

H38 Essentials of Clinical Trial Monitoring

19th November 2019



The day will start at 08:45 with registration and coffee for a prompt start at 09:15. We aim to finish by 17:30.

ICR Training

H38 Essentials of Clinical Trial Monitoring





19th November 2019

H38

Venue - The Institute of Clinical Research Training Suite, White Waltham

The Course

This practical course defines the role of the clinical research monitor and explains the interaction with others in key roles as part of the clinical trial process as well as exploring the more complex aspects of clinical trial monitoring. Guidance will be given on best practices for selecting investigators, conducting monitoring visits and safety reporting. Delegates will explore strategies for dealing with common clinical trial management issues and how to adopt a preventative approach to handling monitoring issues. The course uses a blend of trainer input, exercises and delegate interaction.

Learning Objectives

- Define the roles and responsibilities of Monitors, Sponsors, Investigators and Ethics Committees
- Describe the key criteria for selecting investigators
- Summarise the preparation, conduct and follow-up of the major types of monitoring visit
- Define the requirements for safety reporting
- Prioritise tasks
- Apply risk based techniques to monitoring

Course Outline

- Refresher on the purpose of GCP
- Roles and responsibilities of monitors, investigators, & sponsors
- Investigator selection
- Preparing & conducting prestudy visits
- Preparing and conducting initiation visits
- Conducting monitoring visits
- SDV
- IMP
- Safety reporting
- Informed consent
- Monitoring visit reports
- Prioritising activities
- Preparing & planning close out visits
- Essential Documents

Who would benefit

This course is for all those involved in clinical research monitoring. It covers a wide range of topics and is designed for those who have been in the role for only a few months as well as the experienced CRA.

Course Fees

Guest £550.00 ICR Member £450.00 ICR Member Academic £350.00



Alison Messom

A molecular geneticist by training, Dr. Alison Messom has nearly 20 years of industry experience, working both within Pharmaceutical Companies and Clinical Research Organisations and is the current Chairman of the Board of Directors of the ICR.

With detailed experience of directing global clinical trials, Dr. Messom has worn many different hats during her career. She has had the opportunity to travel far and wide on business; lived in the UK, France and Switzerland, and gained operational experience in over 50 countries and line management in over 30.

During the course of her career she has line managed at all levels from CTAs through to Directors and VPs. At both i3 Research and ICON she helped build a new department of dedicated line managers for CRAs and also Start Up teams. While within the Pharma environment she line managed teams within Study and Project Management, Clinical Supplies, Translational Medicine, Data Management, Statistics, Informatics and IT – showing that line management skills are transferrable across functional areas.



Pre-Course Questionnaire - To be completed by all delegates

Please complete and sent to training@icr-global.org or fax to +44 01628 501 709

Course Title: H38 Essentials of Clinical Trial Monitoring

Date: 19 November 2019
Name:
Company / Hospital:
Position / Job Title:
How much experience of clinical trials do you have? (Years)
What are you hoping to get out of the day?
State one issue/problem you would like discussed at the meeting
Dietary Requirements

^{*} The ICR will work with the venue catering team and endeavour to accommodate specific dietary requirements e.g. Vegetarian/Vegan/Gluten Free - however it may not be possible to cover all requests for dietary preferences.



Address: The Institute of Clinical Research Training Suite, Suite 1, Cedar Court Grove Business Park White Waltham, Berkshire, SL6 3LW

The ICR's training suite is situated near the charming village of White Waltham. Located just off the A404, with connections by road or rail.

The Institute of Clinical Research Training Suite

Suite 1, Cedar Court

Grove Business Park

White Waltham

Maidenhead

SL6 3LW

By Car: Join the A404 either via at J5 of the M40, or J8/9 of the M4. Follow the A404 until you reach J9A signposted to Maidenhead/Reading. Turn right at the first roundabout, then continue on that road, going straight across any subsequent roundabouts. Take the second right past the airfield. If you reach the village, you've gone too far.

