

DIA

Workshop for Excellence in Clinical Development

17-18 May 2017 | Hotel Media Rotana Dubai, UAE



PROGRAMME COMMITTEE

Päivi Itkonen

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Finland

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Chief Medical Writer, KAU Med,
Saudi Arabia

Holger G Adelmann

Senior VP and Managing Director,
DIA EMEA, Switzerland

| OVERVIEW

This workshop will introduce the modern key concepts for preparing and running state-of-the-art clinical trials in humans. A significant part is dedicated to understand the assessment criteria used by Pharma for making partnering decisions and on strategies to increase the attractiveness of clinical trial sites for sponsors. The workshop style will allow in-depth discussions with the expert instructors.

| KEY TOPICS

- Get knowledge about the mandatory regional and international guidelines and prerequisites for clinical investigations in humans.
- Learn how to interpret and apply modern concepts to govern safe human dose selection and dose escalation via concepts such as MTD (maximum tolerated dose) and MABEL (minimum anticipated biological effect level).
- Choose appropriate endpoints according to the trial purpose.
- Key concepts for data capture, data handling, data interpretation and reporting.
- Understand how Pharma companies assess and select their preferred providers.

| OBJECTIVES

This workshop will put you in a position to prepare and run clinical investigations in humans to the highest international standards. You will understand how to attract and build strong business relations with sponsors.

| WHO SHOULD ATTEND?

Clinical trial professionals such as medical doctors / investigators, study nurses, clinical research assistants, data managers, clinical project managers, quality leads, clinical trial site managers, clinical supply professionals, clinical pharmacists.

LEARN MORE

Visit www.DIAglobal.org
or contact EMEA@DIAglobal.org with questions.



Workshop for Excellence in Clinical Development

| DAY ONE | WEDNESDAY, 17 MAY

08:30 REGISTRATION AND WELCOME COFFEE

09:30 SESSION 1

INTERNATIONAL GUIDELINES FOR CLINICAL OPERATIONS & TRIALS IN HUMANS

Moderator: **Päivi Itkonen**, Managing Director, Crown CRO

- ICH
- Declaration of Helsinki / human research ethics
- Regional guidelines

Hamdi Akan, Professor, Ankara University; Chair Clinical Research Association, Turkey

11:00 COFFEE BREAK

11:30 SESSION 2

TRIAL DESIGN AND ENDPOINTS

Moderator: **Holger Adelman**, Senior Vice President and Managing Director, DIA Europe, Middle East and Africa

- Right endpoints for different purposes (safety, pharmacodynamics / biomarker, efficacy)
- Volunteer / patient selection & recruitment
- Regional practices
- International practices

Hamdi Akan, Professor, Ankara University; Chair Clinical Research Association, Turkey

13:00 LUNCH

14:30 SESSION 3

CLINICAL PROJECT MANAGEMENT & TRIAL CONDUCTION

Moderator: **Päivi Itkonen**, Managing Director, Crown CRO

- Solutions and best practices
- MABEL and MTD concepts, study termination criteria

16:00 COFFEE BREAK

16:30 SESSION 4

DISCUSSION ON HOT TOPICS WITH ALL ATTENDEES

Moderator: **Holger Adelman**, Senior Vice President and Managing Director, DIA Europe, Middle East and Africa

Anoud R. Omer, Chief Medical Writer, KAU Med, Saudi Arabia

17:30 NETWORKING RECEPTION

18:30 END OF DAY 1

| DAY TWO | THURSDAY, 18 MAY

08:00 REGISTRATION AND WELCOME COFFEE

09:00 SESSION 5

DATA COLLECTION, DATA MANAGEMENT AND ELECTRONIC DOCUMENT MANAGEMENT

Moderator: **Päivi Itkonen**, Managing Director, Crown CRO

- Data sources, handling and interpretation of data
- Regional recommendations
- International practices

10:30 COFFEE BREAK

11:00 SESSION 6

SAFETY AND PHARMACOVIGILANCE

Moderator: Representative invited

- Adverse Event definition & reporting
- Criteria for assessment of treatment relationship

Semra Sardas, Head of the Toxicology Department and Pharmacogenetics and Drug Safety Unit, Marmara University, Turkey

12:30 LUNCH

14:00 SESSION 7

PARTNERING WITH THE PHARMA INDUSTRY

Moderator: **Holger Adelman**, Senior Vice President and Managing Director, DIA Europe, Middle East and Africa

- Assessment criteria used by Pharma
- Strategies to increase attractiveness for sponsors

Betul Erdogan, Clinical Research Director, GCTO Turkey, Middle East, Egypt, MSD

15:30 END OF CONFERENCE

| Conference Venue

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Workshop for Excellence in Clinical Development

| Group Discounts

Register 3 individuals from the same company and receive a 50% discount for a 4th! All 4 individuals must register and prepay at the same time without exception. DIA will apply the value of the lowest applicable fee to this discounted registration; it does NOT include fees for optional events or DIA membership. You may substitute group participants of the same membership status at any time; however, administrative fees may be incurred.

