

Speakers



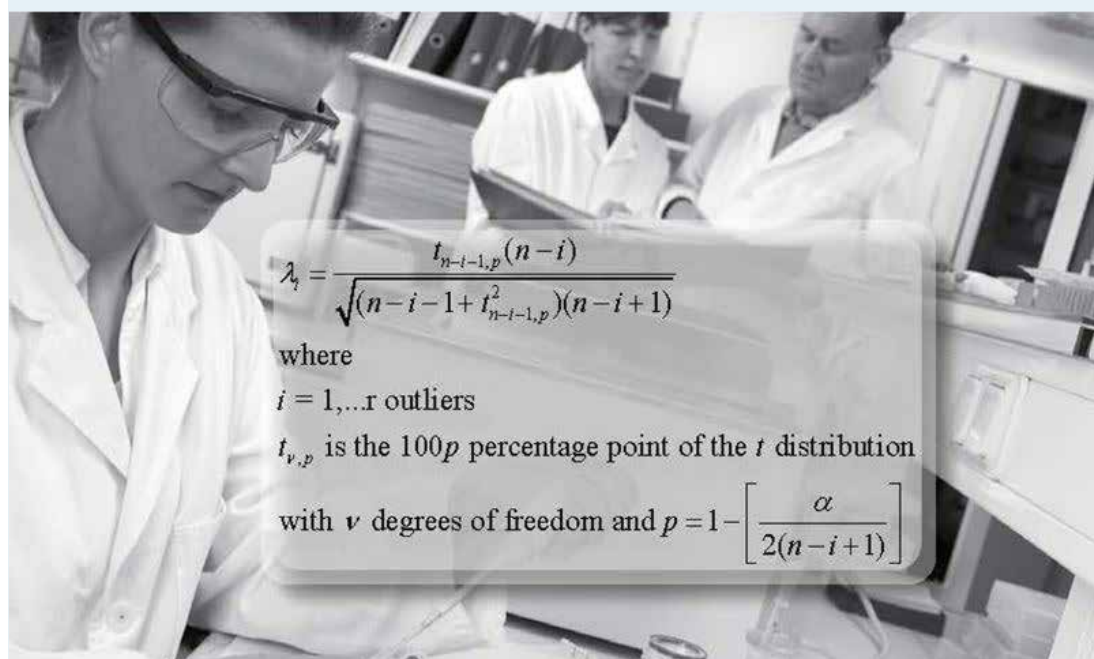
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Ermer Quality Consulting, Germany

Practical Statistical Tools for Analytical Laboratories

Live Online Training on 01/02 December 2021



$$\lambda_i = \frac{t_{n-i-1,p}(n-i)}{\sqrt{(n-i-1+t_{n-i-1,p}^2)(n-i+1)}}$$

where

$i = 1, \dots, r$ outliers

$t_{\nu,p}$ is the $100p$ percentage point of the t distribution

with ν degrees of freedom and $p = 1 - \left[\frac{\alpha}{2(n-i+1)} \right]$

Performance Evaluation and Monitoring for compliant Analytical Procedures and Processes

Highlights

- Participants should gain an understanding of
 - Basic statistical fundamentals
 - Distribution of data and its parameters
 - Accuracy and precision
 - Variability and precision levels
 - Reportable result
 - Linear and non-linear models
 - Performance requirements for analytical procedures
- Participants will be shown how to
 - Apply statistical principles scientifically and pragmatically in their day-to-day business
 - Use statistical simulations
 - Optimise the reportable result for minimum variability
 - Trend data
 - Compare data and methods
 - Establish reliable Reporting/Quantitation Limits

Updated Course for R&D
and QC Laboratories

Objective

Statistical calculations and tools are applied extensively in pharmaceutical analysis including

- Procedure development and validation
- Transfer of analytical procedures
- Setting or verification of specification limits
- Data evaluation, comparison and trending

The ICH Q10 Guideline “Pharmaceutical Quality System”, the FDA Guidances on Process Validation and Methods Validation require monitoring of “process performance and product quality” and “Trend analysis on method performance” throughout the product lifecycle. Hence the appropriate use of statistical trending and evaluation tools has become mandatory.

Consequently, a thorough understanding of statistical fundamentals is essential in order to be able to select parameters and test methods that are ‘fit for purpose’.

Do you speak statistics?

In addition, such an understanding facilitates the communication with other technical and regulatory functions applying statistical tools in order to ensure an overall consistent approach.

Background

The Live Online Training will provide the participants with recommendations, tools and examples to apply scientifically and pragmatically sound statistical principles to their day-to-day business as well as to meet future challenges described above.

