Impact of the New Pharmacovigilance Legislation on Regulatory Affairs

Event ID #13117 4-5 June 2013 Hotel NH Harrington Hall, London, UK



Programme Chair

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Head of Regulatory Affairs, European Medicine Agency, European Union

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Vice-President, Regulatory Affairs, Bristol-Myers Squibb Pharmaceuticals Ltd, France

More Programme Committee members to be invited

Continuing Education

DIA meetings and training courses are generally approved by the Commission for Professional Development (CPD) of the Swiss Association of Pharmaceutical Professionals (SwAPP) and the Swiss Society of Pharmaceutical Medicine (SGPM) and will be honoured with credits for pharmaceutical medicine. All participants are eligible for these credits and certificates are available.

Overview

This two day conference will focus on the impact of the new pharmacovigilance legislation on regulatory affairs, from various perspectives. The changes in legislation will affect the way product assessment is carried out in both pre- and post-authorisation phases, introduce new obligations to Marketing Authorisation Holders (MAH), and bring in continuous benefit risk assessment. This important conference will also be looking at the impact of the newly founded Pharmacovigilance Risk Assessment Committee (PRAC) on the life-cycle management of products and the PRAC's interactions with other committees.

Key Topics

The conference will provide

- An update on the implementation of the new pharmacovigilance legislation with a particular emphasis on the regulatory aspects. Information will be provided on the operation of the PRAC and the new Periodic Safety Update Report/Periodic Benefit-Risk Evaluation Report (PSUR/PBRER) and Risk Management Plan (RMP) requirements will be explored including how assessments will be handled by PRAC
- An overview of the key elements for Post-authorisation Safety Studies (PASS) and Post-authorisation Efficacy Studies (PAES) and sessions will review how the new pharmacovigilance legislation is impacting regulatory affairs and drug development more broadly.

Sessions will include

- Overview of PRAC and new pharmacovigilance requirements
- New pharmacovigilance legislation and how it is impacting drug development
- · Involvement of two sets of rapporteurs in the procedures and impact on regulatory affairs
- Referrals, opinions and conditions
- New pharmacovigilance legislation and the opportunities for regulatory affairs
- Panel discussion on impact on drug development and approval

Objectives

- To provide insight into the regulatory requirements, scientific and operational challenges associated with the implementation of the new pharmacovigilance legislation
- Attendance will offer opportunities to exchange experiences and hear from the regulators directly about how different aspects of the new legislation will be implemented

Who Will Attend

- This conference is aimed at intermediate and experienced professionals from
- Regulatory agencies
- The pharmaceutical industry and service providers
- Academic institutions
- including
- Regulatory affairs personnel
- Pharmacovigilance staff
- Quality assurance personnel for pharmacovigilance and pharmacovigilance inspectors
- Clinical and medical personnel
- Project managers in drug development

This conference is currently in development. Please visit www.diahome.org for regular programme updates or contact the Event Manager on Michael.Hediger@diaeurope.org



TUESDAY I 4 JUNE 2013

08:30	REGISTRATION AND WELCOME COFFEE		Session (
09:30	Session 1		Amendn Governm
09.30	OVERVIEW OF PRAC AND NEW PHARMACOVIGILANCE (PV) REQUIREMENTS (PART 1) Session Chairperson: Industry representative invited		PAES an Industry
	This session will cover overview of the PRAC including remit, membership, new PSUR and RMP requirements.	17:00	DRINKS
	PRAC – General overview including membership, working procedures and transparency Government representative invited	18:00	END OF
	New PSUR Requirements and Experience of Single Assessments/ Work sharing Procedure Government representative invited	WED	NESD
	RMP Requirements and How Assessments Will be Handled by the PRAC Government representative invited	08:30	Session S INVOLVE PROCED Session (
11:00	COFFEE BREAK		This ses rapporte
11:30	Session 2		Marketir
	OVERVIEW OF PRAC AND NEW PV REQUIREMENTS (PART 2) Session Chairperson: Industry representative invited		sets of r Governm
	This session will cover an overview of the key elements for PASS, PAES, annual review of conditions.		Post-Ap process
	Requirements for PASS Including Article 22 Joint Protocols and Registries Government representative invited		Governn Industry Industry
	PAES, Annual Review of Conditions and Renewal Government representative invited	10:00	COFFEE
	Monitoring (Black Symbol) and Change in Scope of Drugs Government representative invited	10:30	Session REFERR
13:00	LUNCH		Session
14.00	Consider 7		Referral Governn
14:00	Session 3 NEW PV LEGISLATION AND HOW IT IS IMPACTING DRUG DEVELOPMENT Session Chairperson: Industry representative invited		Article 2 Industry
	This session will discuss the impact of the new PV legislation on the drug development and pre-approval phase from various perspectives.		Quality o Regulati Governm
	How Does the New PV Legislation Impact the Drug Development and Pre-Approval Phase from the EMA Perspective? Government representative invited	12:00	LUNCH
	How Does the New PV Legislation Impact the Drug Development and Pre-Approval Phase from the Industry Perspective? Industry representative invited		a neutral,
	How Do these Elements Impact the Approval and Post-approval Phase from the Industry Perspective Industry representative invited	manag health) profession gement of p care prod

15:30 COFFEE BREAK

Unless otherwise disclosed, DIA Europe acknowledges that the statements made by speakers are their own opinion and not necessarily that of the organisation they represent, or that of the DIA Europe.

Speakers and agenda are subject to change without notice. Recording of any DIA Europe tutorial/workshop information in any type of media, is prohibited without prior written consent from DIA Europe.

