



International Pharmacovigilance Day

12th June 2019

from 13:15 to 17:50

13th June 2019

from 09:00 to 17:00

Sponsors

WITHIN AND WITHOUT BORDERS

PHARMACOVIGILANCE

Dialog. Solutions



⊙ | Barcelona, Spain

H10 Marina Barcelona

Avinguda del Bogatell, 64-68 - Barcelona, Spain

△ Adis Pharmacovigilance









A portion of the proceeds from this event are donated to the

"Vase of Flowers" project.



ABOUT Official language: English

PHARMACOVIGILANCE

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Yesterday, today and in the future: "Safety first"!

Are you ready to enter the transformed landscape in Pharmacovigilance?

Pharmacovigilance is at the center of the health ecosystem, made up of alliances, partnerships and business cross-contaminations. Therefore, our focus during this conference will be on the importance of collaboration and communication at local, regional and international levels, to ensure pharmacovigilance delivers its full benefits.

The world of medicines and device safety is rapidly changing. The role of patients is more important than ever but how do we bring patient insight into drug safety? How can we make safety information more accessible? How will the device regulation be implemented in the face of recent crises? How digital tools will change the face of PV? What new skills are needed to grow and become the Pharmacovigilance professionals of the future? Also, new global PV legislation is being issued at a fast rate from emerging markets, bringing new harmonization challenges and more regulatory authority inspections from brand new inspectorates. Is industry prepared?

The International Pharmacovigilance Day conference brings together senior leaders and technical experts to discuss the hot topics of today and tomorrow, share learnings and explore solutions. It promises to be a lively exchange of ideas, learning and concepts during a one day and a half, where the topics to be treated, among many others, include:

- How can Patient Support Programs and Risk Management become more patient centered?
- What could be Risk minimization measures and how to be compliant with the guality requirements?
- What opportunities and challenges does the EU Electronic health record bring to PV?
- Preparing for regulatory inspections with new global inspectorates and in post-Brexit?
- Digital Pharmacovigilance: What is hype and what is hot?
- EudraVigilance new approaches and new challenges
- How are companies expected to fulfil their requirements towards new regulations?
- How the future will look like, and what could be new skills for a new PV future?
- These are some of the questions that will make the future of PV, but how to be ready to embrace it?

We warmly welcome you to join us for this event of the future.



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Scientific Board



Jabeen Ahmad Regional Pharmacovigilance Director, Eastern Europe, Middle East & Africa (EEMEA) at AbbVie



Raffaele Di Marzo FU QPPV at Kedrion



Martin Holm-Petersen CEO at Insife





Marco Anelli Head of "Data, Information, Knowledge & Intelligence" Platform - PLG (Product Life Group)



Dr. Leonardo Ebeling Managing Director at Dr. Ebeling & Assoc. GmbH



Betina Østergaard Eriksen Vice President, Safety Surveillance at Novo Nordisk



Dr Solange Corriol-Rohou Sr. Director Regulatory Affairs & Policy, Europe at AstraZeneca

Who should attend?

This conference is designed to benefit functional/technical professionals working in the pharmaceutical and health care area dealing with the Pharmacovigilance system, such as:

- Safety and Pharmacovigilance dept.
- Clinical operation dept.
- Statistic dept.
- Medical Affairs dept.
- Medical Information dept.
- Regulatory Affairs dept.
- Quality & Compliance dept.
- Legal dept.
- Software Developing dept.
- Medical Devices Manifacuring Companies
- University Faculties scientists who are related to clinical and medical research (Senior, Associate and Assistant Professors, Research Scholars, Phd students).

Also, Directors/Seniors Directors/Executive Directors and Vice Presidents / Senior Vice Presidents/Exe cutive Vice Presidents and Heads/Leaders/Partners of: CROs and CMOs, Clinical Research Sites, Pharma, Biotech and Medical Devices Industries.



PHARMACOVIGILANCE WITHIN AND WITHOUT BORDERS

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Day 1

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from 13.15 to 17.50

from 09.00 to 17.00

13:15 - 13:45	Registration and Welcome coffee
13:45 - 14:00	Setting the scene with the new Regulations: CT Regulation
	Dr Solange Corriol-Rohou - Sr. Director Regulatory Affairs & Policy, Europe at AstraZeneca
14:00 - 14:40	The New MDR: Safety By Design And By Vigilance
	Jan Bart Hak - Head Medical Device Team at ProPharma Group
14:40 - 15:20	Structured Signal Management process as a tool to ensure patients' safety in clinical trials
	Mircea Ciuca - Global Head Medical & Clinical Drug Safety at Vifor Pharma
15:20 - 15:50	Coffee break
Signal Manag	gement using EVDAS
15:50 - 16:10	The EViSiMa® Project
10.00 10.10	Claudia Nowak - Pharmacovigilance Advisor at Dr. Ebeling & Assoc. GmbH
16:10 - 16:30	A company's experience
	Cathrine Lange - Safety Surveillance Principal Specialist at Novo Nordisk A/S
16:30 - 17:10	Safety in clinical trials and postmarketing in Europe: Major challenges
	Elena Prokofyeva - Head of Drug Safety Unit, Department of Research & Development, DG PRE at Federal Agency
	for Medicines and Health Products (FAMHP), Belgium
47.40 47.75	Panal Biassasian I Cosa musassing in Ell, quasial tanics for EV assas availability.
17:10 - 17:45	Panel Discussion Case processing in EU: special topics for EV cases availability
	CHAIR: Dr. Leonardo Ebeling - Managing Director at Dr. Ebeling & Assoc. GmbH
17:45 - 17:50	Concluding remarks Day One



