviews
 - Quality Metrics

Objectives

This course examines regulatory requirements, provides insight into inspectors' expectations and explains tools for improving your documented review processes.

Based on real examples you will learn how you can implement and improve your Quality Reviews and use them more efficiently.

Background

Quality Reviews and Metrics are critical GMP elements. They are an integral part of a pharmaceutical quality system and provide an opportunity to assess and control relevant processes.

Both parts of the EU-GMP Guidelines require the Product Quality Review (PQR) to verify the consistency and appropriateness of existing processes, but also to identify product and process improvement opportunities.

The FDA 21CFR 211 requires an Annual Product Review (APR) to evaluate annually the quality standards of each drug product.

All relevant guidance do also consider a Management Review to be an appropriate instrument to assess adequacy and effectiveness of quality systems.

All these different reviews could result in a tremendous work load or they can be performed in an efficient way with useful results – depending on how they are organised. Therefore it is very important to understand the requirements and the idea behind it and to see how these tools can be used more efficiently.

Target Audience

This Education Course is designed for managers, supervisors and all other staff members in the pharmaceutical and API industry who are involved in preparing and compiling Quality Reviews and Metrics.

Moderator

Wolfgang Schmitt
CONCEPT Heidelberg (on behalf of ECA)

Programme Thursday, 03 September

All times in CEST:

9.00 – 9.10 h
Introduction

9.10 – 10.00 h
Quality Reviews in the Context of FDA, EU and ICH Requirements and Expectations

- EU-GMP: which types of Quality Reviews are required?
- EU Quality System Review (overview)
- How to achieve EU-GMP compliance
- ICH/US-FDA view on the situation (overview)
- EU Product Quality Review (PQR)
 - Technical terms and aims of PQR
 - What documents and data should be reviewed?
 - Are EU-requirements the same for APIs & medicinal products?
 - What about US-FDA and ICH?

10.00 – 11.00 h
PQR and APR

- How to combine PQR and APR in an efficient way
- Well-proven PQR/APR designs
- Interface to Regulatory Affairs
- Certainties (PQR/APR in Custom Manufacturing, how to deal with limited numbers of batches ...)



11.00 – 11.10 h
Time for Live Q&As

11.10 – 11.30 h
Break

11.30 – 12.30 h
Quality Reviews in the Light of Inspections - Expectations of the Agencies

- Inspectors view on critical parts of EU-PQR
- Practical implementation and inspection
- PQR and contract manufacturing
- Comparison EU-PQR and US-APQR (inspectors point of view)




12.30 – 13.00 h
Time for Live Q&As

13.00 – 14.00 h
Break

14.00 – 15.15 h
Discussion of given PQR-Examples
Based on real examples, the speaker will discuss the content and lay-out of PQRs:


- What is useful?
- What is ambiguous?
- What could be improved?

 15.15 – 15.30 h
Time for Live Q&As

15.30 – 15.45 h
Break

15.45 – 16.45 h
Set up of efficient PQRs and APRs

- How to profit from existing QA Systems in PQR/APR and vice versa
- Best practices
- Time/efforts needed
- Ongoing data collection
- Foreseeable complications/advantages
- Well-proven examples

 16.45 – 17.00 h
Time for Live Q&As

Programme Friday, 04 September

All times in CEST:

08.30 – 09.40 h
Management Review

- Definition, scope, objectives
- Organisation
- Participants, responsibilities
- Topics to be presented: input and output
- KPIs per system
- Examples and experience


09.40 – 10.00 h
Break

10.00 – 10.40 h
Using KPI in Quality Reviews and in Communication with Authorities

- Current status of the requirements
- Key areas and data to be submitted
- How industry can prepare to meet the expectations

10.40 – 11.40 h
Quality Reviews in Contract Manufacturing

- Customer QMRs - content, scope, frequency, organisation
- Interface with Business Management Reviews
- Assessment of data, trending and decision making
- Actions, follow-up
- „Face to Face“ or telecon?

 11.40 – 12.00 h
Time for Live Q&As


12.00 – 13.00 h
Break

13.00 – 14.00 h
Kite Pharma Case Study: Management Review - from Data Collection to Evaluation and Reporting

- Collection and preparation of data: time/efforts needed, automatic vs. manual data capture
- Evaluation of deviations and changes
- Interpretation of data: what is the data telling us?
- How to report the data and information gained

14.00 – 15.00 h
Review Management: Bringing them all together in an efficient Way

- How to set up an integrated data, review and report management
- How to avoid double work

 15.00 – 15.30 h
Time for Live Q&As

Speakers



Dr Rainer Gnibl,
GMP Inspector, District Government of
Upper Bavaria, Germany

Dr Rainer Gnibl is pharmacist and GMP Inspector for the District Government and the EMA and performs GMP inspections worldwide. Before that, he was working for the Bavarian Ministry of Environment and Health. Rainer Gnibl also holds a lectureship at the University Erlangen-Nürnberg.



Arno Hoekstra
Kite Pharma EU B.V., Netherlands

Arno Hoekstra is Senior Manager Quality Systems and Chair of the Quality Management Review Board, Change Control Board and Deviation Board. He has more than 20 years QA experience within different pharmaceutical companies.



Dr Andreas König
Fidelio Healthcare Limburg GmbH, Germany

Dr Andreas König is General Manager of Fidelio Healthcare Limburg GmbH. Before that he was amongst others Senior Vice President Corporate Quality & HSE at Aenova Holding GmbH and Vice President Global Quality Operations Animal Health at Schering Plough.



Dr Jens-Uwe Rengers
JeRo Consulting, Switzerland

Prior to the funding of his consultancy business, Jens-Uwe Renger acted as General Manager at Akorn AG. Before that he was Director Quality and QP and held different other roles at Byk Gulden (now Takeda), Cytos Biotechnology AG and Siegfried Ltd.

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Reservation Form (Please complete in full)



Improve your Quality Reviews – Live Online Training on 03/04 September 2020

Title, first name, surname

Department

Company

Important: Please indicate your company's VAT ID Number

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Country

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2. If you have to cancel entirely we must charge the following processing fees:
 - Cancellation until 2 weeks prior to the conference 10 %
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German law shall apply. Court of jurisdiction is Heidelberg.

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Date of the Live Online Training

Thursday, 03 September 2020, 9.00 – 17.00 h CEST

Friday, 04 September 2020, 8.30 – 15.30 h CEST

Technical Requirements

For our webinars, we use Cisco WebEx, one of the leading suppliers of online meetings. At <http://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)

ECA Members € 1,490

QP Association Members € 1,490

APIC Members € 1,590

Non-ECA Members € 1,690

EU GMP Inspectorates € 845

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

Registration

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

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 book the recording of the Live Online Training at any time at <https://www.gmp-compliance.org/gmp-webinars/recorded-gmp-webinars>.

Organisation and Contact

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

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