GMP for Vaccine Manufacturers
Current Regulatory Requirements and Practical Implementation

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

SPEAKERS:

Dr Robert Eskes
Novartis Technical Operations

Mag. Petra Falb
AGES - Austrian Agency for Health and Food Safety

Dr Andreas Neubert
IDT Biologika

Dr Volker Öppling
PEI, German Federal Institute for Vaccines and Biomedicines

Robert Schwarz
Shire, Austria

Dr Jörg Weyermann
GlaxoSmithKline

HIGHLIGHTS:

- Four Case Studies:
  1. New building of a multipurpose vaccine production facility
  2. New filling facility in direct cooperation with an existing bulk production
  3. Different fogging and gassing systems for decontamination of isolators, lyophilizers and clean rooms
  4. GMP manufacturing of recombinant viral vaccines for clinical trials

- Peculiarities of viral and bacterial vaccines
- GMP issues for upstream and downstream processing
- Staff Safety

With case studies on new manufacturing buildings and the validation of fogging/gassing systems.

04-05 December 2018, Barcelona, Spain
Objectives

The development and production of vaccines makes high demands on the manufacturing pharmaceutical industry. The special requirements on handling and safety with live organisms necessitate measures which exceed the requirements of classic pharmaceutical manufacturing.

Topics like the enhanced risk of cross-contaminations, questions about individual safety of staff and the issues of cleaning and disinfection of rooms and equipment concern a vaccine manufacturer in a considerable scale.

Specifically the demands of the necessary bio safety classes with negative pressure of rooms versus that of aseptic processing with positive pressure requires a well thought-out design of vaccine facilities.

Also, the safety of environment and waste disposal should receive proper attention already in the design phase. But the dedicated requirements on staff safety are also a challenge in vaccine manufacturing.

This Course will give you the possibility to see the theoretical background as well as the practical implementation of GMP requirements in the vaccine production. A combination of theoretical requirements and practical case studies is the best way to learn this.

Speakers from regulatory bodies, consulting and practising experts will give you the chance to get to know the different views and you will have ample opportunity to discuss with speakers and other participants about specific issues.

Background

“Vaccines are expected to be very safe” is one of the headlines in the presentation of the CBER “Vaccine safety team”. At the same time, new vaccines are needed for diseases for which currently no vaccine is available, and production technologies need improvement to deal with the shortage of certain types of vaccines. This has led to the emergence of new technologies. One of the important questions from the authorities however is “How safe are the new technologies”. The FDA has issued a draft guideline on new cell substrates for vaccine manufacturing to detail requirements in this area.

In the development of new technologies for the pharmaceutical and biopharmaceutical production of vaccines again the question of GMP compliance and safety is emphasised.

Furthermore, with the Quality Initiative for the 21st Century from the FDA new guidelines have been issued, which have an impact not just on the conventional pharmaceutical industry but also on vaccine manufacturers. Risk management and quality in design are essential in the implementation of new technologies and the introduction of new vaccines. Ensuring the expected safety is one of the greatest challenges of all vaccine producers.

Target Audience

The course is designed for personnel of pharmaceutical industries, their suppliers and regulatory bodies who are responsible for quality control and/or quality assurance in vaccine/biopharmaceutical production, manage the vaccine production, establish the operator protection, audit vaccine manufacturers, design or operate vaccine production sites.

Programme

GMP for vaccines: What are the Issues?
- Differences between vaccines and conventional products
- Inactivated and live vaccines
- Control of vaccine strains and cell lines
- Risk of (cross-)contamination
- (Bio)safety issues

The Peculiarities of Bacterial Vaccines
- Types of vaccines available
- Manufacturing of classical/modern bacterial vaccines
- Challenges in manufacturing (quality/regulatory issues)
- New technologies and products

The Peculiarities of Viral Vaccines
- from viral seeds to finished products
- Requirements for raw and starting materials
- Efficient process and product control
- Setting specifications adequately
- Appropriate tests and assays for product release
- Stability testing
- Viral safety aspects
- TSE compliance
- How to deal with OOS results?
- Requirements for early and late clinical trial phases

Containment, Biological Safety and Product Protection
- Containment, product safety versus environmental safety
- Primary containment and additional measures
- Negative pressure areas in aseptic manufacturing
- Decontamination of facilities
- Personnel as critical component in containment

Case Study: New Building of a Multipurpose Vaccine Production Facility
- Practical issues with flow of material, personnel and waste material
- Clean room qualification
- Segregation of cell preparation, virus production and downstream processing
- Change over procedures for manufacturing campaigns

