Quality of Pharmaceutical Packaging Systems

From Development to Routine Control

21 – 22 May 2014, Berlin, Germany

SPEAKERS:
Dr Helmut Gaus  
Rentschler Biotechnologie
Torsten Kneuss  
Bayer Pharma
Dr Jörg Zürcher  
Bayer Pharma

LEARNING GOALS:
- Regulatory Requirements: EU and US
- Packaging Related Topics of the Common Technical Document (CTD)
- Specifications for Container Closure Systems
- Defect Evaluation Lists
  - Random Sampling Tables
  - Acceptable Quality Levels (AQL)
- Quality Control
  - Receipt, Identification, Sampling, Testing, Approval, and Rejection of Packaging Materials
- Strategies for Reduced Testing
- Control of Printed Packaging Materials
- Reference Samples of Packaging Materials According to EU GMP Guide Annex 19
- Dimensional Checks: Measurement Equipment and Application Ranges
- Testing of Extractables / Leachables

Participate in 2 Workshops:
- Examples of Defective Packaging Materials
- FMEA: Critical Parameters for a Suitable Quality Control System for Packaging Materials

This education course is recognised for the ECA GMP Certification Programme „Certified Quality Control Manager“. Please find details at www.gmp-certification.eu
Objectives

The focus of this GMP Education Course is on the development and routine control of pharmaceutical packaging systems. Participants will learn how to develop pharmaceutical packaging materials systematically, and how to translate the requirements of the Common Technical Document (CTD) to regulatory documents for packaging materials.

The course will also focus on the testing of a variety of different packaging materials, as carried out in every incoming-goods laboratory in quality control.

This includes the setting of sound and scientific specifications and Acceptable Quality Levels (AQLs), the control of dimensions, and the control of printed packaging materials.

The impact of EU Annex 19 on packaging material samples will also be discussed.

Background

There is a great number of regulatory requirements on pharmaceutical packaging materials, in the pharmacopoeias, the GMP regulations, in the FDA guidances, etc. Packaging materials also have to be described in the registration process of a drug product according to the requirements of the CTD.

And the pharmaceutical manufacturer has to guarantee that only such packaging materials are used that are correctly printed, in conformity with the specifications and in compliance with the regulatory requirements. The quality control unit is responsible for the control of pharmaceutical packaging materials including the receipt, identification, sampling, testing, and approval or rejection of drug product containers and closures.

In order to determine the scope of the tests for the quality control of pharmaceutical packaging materials, the „Defect Evaluation Lists“ have proved efficient. The responsibility for the tests lies now more and more with the manufacturers of packaging materials, while the pharmaceutical industry tries to reduce testing at the same time. However, as a precondition for this, additional QA measures, like supplier qualification, audits and supply agreements, have to be taken.

Target Audience

This GMP Education Course is directed at employees working in pharmaceutical research and development, regulatory affairs, quality control, incoming goods control of packaging materials, and quality assurance departments. The Course is also intended for staff of manufacturers and suppliers of packaging materials.
Quality Control of Primary Packaging Material
- What is a suitable QC system for Primary Packaging Materials
- Definition of critical parameters
- Best practice in testing
- AQL testing, skip lot
- Must to have QA systems (i.e. OOS, complaints)
- Sample management incl. reference samples

Dr Helmut Gaus, Rentschler Biotechnologie GmbH

Supplier Management
- Supplier qualification and audits
- Supply agreements and supplier qualification
- Quality standards for suppliers
- Cascade of Quality Control, reduced testing
- Sampling Plans

Dr Helmut Gaus, Rentschler Biotechnologie GmbH

WORKSHOP II
Risk Management (Focus: FMEA)

The aim of this workshop is to define in small discussion groups the critical/major parameters to build up a suitable quality control system for your packaging materials. Focus will be on the practical application in a FMEA. The groups will evaluate:
- What, why, and where to test
- Value of FMEA in the Quality Control concept for packaging materials
- Strategies for reduced testing

Moderator: Dr Helmut Gaus

Dimensional Checks in Packaging Development and Quality Control
- Measurement equipment: overview
- Application ranges
- Practical examples

