Speakers



Dr Martin Becker ECA Visual Inspection Group



Haluk Dönmez B. Braun



Sabine Eggensberger Vetter Pharma-Fertigung



Dr Helmut Gaus ECA Visual Inspection Group



Felix Krumbein Head ECA Visual Inspection Group



Christof Langer OSConsulting



Dr Stephen Langille formerly FDA



Dr Daniel Müller GMP-Inspector



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Visual Inspection of Parenterals

State-of-the-art Visual 100% Inspection

6/7 November 2024, Berlin, Germany



- Fundamentals of Visual Inspection on 5 November
- All participants receive the new version of ECA's Guide on "Visual Inspection" for free

Highlights

- Current FDA Requirements for Visual Inspection of Injectables
- Compliance with EU Annex 1 and the Pharmacopeias
- Requirements from an EU GMP Inspector's View
- Usage of AI in Automated Visual Inspection
- Semi-Automated Visual Inspection
- Approaches for the Inspection of hard-to-inspect Container-Systems
- Transfer of a Visual Inspection Process
- Case Study Vetter Pharma: A Particle Lifecycle Concept

Fundamentals of Visual Inspection

5 November 2024

The training course on visual inspection which takes place before the Particles Conference gives you an understanding of the fundamentals of visual inspection of injectable products, applicable to manual and automated inspection. You will also learn how to implement an automated system on the basis of the manual inspection and how to qualify it.

Skills you will develop through the course:

- Ensuring GMP-compliance in manual inspection
- Setting up a qualification strategy for automated systems
- GMP-compliant routine operation of automated systems

Content

General requirements

- Requirements of the Pharmacopeia
- Defect categorisation
- Test kits for training, qualification and routine

Manual Inspection

- Qualification and training of personnel
- Standardisation of working conditions
- AQL in the manual inspection

From Manual to automated inspection

- Usage of the Knapp and the modified Knapp test
- Cross validation during the PQ phase
- Evaluation matrices

Automated inspection

- Importance of particle detection rates
- System-Suitability, Requalification and revalidation

Evaluation of inspection data & batch release

- Trending of inspection results
- AQL Testing
- Re-inspection allowed or not?

Course Trainers

Dr Helmut Gaus, formerly Boehringer Ingelheim Felix Krumbein, Head ECA Visual Inspection Group

ECA Guide on Visual Inspection of Parenterals



All participants receive the new version 4.0 of ECA's Guide on "Visual Inspection of medicinal products for parenteral use" for free.

Visual Inspection of Parenterals

6/7 November 2024

Objective

Main topic of this conference is the detection of defects like particles in injectables and their evaluation. Besides the current regulatory requirements with regards to particulate matter, routine 100% inspection of injectables will be addressed. Manual inspection as well as automated inspection systems will be covered, including validation, training, defect categories, AQL testing and trending.

Background

In most cases particles found in parenteral medicines will lead to a quarantined product or even to the recall of the product – as we have seen in the last years in the cases of several pharmaceutical companies. Responsible staff in charge will have to start root cause analysis to find the source of the particles and will have to do an evaluation of batches already shipped.

There is still confusion within the global pharmaceutical industry with regard to the requirements for testing for visible particles. After the USP chapters <790> and <1790> were published, things have become much clearer, at least for the US.

In Europe chapter 5.17.2 of the European Pharmacopoeia now also gives further advice. However, many questions remain, e.g. concerning training, re-testing, detection capabilities and revalidation of inspection systems.

Furthermore, there has been a recognisable trend towards automated inspection machines throughout the last years. High expectations are also placed on the use of artificial intelligence. The challenge for pharmaceutical companies is to find a suitable machine for their products and to determine reasonable inspection parameters during qualification and validation. But also during routine process there are questions arising like re-testing and the usage of test-sets, doing AQL-Testing as well as the adjustment of parameters of the vision system.