The relevance of such statistical tools is also increasingly recognised by the Compendia, as reflected, for example, in the USP General Information Chapter <1010> “Interpretation and treatment of analytical data” and the recently introduced <1033> “Biological assay validation” together with the proposed General Chapter <1220> on Analytical Procedure Lifecycle.

Statistical tools are needed, for example, to evaluate:

- Distribution of data and its parameters
- How to detect outliers and trends?
- How to establish the total variability of the method?
- How to identify method parameters that must be controlled?
- Method performance and specification limits
 - Which accuracy and precision is needed to achieve an acceptable risk of OOS results?
 - Scientifically based justification and optimisation of the reportable result (single or average?)
 - What are the requirements for impurity methods?
- Comparison of methods and data
- What are the requirements for calibration models?
- How to optimise the number of calibration replicates on a scientific basis?

A brief discussion of supporting software tools (e.g. Excel, Minitab, JMP) to facilitate the generation of statistical information in a consistent manner will be undertaken.

One of the main features of this Live Online Training is the balance of presentations and more than five hours of practical exercise workshops which will allow participants to gain ‘hands on’ practical experience in applying the statistical methods described. By means of statistical simulation tools, the participants will gain intuitive understanding of the consequences of appropriate and inappropriate performance parameters, for example the relationship between precision and OOS results.

In order to fully benefit from the workshops, attendees should preferably have a notebook or computer with Excel® 2007 or later available.

Target Audience

This best practice oriented course is designed for analytical laboratory managers and their colleagues charged with the day to day management and evaluation of laboratory data throughout the lifecycle, i.e. in method development, validation, transfer, specification setting, batch release and stability, continuous performance verification and change control.

QA, manufacturing and regulatory affairs professionals will benefit from participation by gaining a clear understanding of the statistical fundamentals which are important to implement scientifically sound and pragmatic tools to conform to GMP and regulatory requirements for example Product Quality Review.

Moderator

Dr Christopher Burgess, Burgess Analytical Consultancy Ltd., UK

Programme

Analytical Procedure Lifecycle Management (USP & ICH initiatives)

- Principles of APLM
- Proposed USP <1220>
- Risk based approach
- Target Measurement Uncertainty
- Decision rules

(Normal) Distribution of Data and its Parameters

- Data shape and its importance
- Characterisation of distributions (Location and Dispersion)
- Probability considerations; all measurements are subject to error
- Populations and samples
- Confidence intervals
- What is an outlier?
- Error of the error



WORKSHOP I

Understanding the Variability (Statistical Simulations)

- Range of expected data
- Variability of standard deviations
- Number of data and reliability of calculated standard deviations

Calculation and Evaluation of Precision Levels

- System precision, repeatability, intermediate precision, reproducibility
- ANOVA: Identification of relevant variance components from injection, measurement, sample preparation, intermediate conditions
- Total variability: precision of the reportable result and its optimisation
- Optimisation of single-point calibration
- Relationship between precision and probability of OOS results
- Practically relevant acceptance criteria for precision



WORKSHOP II

Optimisation of Variability

- Statistically based format of the reportable result (single or average)
- Number of determinations for various levels
- Probability of results outside established limits

Trending of Data

- Why trend?
- Evaluation; do we expect a trend or not?
- Statistical Process Control principles
- Types of Control charts and their application
- Application to stability testing



WORKSHOP III

Control Charts & Trending

- Interactive workshop based on supplied real data sets for interpretation
- Use of Minitab for control charting
- Team working on evaluation and interpretation of trend data

Monte Carlo Simulation of Analytical Procedures

- Principles of Monte Carlo simulation
- Understanding variance contributions and how they combine
- Measurement uncertainty
- Application to analytical procedures
- Examples of unit and complete procedures using Companion by Minitab

Comparison of Data & Accuracy

- Significance (F- and t-test) and equivalence tests
- Statistical significance and practical relevance
- Differences caused by random variability: observed and true bias
- Applications in transfer and cross-validation



WORKSHOP IV

Comparison of Data (Statistical Simulations)

- Significance and equivalence tests: influence of number of data and series
- Differences between means and variability

Calibration Models, Linear and non-Linear

- What is a calibration model?
- What is the difference between linear and non-linear models?
- The principle of least squares and why it is important
- Applying the principles to linear and non-linear models



WORKSHOP V

Linearity (Statistical Simulations)

- Regression range and evaluation of the intercept
- Extrapolation effects

Performance Requirements for Impurity Procedures

- Concentration dependence of precision (Horwitz relation)
- Detection and Quantitation Limits



WORKSHOP VI

Quantitation Limit

- Basics to consider for calculation from linearity
- How to determine appropriately from precision

Summary Workshop & Discussion: Appropriate Choice of Tests/Calculations

- Practical objectives and data sets are provided
- The participants will discuss and define appropriate tests and parameters to be calculated
- The participants are given the calculation results and are asked to make an evaluation
- The defined tests and results are discussed in the audience



Optimised as Online Event – Interaction ensured

Q&A sessions ensure interaction and that all questions are answered.