Torsten Kneuss, Bayer Pharma AG

Control of Printed Packaging Materials
- Legal requirements
- Level of certification
- Sample size & test procedures
- Reference samples vs. retention samples (Annex 19)

Dr Helmut Gaus, Rentschler Biotechnologie GmbH

Technical Specifications
- Scope & content
- Concept (proposal)
- Template and practical example

Torsten Kneuss, Bayer Pharma AG

Testing of Extractables/Leachables
- Regulatory Background
- Principles of Extractable and Leachable testing
- Potential Extractables of different Container Materials

Dr Jörg Zürcher, Bayer Pharma AG

Moderator

Dr Jörg Zürcher, Bayer Pharma AG, Germany

Speakers

Dr Helmut Gaus
Rentschler Biotechnologie GmbH, Laupheim, Germany

Dr. Gaus started at Merckle/ratiopharm, in 2001 he took over at Novartis-Generics, the position of Qualified Person and Head of Quality Control. From 2003 to 2006 he was Head of Quality Control at Vetter Pharma. Since 2006 he is Qualified Person and Vice President Quality Control at Rentschler Biotechnologie. Within his various positions in the pharmaceutical industry the incoming inspection of packaging components was always part of his responsibility.

Torsten Kneuss
Bayer Pharma AG, Berlin, Germany

Torsten Kneuss studied Business Administration and Engineering. He joined Schering AG / Bayer Schering Pharma AG in 1996. Since 1999 he has been working with pharmaceutical materials, including several years within the field of packaging quality control. Since 2007 he has been employed in packaging development, lately as an Application System Development expert. Since November 2010 he has been working as project coordinator within Contract Manufacturing Biotech, and as Operations Manager he is responsible for pre-filled syringes and medical devices.

Dr Jörg Zürcher
Bayer Pharma AG, Berlin, Germany

Dr Zürcher joined Schering (since 2007: Bayer Pharma) in 1990. Starting with systems for solid and semi-solid formulations his focus is now on the development of state-of-the-art container closure and application systems for liquid dosage forms, sterile products, inhalatives and ophthalmics.

Social Event

On 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.
Reservation Form (Please complete in full)

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*Title, first name, surname
Company
Department
Important: Please indicate your company's VAT ID Number
P.O. Number, if applicable
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City
Zip Code
Country
Phone/Fax
E-Mail (please fill in)

General terms and conditions
1. We are happy to welcome substitute colleagues at any time.
2. If you cannot attend the conference and notify us by fax/phone prior to the conference, we will hold your registration until the day of the conference. After that, we will cancel your registration. The registration fee is non-refundable.
3. No cancellation due to non-payment or non-arrival is accepted.
4. Conference fees are payable in advance after receipt of invoice and includes conference documentation, dinner on the first day, lunch on both days and all refreshments.
5. VAT is reclaimable.
6. Registration will be confirmed upon receipt of payment.
7. The official conference language will be English.
8. For questions regarding content: Dr Günter Brendelberger (Operations Director) at phone +49-62 21 / 84 44 40, or by e-mail at brendelberger@concept-heidelberg.de.
9. For questions regarding reservation, hotel, organisation etc.: Ms Marion Weidemaier (Organisation Manager) at phone +49-62 21 / 84 44 46, or by e-mail at weidemaier@concept-heidelberg.de.

Date
Wednesday, 21 May 2014, 09.00 h - 18.15 h
(Registration and Coffee 8.30 h - 9.00 h)
Thursday, 22 May 2014, 08.30 h - 16.30 h

Venue
Steigenberger Hotel Berlin
Los-Angeles-Platz 1
10789 Berlin, Germany
Phone +49 / (0) 030 2120 - 0
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Fees
ECA Members € 1,490.- per delegate plus VAT
APIC Members € 1,590.- per delegate plus VAT
Non-ECA Members € 1,690.- per delegate plus VAT
EU GMP Inspectorates € 845.- per delegate plus VAT

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 when you have registered for the event. Please use this form for your room reservation to receive the specially negotiated rate for the duration of your stay. 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
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