We will address those topics during the conference and discuss and answer questions on

- The latest compendial requirements concerning particulate matter (EU & US)
- How to comply with FDA's new guidance on Visual Inspection
- Compliance with the new EU Annex 1
- Implementation of artificial intelligence in the automated inspection process
- Reduction of false rejects in automated inspection systems
- How to inspect hard-to-inspect containers
- How to set up a company-wide particle reduction programme
- How to transfer a visual inspection process during a site change

The fundamentals, such as training of operators in manual inspection, AQL testing, trending and the validation of an AVI system are content of the Pre-Conference course on 5 November.

Target Audience

This course is directed at staff from sterile operations involved in the 100% inspection process, that is production, quality and engineering. But also suppliers of primary packaging materials and inspections technology are target group of this event.

Moderator

Christof Langer, OSConsulting

Programme

Regulatory Requirements for the Visual Inspection of Parenterals

- Compendial requirements
 - 100% visual inspection & AQL testing
 - PharmEur, USP, JP similarities and differences
- News from the Annex I
- Risk management considerations

Current FDA Requirements for Visual Inspection of Injectables (remote presentation)

- Difference between current FDA and EU requirements
- Use of quality risk management for the prevention of visible particulate contamination
- FDA Visible Particulate case studies
- Drug-Device combination product considerations

Visual Inspection of Parenterals – a GMP Inspector's View

- Applicable regulations & guidance
- Current requirements for pharmaceutical industry
- Expectations of a GMP Inspector
- Examples of observations

Presentation of the new Version 4.0 of the ECA Guide

The new version 4.0 of the ECA guide has recently been released. The presentation will introduce the guide itself and the changes in the new version. Among other things a chapter on semi-automated visual inspection has been added. Also, the description of uninterrupted inspection times & breaks in manual visual inspection has been adapted. And the Annex 1 requirement for 'performance checks at regular intervals throughout the batch' for AV inspection was taken into account.

Semiautomated Inspection – Advantages and Risks

- Regulations for the use of SAVI
- Technical setup of a SAVI system
- Differences manual and semi-automated inspection
- Qualification of inspection personal
- Typical inspection errors and walkarounds

Practical Approaches for the Inspection of hard-toinspect Container Systems

Part I

- Inspection of Bags
- Single chamber and multi-chamber bags
- Inspection of Blow-Fill-Seal containers
- Inspection of Form-Fill-Seal containers
- Manual, semi-automated and fully-automated approaches
- Two step inspection

Part II

- Use of artificial intelligence
- General approach
- Training and Machine Learning
- Testing and Validation
- Limitations

Particle Lifecycle Concept

- What does it tell about the product?
- What can we learn about the visual inspection process and operator qualification?
- Challenges and first experiences

Transfer of a Visual Inspection Process

- Manual inspection
 - Manual process to manual process (same manufacturing location)
 - Manual process to manual process (different manufacturing location)
 - Test set transfer?
 - Create new test set? How, why?
 - What does validation look like?
 - End goals and success (what are we trying to achieve?)
- Automated inspection
 - Automated transfer: same machine (pitfalls)
 - Automated transfer: different machine (pitfalls)
 - End goals and success (what are we trying to achieve?)

Application, Project Planning and Qualification of AI in fully automated Visual Inspection

- Development of robust, reliable and production-ready models in 4 phases
 - Phase 1: Problem identification & description
 - Phase 2a: Specification of inspection concept
 - Phase 2b: Definition of the sample sets (artificial and production samples), creation of the datasets, clarification of the labelling strategy
 - Phase 3: Model design, training and verification a risk-based approach
 - Phase 4: Qualification & validation
 - Processes & technologies
 - Technologies for efficient image data acquisition, variable model technologies, transfer learning / pre-trained models, labelling application
 - Documentation of model development: traceability, risk minimisation and build-up of confidence

Social Event

On 6 November 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.

Speakers



Dr Martin Becker ECA Visual Inspection Group

Martin Becker has many years of experience in the pharmaceutical industry. He worked in analytical

development, QA and production at IDT and Sandoz, among others. He was Head of Technical Operations at Siegfried Hameln GmbH and Director Manufacturing at Baxter Oncology in Halle.



Haluk Dönmez, B. Braun Haluk Dönmez has 23 Years work experience in Life Sciences. His current position is "Head of QM Digital Transformation" in global QM of B.Braun

Melsungen AG.