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Case Study: Different Fogging and Gassing Systems for decontamination of Isolators, Lyophilizers and Clean Rooms

- Different gassing systems
- System qualification
- Validation of a dry fog detergent

Issues of Staff Safety

- Requirements and Guidelines
- Differences Vaccines Products and Plasma Products
- Use of S3 Coveralls
- Environmental Health and Safety challenges
- Examples from daily business

Decontamination, Virus Inactivation and Virus Removal Techniques

- Decontamination of surfaces
- Validation of decontamination procedures
- Virus inactivation: principles and methods
- Virus removal methods
- GMP issues on virus inactivation and virus removal techniques

cGMP Issues for Upstream Processing

- General GMP concerns for upstream processing
- Raw materials and media preparation
- Cell culture
- Virus culture
- Inactivation of microorganisms

GMP Manufacturing of Recombinant Viral Vaccines for Clinical Trials

- Regulatory expectations for vaccine batches for phase 1/2/3 clinical trials
- Development vs. validation
- Regulatory expectations for implementation of analytical methods – qualification and validation
- Contract manufacturing of IMPDs

cGMP Issues for Downstream Processing

- General GMP concerns for downstream processing
- (ultra)filtration techniques
- (ultra)centrifugation techniques
- Sterile filtration and aseptic processing

Case Study: Planning and Realization of a new Filling Facility in direct Cooperation with Existing Bulk Production

- Requirements of design
- Issues of construction
- Qualification challenges

Speakers

De Robert Eske, Novartis Technical Operations, Austria
Currently Robert Eske is Head MS&T Unterach / Novartis Technical Operations – Aseptics at EBE-WE Pharma in Unterach, Austria. From 2014 to 2016 he was Quality Assurance head for Third Party Manufacturing at GSK, former Novartis Vaccines and Diagnostics. GmbH. Before that, he held positions as Quality Assurance head for Rabies and FSME bulk production and in Manufacturing Science and Technologies leading the manufacturing support and tech transfer of Rabies and FSME bulk production also at Novartis Vaccines and as Validation manager at CSL Behring GmbH.

Petra Falb, AGES - Austrian Agency for Health and Food Safety
Petra Falb studied at Veterinary University Vienna, (Austria). From 1998 to 2001 she worked as scientist at the Institute for Virology and later at the Institute for pathology. 2001-2003 she was self-employed as veterinary surgeon. In 2003 she joined the AGES with responsibilities in quality assessment of human and veterinary vaccines (national, decentralised and centralized procedures). Until 2016 her focus was on viral vaccines. In 2017, she took over new responsibilities for veterinary vaccines.

Dr Volker Öppling, Paul Ehrlich Institut, Germany
Head of section “Microbiological Vaccines” (all bacterial, fungal and parasitic vaccines). Main responsibilities: assessment of marketing authorisation applications (quality, preclinical and clinic), batch release, assessment of clinical trial applications, provision of regulatory and scientific advise, managing of regulatory affairs issues.

Dr Andreas Neubert, IDT Biologika, Germany
Andreas Neubert completed his study of veterinary medicine with graduation. He works since several years in different positions at IDT and is currently head of production there.

Robert Schwarz, Shire, Austria
Robert Schwarz studied biotechnology. After working in a medicinal lab as medical/technical analyst he joined Baxter, Vienna in 2001. From 2001 to 2005 he was coordinator of environmental monitoring at Baxter, Vienna. Since 2005 he is validation specialist responsible for equipment qualification, sterilisation validation and cleaning validation.

Dr Joerg Weyermann, GlaxoSmithKline, Germany
Joerg Weyermann is head Head Quality Assurance at GSK, former Novartis Vaccines and Diagnostics GmbH. Until 2009 he was the Head Quality Operations for Sandoz Industrial Products GmbH. Before that he was Head Quality Control at Sandoz.

Moderator

Axel H Schroeder, Concept Heidelberg
Reservation Form (Please complete in full)

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04-05 December 2018, Barcelona, Spain

Date

Tuesday, 04 December 2018, 09.30 h – 18.00 h
(Registration and coffee 09.00 h – 9.30 h)

Wednesday 05 December 2018, 08.30 h – 17.00 h

Venue

Barcelo Sants Hotel
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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, lunch on both days and dinner on the first day 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 event. 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.

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.

Conference Language

The official conference language will be English.

Organisation and Contact

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

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