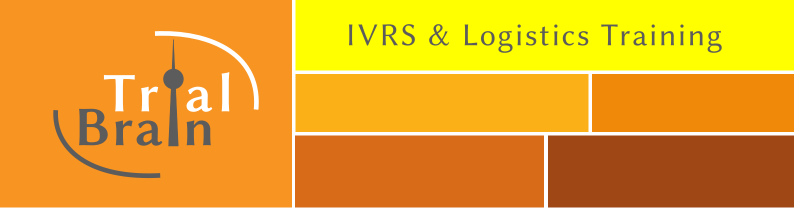
Seminar with practical Exercise

Understand and optimize IVRS User Requirements (IVRS-03)

10 & 11 April 2014, Berlin



Gain Understanding of IVRS User Requirements and their Impact on the Supply Chain!



Instructor

Nimer Yusef, PMP

Content

IVRS Parameters and their Significance

Characteristics of User Requirement Specifications for professional IVR(S)-Systems



Standardized IVRS-Glossary

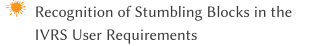
Optimization of IVRS User Requirements

Limited to 12 Attendees



Training Objectives

Training will provide you with This an Understanding from a logistical Perspective to set up a IVR(S)-System successfully.

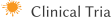


Understand and Communicate the Nature of IVRS User Requirements

Optimization of IVRS User Requirements by Avoidance of typical Errors

Who should attend?

The Training is designed to benefit Clinical Trial Professionals and Executives within Pharma Companies and CROs. The target audience is:



Clinical Trial Supply / Clinical Logistics



QA-Manager / Inspector / Auditor

- Consultant
- **IVRS-Expert**

Project Manager

Newsletter

If you are interested in being informed of further trainings, please subscribe to my Newsletter. (E-Mail: Newsletter@trial-brain.com).

Instructor

Nimer Yusef, PMP

is a Mathematician and certified Project Manager (PMP) with a strong background in Information Technology. His first experiences within the clinical research industry he gathered with the CRO ICRC Weyer more than 10 years ago. Subsequently, he worked for Perceptive Informatics, where he was the responsible Project Manager for IVRS-System Development with a focus in the area of Inventory Management.

As Technical Logistics Leader at Parexel Clinical Logistics Services he is responsible for training and consulting in IVRS-technology concepts since 2008. To close the competence gap between IVRS providers and end users in clinical study teams he founded the Company Trial Brain in 2010 to offer Seminars, Trainings and Workshops for IVRS and Clinical Logistics.

Date

Thu 10-Apr-2014, 09:00 AM to 05:00 PM Fri 11-Apr-2014, 08:00 AM to 04:00 PM

Location

Kempinski Hotel Bristol, Berlin near the Kurfuerstendamm and the City West

Costs

1.600 Euro + VAT per attendee (fee includes documentation and catering). Register 1 month in advance for a 10% rebate!

IVRS-03







Registration

Details

Fax: +49 / 30 / 81 45 84 99 E-Mail: events@trial-brain.com Online: at trial-brain.com

Registration

Registration for the Seminar "Understand and optimize IVRS User Requirements" (IVRS-03) 10 & 11-Apr-2014 in Berlin, Germany.

Vame	 	
Position		
Department		
Company	 	
Street	 	
Postal Code / City		
Country		
Phone		
E-Mail	 	
Notes		
Date & Signature		

Date & Time

Thu 10-Apr-2014, 09:00 AM to 05:00 PM Fri 11-Apr-2014, 08:00 AM to 04:00 PM

Location

Kempinski Hotel Briston Berlin Kurfuerstendamm 27 10719 Berlin, Germany Tel.: +49 /30 / 88 43 40

Accommodation

The hotel rooms are at the costs of 139,00 euro per night for participans of this seminar available. You can reserve a room at the hotel for the key-word "Trial Brain" until 10-Mar-2014.

Costs

1.600 Euro + VAT per Attendee (Fee includesDocumentation and Catering).Register 1 month in advance for a 10% rebate!

Cancellation by Registratants will be associated with the following Costs: - until 1 Month, 100 Euro + VAT - until 2 Weeks, 700 Euro + VAT A Substitution of a Registrant is possible anytime free of Charge.

Contact

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events@trial-brain.com +49 / 30 / 555 7777 9 www.trial-brain.com

IVRS-03