Sabine Eggensberger Vetter Pharma-Fertigung

Sabine Eggensberger is Director Quality Assurance for Supplier Quality Management/IPC & VI System

at Vetter Pharma-Fertigung. She is also responsible for In-Process-Control/Visual Inspection Systems, Supplier Quality Management Packaging Materials and Single Use Systems.



Dr Helmut Gaus, ECA Visual Inspection Group & former Director Quality Control at Boehringer Ingelheim

Dr Gaus was Head of Quality Control Service at Boehringer Ingelheim, Biotechnology. He has also been working as Vice President Quality Control and Qualified Person for Novartis Generics, Vetter-Pharma and Rentschler Biotechnologie. In 2018 he founded his own company WinSol. He is also member of the ECA Visual Inspection Board.



Felix Krumbein, Head ECA Visual Inspection Group

Felix Krumbein studied optotechnics and image processing and was Head of Inspections-Systems-

Support at Roche Mannheim, where he was responsible for the qua-lification of visual inspection systems. He also was Head of Visual Inspection at INSPECTIFAI where he was responsible for the development of AI-based solutions for fully automated inspection machines. Mr Krumbein is Head of the ECA Visual Inspection Group.



Christof Langer, OSConsulting

Christof Langer studied Biotechnology and is certified Risk Manager as well as a Lean Six Sigma Black Belt. He has been working as Managing Director at

Baxter BioScience, responsible for Operations. Since 2009 he runs his own consultancy business.



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Speakers (cont.)



Dr Stephen Langille, ValSource, formerly FDA

Dr Langille is a senior microbiology Consultant at ValSource. He worked for the US FDA for 19 years in

the Office of Pharmaceutical Quality's Division of Microbiology Assessment as a reviewer, branch chief and division director. He is a member of the United States Pharmacopeia's expert panel on visual inspection and helped author the FDA's Draft Guidance for Industry on the Inspection of Injectable Products for Visible Particulates.



Dr Daniel Müller, GMP Inspector, Germany Daniel Müller is head of a local GMP inspectorate in Germany. Since 2001 he has been working as GMDP inspector, conducting national inspections as well as

EMA and overseas inspections. He is member of Germany's expert groups 'biotechnology & tissue' and 'quality assurance'.

Date of the Conference

Wednesday, 6 November 2024, 09.00 to approx. 17.45 h (Registration and coffee 08.30 – 09.00 h)
Thursday, 7 November 2024, 08.30 to approx. 15.00 h

Venue

DoubleTree by Hilton Berlin Ku'damm Los-Angeles-Platz 1 10789 Berlin

Phone: +49 030 21270

E-Mail: reservations@doubletreeberlinkudamm.com

Fees (per delegate, plus VAT) Conference

ECA Members € 1690 APIC Members € 1790 Non-ECA Members € 1890 EU GMP Inspectorates € 945

The conference fee is payable in advance after receipt of invoice and includes (electronic) conference documentation, dinner on 6 November, lunch on both days and all refreshments. VAT is reclaimable.

Pre-Conference Workshop

ECA Members € 790 APIC Members € 890 Non-ECA Members € 990

The fee is payable in advance after receipt of invoice and includes (electronic) conference documentation, lunch and all refreshments. VAT is reclaimable



Save \leq 300 when booking pre-conference course and conference together.

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 – or search and register directly at www.gmp-compliance.org under the number 21379. To avoid incorrect information, please give us the exact address and full name of the participant.

Presentations/Certificate

The presentations for this event will be available for you to download and print before and after the event. Please note that no printed materials will be handed out on site and that there will not be any opportunity to print the presentations on site. After the event, you will automatically receive your certificate of participation.

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
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 please contact:

Dr Robert Eicher (Operations Director) at +49(0)62 21/84 44 12, or per e-mail at eicher@concept-heidelberg.de.

For questions regarding organisation please contact:

Mr Rouwen Schopka (Organisation Manager) at +49(0)62 21/84 44 13, or per e-mail at schopka@concept-heidelberg.de.