

PharmaFORUM

Webcast Clinical Trials

THE UPCOMING WEBCASTS AT A GLANCE

- Contracts & Agreements
- IT Systems: Risk Management and Validation
- Practical Advice for Risk-based Monitoring
- Safety Reporting in Clinical Trials
- GCP Audits and Inspections
- Data Protection

YOUR BENEFITS

- One live webcast with clinical trial experts every month
- Consolidated information in a short period of time at your work place
- Possibility to directly interact with the speaker via chat

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Concept

Do you work in clinical trials? Then we would like to invite you to our live webcast series in which experts from clinical research will inform you about the latest updates and trends in clinical studies within and beyond the ICH region. You will receive information on the design and conduct of clinical trials, as well as practical tips for successful collaboration at the interfaces to other areas.

You will meet our experts in a virtual conference room. Each meeting will be held as a 1.5 - 2 hour live webcast, presenting the latest news with supporting presentation slides (also for your personal download). Your practical questions will be addressed directly via the chat function, coordinated by the meeting's chairperson.

Are you unable to attend one of the webcasts? No problem! After each live meeting, you will be able to retrieve the recorded webcast (audio and presentation) from our eLearning Centre, using your personal password. This allows you to review each presentation at any time and as often as you like. An optional multiple-choice test finalises each webcast, giving you the possibility to receive a personal certificate.

Your experts in 2020

Dr Andreas Jabs

Alegri Intl. Service GmbH, Frankfurt
Principal Consultant/Manager

Alexander Maur

Kanzlei am Ärztehaus Frehse Mack Vogelsang,
Cologne
Lawyer

Melanie Blessing

Boehringer Ingelheim Pharma GmbH & Co. KG,
Biberach
Head of Supply Chain Management
CardioMetabolic, Global Clinical Trial Supplies Unit

Bettina Schopf

PHARMALOG, Institut für klinische Forschung
GmbH, Ismaning
Head of Clinical Monitoring

Dr Sybille M. Eibert

Senior Expert Medical Writing, Basel

Angela Hartmann

Senior Expert Pharmacovigilance, Darmstadt

Prof. Sebastian Harder

Ethikkommission der Landesärztekammer
Hessen, Frankfurt

Silja du Mont

Regierungspräsidium Freiburg

Karsten Sternickel

PAREXEL International GmbH, Mörfelden-Walldorf
Director, Project Management, Real World Evidence

Dr Liana Iatridou-Oster

PRA Health Sciences, Mannheim
Director of Quality Assurance

Dr Carsten Schwenke

SCO:SSIS, Berlin
Statistical Consultant

Prof. Kurt Racké

Medizinische Fakultät der Universität Bonn, Bonn
Chairman of the Ethics Committees

Your Programme	Date (Starting 10:00 German time)	Your Expert
IT Systems in Clinical Trials: Risk Management and Validation	23 March 2020	Dr Andreas Jabs
Contracts & Agreements for Clinical Trials <ul style="list-style-type: none">Structure and content of contracts for beginners	1 April 2020	Alexander Maur
Investigational Medicinal Product: EU GMP Annex 13 Versus Regulation EU No 536/2014 <ul style="list-style-type: none">Present and future challenges for clinical investigational products	7 May 2020	Melanie Blessing
Practical Advice for Risk-based Monitoring	15 June 2020	Bettina Schopf
Clinical Data Publication – EMA Policy 0070 <ul style="list-style-type: none">Protected Personal Data (PPD) and Commercially Confidential Information (CCI)	7 July 2020	Dr Sybille M. Eibert
Data Protection in Clinical Trials: The Point of View of the Ethics Committee	10 August 2020	Prof. Sebastian Harder
GCP Audits and Inspections: What is Currently in Focus?	17 September 2020	Silja du Mont
Digital Data Generation – Using Apps und Wearables in Clinical Trials	6 October 2020	Karsten Sternickel
Trial Master File <ul style="list-style-type: none">Implications of the new EMA-guidelinesRequirements for electronic data archiving	25 November 2020	Dr Liana Iatridou-Oster
Update Statistics: Effects of the ICH E9 (R1) Addendum on the Study Design <ul style="list-style-type: none">Estimands and sensitivity analysis in clinical trials	8 December 2020	Dr Carsten Schwenke
Cooperation with Ethics Committees in Germany and Europe	22 January 2021	Prof. Kurt Racké
Safety Reporting in Clinical Trials <ul style="list-style-type: none">Reporting in clinical trials vs. non interventional studies (NIS)	5 February 2021	Angela Hartmann

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HOW TO REGISTER

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REGISTRATION

Yes, I want to join the

- PharmaFORUM Webcast Clinical Trials
(you will receive a confirmation email
with your login details)
- Yes, I agree that FORUM Institut may inform me
about events and relevant expert content by:
 email; and/or telephone.
I may withdraw my consent at any time.

Name

E-Mail (required for your login details)

Position

Company

Street address

Postal Code/City/Country

Tel. No.

Date, Signature

Fee:

Membership of the PharmaFORUM Webcast
Clinical Trials is available for one year.

The annual membership fee of € 1.800

(plus German VAT) for twelve webcasts is due upon
registration.

Membership is automatically extended by one year,
unless written notice has been submitted no later
than six weeks before the end of the membership.
A 12-month membership may be started at any time.

**If you are interested in a group account,
please contact us.**

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CANCELLATION POLICY

Our general terms and conditions apply
(as of 01.01.2016) and are available upon request.
We can send them to you anytime or you can find
them online at www.forum-institut.com/t&c

YOUR CONTACT



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