Once inside Grove Park, turn left at the first mini roundabout, then right at the T junction. Turn right at the red brick building, and we are in the one furthest from the road. Park in the large car park on your right. Please contact the Secretariat using the buzzer by the door - type in 12 and then press the small bell.

By Train: The nearest station is Maidenhead, located on the Paddington - Reading line. We recommend getting a taxi from the station, as bus routes are sporadic. It should take roughly 15 minutes.

Local Taxi Companies

Dot to Dot + 44 01628 200 999 Golden Executive + 44 1628 622955 Imperial +44 1628 771777

Accommodation

The ICR does not specifically recommend any accommodation - however the following are within easy travelling distance of the training venue

Hotel or Bed & Breakfast Name	Contact number	Website	Approximate Distance to The Institute of Clinical Research
Premier Inn	08715	http://www.premierinn.com	4 miles
Maidenhead	279520		14 minutes
Taplow	01628	https://taplowhousehotel.com	6 miles
House Hotel	670056		16 miles
Bel & The	01628 521263	https://belandthedragon-	7 miles
Dragon		cookham.co.uk/	18 minutes
CIM Moor	+44 1628	https://moorhall.cim.co.uk/	7 miles
Hall	427500		18 minutes

There are also a number of travel websites which may allow you to identify local accommodation and special offers - e.g. expedia

The Small Print

As a matter of policy we do not issue electronic copies of the slides used.

All ICR materials are copyrighted.

All delegates receive a delegate book.

Payment must be received in advance of a training course commencing. The ICR has the right to refuse entry for non-payment. Payment by invoice must be settled within 14 days from the date of invoice.

We understand that occasionally circumstances may change and that you will be unable to attend your chosen course. Notification of cancellation must be made in writing. If you cancel **more than 14 days prior to the event**, we will refund the course less £50 to cover administration costs. If you cancel within 14 days, no refund will be payable, but we will allow you to transfer to another course of your choice.

We will accept a change of delegate at any time without you incurring a penalty. The Institute of Clinical Research reserves the right to cancel any course that is under-subscribed but will give you 7 days notice in writing and will refund your course fees without any liability for any consequential or indirect loss.

At anytime, you may transfer to the same course within 12 months, or to another course of your choice within 6 months; a £25 administration fee will be charged for such transfers.

We may also need to change the venue but will give you 7 days notice in writing of the new location.

Programmes as published are correct, however due to circumstances beyond our control, trainers, speakers and/or the programme may need to be altered occasionally.

The ICR will work with the venue catering team and endeavour to accommodate specific dietary requirements e.g. Vegetarian/Vegan/Gluten Free – however it is not possible to cover all possible requests for dietary preferences.

Registration Form

Registration Form

Please photocopy this form for further registrations

Course Title:	H38 Essentials of Cl Monitoring	inical Trial	Course Da	ite:	19 November 2019		
Membership No.:		Title(Dr,Mr,Mrs,etc)):	First Name	2:		
Surname:			Job Title:				
Company Name:							
Email Address:							
Confirmation of booking will be sent by email, unless you request here that it is sent by post							
Correspondence A	ddress						
Address:							
Postcode:		Country:			Telephone Number:		
Special Dietary Requirements							
Declaration	and conditions of bool	king Signatur e	e:				
Method of Paymer		firmed when payment	has been re	caived (pla	ase tick as required)		
		illilled when payment	nas been re	cerved (pred	ase lick as required)		
I wish to pay the fee							
I enclose a chequ OR I wish to pay by	ue payable to "The Ins	titute of Clinical Resea	arch"				
VISA	MASTERCARD	☐ DELTA	EURO	CARD			
Card Number							
Start Date		Expiry Date					
Name (as it appears	on the card)						
Signature of card ho	older						
Please invoice m	y company using Purc	hase Order Number		Invoices	can only be raised when a PO no. is		
Correspondence A	ddress						
Address:							
Postcode:		Country:					