During the workshops, the participants will be invited to switch on their microphone to join the discussion. **If you want to take part in it, please make sure that you have a headset or that your computer has a built-in microphone.**

In addition, you will have the opportunity to ask questions per chat, which the speakers will then answer. Furthermore, the speakers will work with live poll questionnaires for gathering feedback and asking questions.

Please note: The chat and poll questionnaires are anonymous. The names of the people discussing during the workshops will be visible for all participants. The Live Online Training will not be recorded.



Dr Christopher Burgess
Burgess Analytical Consultancy Ltd., UK

He is a Chartered Chemist and has more than 44 years experience in the pharmaceutical industry initially with Glaxo in Quality Assurance and Analytical R&D and then 24 years in international consultancy. He is a “Qualified Person” in the European Union. He has been appointed to the United States Pharmacopoeia’s Council of Experts 2010 to 2020 and is a visiting professor of the University of Strathclyde’s School of Pharmacy and Biomedical Sciences (SIPBS). In addition, he is the chairman of the ECA Analytical Quality Control Group and a member of the Extended board of European Compliance Academy Foundation. He is also a member of the USP Expert Panel on Validation and Verification entrusted to revise General Chapters <1224>, <1225> and <1226>.

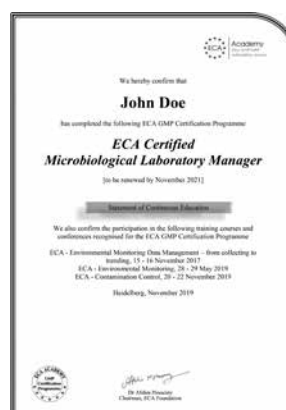


Dr Joachim Ermer
Ermer Quality Consulting, Germany

He has 30 years of experience in pharmaceutical analytics including development products, global responsibilities as Director of Analytical Processes and Technology, Head of Quality Control, and Head of QC Lifecycle Management Frankfurt Chemistry at Sanofi. From 2010 till 2020, he was also responsible for the central reference standard group of Sanofi in Frankfurt. He is member of the USP Expert Committee Measurement and Data Quality, of the Chromatographic Separation Techniques Working Party of the European Pharmacopoeia, and of the EFPIA support team for the update/establishment of ICH Q2/Q14.

GMP/GDP Certification Scheme

Building on your education the ECA GMP/GDP certification programmes provide you with the appropriate supplement to acquire this qualification. Simply choose any three courses within the programme according to your professional interest. Your certificate is then valid for two years. To renew it, you can pick any training from the ECA courses and conferences list within that two-years period – allowing you to broaden your knowledge in GMP and GDP compliance.



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Reservation Form (Please complete in full)



Practical Statistical Tools for Analytical Laboratories Live Online Training on 01/02 December 2021

Title, first name, surname

Department

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P.O. Box 101764
Fax +49 (0) 62 21/84 44 34

D-69007 Heidelberg
GERMANY

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cellation or non-appearance. If you cannot take part, you have to inform us in writing. The cancellation fee will then be calculated according to the point of time at which we receive your message.

In case you do not appear at the event without having informed us, you will have to pay the full registration fee, even if you have not made the payment yet. Only after we have received your payment, you are entitled to participate in the conference (receipt of payment will not be confirmed) (As of January 2012).

German law shall apply. Court of jurisdiction is Heidelberg.

Privacy Policy: By registering for this event, I accept the processing of my Personal Data. Concept Heidelberg will use my data for the processing of this order, for which I hereby declare to agree that my personal data is stored and processed. Concept Heidelberg will only send me information in relation with this order or similar ones. My personal data will not be disclosed to third parties (see also the privacy policy at: http://www.gmp-compliance.org/eca_privacy.html). I note that I can ask for the modification, correction or deletion of my data at any time via the contact form on this website.



Date of the Live Online Training

Wednesday, 01 December 2021, 09.00 - 18.00 h

Thursday, 02 December 2021, 09.00 - 17.00 h

All times mentioned are CET.

Technical Requirements

For our Live Online Training Courses and 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 € 1,590

APIC Members € 1,690

Non-ECA Members € 1,790

EU GMP Inspectorates € 895

The conference fee is payable in advance after receipt of invoice.

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 Recordings

Independent from the Live Online Training Courses, you can also order recordings of selected Live Online Training Courses at the same conditions – at www.gmp-compliance.org/recordings. These recordings will be provided on our media server. All you need to watch it is an Internet browser – no additional software.

Organisation and Contact

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

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