Group registration is not available online and only available for the industry rate.

To take advantage of this offer, please print the registration form for each of the four registrants from your company. Include the names of all four group registrants on each of the forms and return them together to DIA.

For groups of 5 or more individuals, please contact Vishal.Bharadwaj@DIAglobal.org for a custom group rate.

DON'T MISS THESE UPCOMING OFFERINGS IN YOUR REGION

10-11 Apr 2017 | Dubai, United Arab Emirates | #17560

Building the eCTD - Practical Approaches to Compiling Electronic Submissions

This course will offer insight into the compilation of the eCTDs, share experience and best practice gained during eCTD submissions in the EU, and explain the eCTD review and lifecycle process. Especially eCTD submissions in the GCC region will be addressed in detail.

www.diaglobal.org/ectd

10-11 Apr 2017 | Dubai, United Arab Emirates | #17563

Practical Guide for Pharmacovigilance: Clinical Trials and Post-Marketing

The course walks you through the essentials of pharmacovigilance practices in pre- and post-marketing phases to enable you hit the ground running when implementing safety reporting. Regional developments in the GCC countries will also be highlighted.

www.diaglobal.org/PracticalPhV

20-21 November 2017 | Kuwait City, Kuwait | 17102

Middle East Regulatory Conference (MERC)



Early-bird discount available for members: Register by 5 April 2017

To qualify for the early-bird discount, registration form and accompanying payment must be received by the date above.

Early-bird fee applies to industry members only.

€ 1'230.00

CATEGORY	Member *	Non-Member*
Industry	€ 1'430.00 <input type="checkbox"/>	€ 1'585.00 <input type="checkbox"/>
Government/Charitable/Non-profit/Academia (Full-Time)	€ 715.00 <input type="checkbox"/>	€ 870.00 <input type="checkbox"/>

If DIA cannot verify your membership upon receipt of registration form, you will be charged the non-member fee. Group discount/SME rates available. Special rates for students and patient representatives on offer, subject to availability. Please contact DIA EMEA for more information.
Registration fee includes: refreshments, lunches, reception and meeting materials.

*All fees are subject to the applicable VAT. Payment due 30 days after registration and must be paid in full by commencement of the event.

TOTAL AMOUNT DUE: € _____

ATTENDEE DETAILS

PLEASE COMPLETE IN BLOCK CAPITAL LETTERS OR MAKE REGISTRATION EVEN SIMPLER BY ATTACHING THE ATTENDEE'S BUSINESS CARD HERE

Prof Dr Ms Mr

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First Name

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Please provide your European VAT number

PAYMENT METHODS

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Please charge my VISA MC AMEX

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Bank transfers: When DIA completes your registration, an email will be sent to the address on the registration form with instructions on how to complete the bank transfer. Payments in EURO should be addressed to "Account Holder: DIA." Please include your name, company, Event ID#17121 as well as the invoice number to ensure correct allocation of your payment.

Payments must be net of all charges and bank charges must be borne by the payer. **If you have not received your confirmation within five working days, please contact DIA Europe, Middle East and Africa.**

By signing below, I confirm that I agree with DIA's Terms and Conditions of booking. These are available from the office or online by clicking [here](#).

Date	Signature
<input type="text"/>	<input type="text"/>

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DIA offers one year complimentary membership against event registration at non-member rate

I do not want complimentary membership

TERMS AND CONDITIONS

Cancellations

All cancellations must be made in writing and be received at the DIA Europe, Middle East and Africa office four weeks prior to the event start date. Cancellations are subject to an administrative fee:

- Industry (Member/Non-member) € 200.00
- Academia/Charitable/Government/Non-profit (Full-time) (Member/Non-member) € 100.00

For cancellations after this date, or if the delegate fails to attend the meeting, no refund of fees will be given and be responsible for the full registration fee. DIA EMEA reserves the right to alter the venue and dates if necessary. If an event is cancelled, DIA EMEA is not responsible for airfare, hotel or other costs incurred by registered attendees. Registered attendees are responsible for cancelling their own hotel and travel reservations.

Transfer Policy

You may transfer your registration to a colleague prior to the start of the event but membership is not transferable. Substitute attendees will be responsible for the non-member fee, if applicable. Please notify the DIA EMEA office of any such substitutions as soon as possible.

Photography and Video Policy

By attending the event, you give permission for images of you, captured during the conference through video, photo, and/or digital camera, to be used by DIA in promotional materials, publications, and website and waive any and all rights including but not limited to compensation or ownership.

The DIA Europe, Middle East & Africa Contact Center will be pleased to assist you with your registration from Monday to Friday between 08:00 and 17:00 CET.

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