16:00	Session 4		
	WHAT IS NEXT IN 2013?		
	Session Chairperson: Industry representative invited		

nent to the PV legislation and PV fees nent representative invited

d Future of Art 57 (2) representative invited

RECEPTION

DAY ONE

AY I 5 JUNE 2013

	INVOLVEMENT OF TWO SETS OF RAPPORTEURS IN THE PROCEDURES AND IMPACT ON REGULATORY AFFAIRS Session Chairperson: Industry representative invited
	This session will discuss procedural aspects involving two sets of rapporteurs.
	Marketing Authorisation Application (MAA) - Evaluation with two sets of rapporteurs; from pre-submission to opinion Government representative invited
	Post-Approval Evaluation with Two Sets of Rapporteurs: Role and process of interaction between the rapporteurs and committees Government representative invited
	Industry Experience: Evaluation with Two sets of rapporteurs
	Industry representative invited
00	Industry representative invited
00	
	COFFEE BREAK
	COFFEE BREAK Session 6 REFERRALS, OPINIONS AND CONDITIONS
	COFFEE BREAK Session 6 REFERRALS, OPINIONS AND CONDITIONS Session Chairperson: Industry representative invited Referral Procedures: Art 20, 31 and UUP (107i), public hearings
	COFFEE BREAK Session 6 REFERRALS, OPINIONS AND CONDITIONS Session Chairperson: Industry representative invited Referral Procedures: Art 20, 31 and UUP (107i), public hearings Government representative invited Article 20: Recent industry experience with procedures

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For more information, visit www.diahome.org or call DIA Europe +41 61 225 51 51.

3:00	Session 7	TRAVEL INFORMATION		
	NEW PV LEGISLATION AND THE OPPORTUNITIES FOR REGULATORY AFFAIRS			
	Session Chairperson: Government representative invited	From the airport:		
	This session will discuss the opportunities the new PV legislation provides and initiatives and status around adaptive licensing.	Heathrow: Take the Central Piccadilly Line to Gloucester Road station.		
	Adaptive Licensing: Overview of initiatives and status Government representative invited	Standsted: Take the Standsted Express to Liverpool Street. From there, take the Circle or District line to Gloucester Road station.Luton: Take the 757 bus to Victoria Station. From there, take the Circle or District tube line to Gloucester Road station.		
	Data Elements around Adaptive Licensing and Which Tools of the New PV Legislation Could be Used Government representative invited			
	Potential Pilot Projects – What could this look like?			
		London City: Take the DLR to Bank station. From there, take the Circle or		
4:30	COFFEE BREAK	District tube line to Gloucester Road station.		
5:00	PANEL DISCUSSION ON IMPACT ON DRUG DEVELOPMENT AND APPROVAL Panel Chair: Industry representative invited	Gatwick: Take the Gatwick Express to Victoria station. From there, take the Circle or District tube line to Gloucester Road station.		
	Panel will discuss impact of the new pharmacovigilance legislation on drug development, approval review and post-approval plans. Panellists have been invited			

16:00 END OF CONFERENCE

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HOTEL INFORMATION

The DIA has blocked a limited number of rooms at the following hotel:

NH Harrington Hall Hotel

5-25 Harrington Gardens South Kensington, London SW7 4JW UK Tel.: +44 207 396 96 96 Fax: +44 207 398 46 61

Email: bookings@nh-hotels.com

at the rate of: GBP 180.00 per room/night inclusive of breakfast and VAT.

To make your reservation, please contact the hotel directly at: bookings@nhhotels.com or by phone: +44 870 735 0358.

Please quote the booking reference: Group name: DIA Group code: 19429982

IMPORTANT: Please complete your reservation by 4 May 2013. Reservations received after this date will be subject to hotel availability and room rate may vary.

IN CASE OF CANCELLATION:

Cancellation of the hotel booking must be made in writing directly to the hotel. Cancellations made at least 7 days prior to arrival will not incur any cancellation charges. In case of no show or late cancellation the costs will be charged to the credit card of the guest.

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REGISTRATION FORM

Impact of the New Pharmacovigilance Legislation on Regulatory Affairs 4-5 June 2013 I Hotel NH Harrington Hall, London, UK



Early-bird rates available for members: Register by 23 April 2013							
Join DIA now to qualify for the Early-bird member fee! The Early-bird registration form and a Early-bird industry fee applies to industry members only. <i>(www.diahome.org/membership)</i>	accompanying payment must be re	ecceived by the date above. € 1'165.00 □					
FEES (after 23 April 2013) Industry		Member* Non-Member* € 1'365.00 □ € 1'480.00 □					
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	Registration fee includes: refresh	nments, lunches and meeting material.					
TOTAL AMOUNT DUE:	Payment is due 30 days after registration and must be paid in full by commencement of the event.						
ATTENDEE DETAILS	PAYMENT METHODS						
PLEASE COMPLETE IN BLOCK CAPITAL LETTERS OR ATTACH THE ATTENDEE'S BUSINESS CARD HERE	Credit cards: Payments by VISA, Mastercard or AMEX can be made by completing the details below. Please note that other types of credit card cannot be accepted.						
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*(Required for confirmation)	Date	Signature					
DIA reserves the right to include your name and affiliation on the attendee list.							

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All cancellations must be made in writing and be received at the DIA Europe office five working days prior to the event start date. Cancellations are subject to an administrative fee: • Full Meeting Cancellation: Industry (Member/Non-member) € 200.00.

- Academia/Charitable/Government /Non-profit (Full-Time) (Member/Non-member) € 100.00.
- Tutorial cancellation € 50.00.

If you do not cancel five working days prior to the event start date and do not attend, you will be responsible for the full registration fee. DIA Europe reserves the right to alter the venue and dates if necessary. If an event is cancelled or postponed, DIA Europe 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 Europe office of any such substitutions as soon as possible.

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