Day 2

from 13.15 to 17.50

from 09.00 to 17.00

Official Language: English

09:00 - 09:10 **Start Day Two**

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Global	Pharma	acovidilan	ce Session

CHAIR: Jabeen Ahi	mad - Regional Pharmacovigilance Director, Eastern Europe, Middle East & Africa (EEMEA) at AbbVie
09:10 - 09:30	Pharmacovigilance in Pakistan Challenges and Opportunities
07:10 - 07:30	Pharmacovigilance in Pakistan, Challenges and Opportunities Farrukh Malik - Drug Safety & Regulatory Consultant at SYNTOF, France
	Turrant Hatte Drug Salety & Regulatory Consultant at STITTOT, Transec
09:30 - 09:50	Chinese FDA: Implementation of the ICSR guidelines and local language issue
	Jabeen Ahmad - Regional Pharmacovigilance Director, Eastern Europe, Middle East & Africa (EEMEA) at AbbVie
09:50 - 10:10	Pharmacovigilance in India – A Success Story
	Dr J Vijay Venkatraman - Managing Director & CEO at Oviya MedSafe Pvt Ltd
10:10 - 10:30	Pharmacovigilance in Latin America: challenges and synergistic partnerships between regulators, industry
	associations and marketing authorization holders towards a harmonized and robust PV system in the region
	Renata Yoshida Shihomatsu - Head of International Pharmacovigilance for Latin America at Roche (Hoffman-La Roche
10:30 - 10:50	Q&A session
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10:50 - 11:20	Coffee break
11:20 - 12:00	Pharmacovigilance Inspections what's new?
	Dr Ernesto Vera-Sanchez - Head of GCP and PHV Inspectorate at AEMPS - Spanish Medicines and Medical Devices Agency
12:00 - 12:40	Risk Minimisation Measures (RMM): How do we ensure that these measures are effective? Can real-world data help?
	Paola Nasuti - Associate Principal at IQVIA
12:40 - 13:40	Networking lunch
13:40 - 14:20	Al and Machine Learning literature monitoring: improve quality and save time!
	Mark Drinkwater - Co-founder of Pi2 Solutions at ProQuest
14:20 - 15:00	Electronic Product Information: How to change our communication with the patients?
	Dr Rüdiger Faust - Regulatory Strategy & Intelligence Lead, Innovation Unit Devices & Technologies at Grünenthal GmbH
15:00 - 15:30	Coffee break
15:30 - 16:10	WEB-RADR: App and Patient Support Programs experience
	Jabeen Ahmad - Regional Pharmacovigilance Director, Eastern Europe, Middle East & Africa (EEMEA) at AbbVie
	Phil Tregunno - Vigilance, Intelligence and Research Group Manager at MHRA
16:10 - 16:40	Al in PV: lesson learned in case studies
	Phil Tregunno - Vigilance, Intelligence and Research Group Manager at MHRA
16:40 - 17:00	Digital Media: A Pharmacovigilance Challenge
	Sonia López - Qualified Person Responsible for Pharmacovigilance at Asphalion
	Nuria Cabello - Senior Drug Safety Officer at Asphalion
17:00	Concluding remarks

HOW TO REACH THE CONFERENCE VENUE



H10 Marina Barcelona

Avinguda del Bogatell, 64-68 - Barcelona, Spain

By taxi

A taxi from the airport should take about 25 minutes.

By train

The RENFE train leaves the airport every half an hour and will take you to Passeig de Gràcia station, in about 25 minutes. Then you have to change at Passeig de Gràcia for the Yellow Metro for Bogatell and your hotel is 3 minute walk from the metro stop.

By bus

The Aerobus will also take you from the Airport to Catalunya station. The journey time is around 30 minutes. From Catalunya you need to change for the Green line Metro to Passeig de Gràcia; at Passeig de Gràcia you need to change again for the for the Yellow Metro for Bogatell and your hotel is 3 minute walk from the metro stop.

By tube

The closer metro station is Bogatell.







For further information

Please visit the conference website or contact the organisational offices:

Ilaria Butta Phone: +39 (0)35.4123594 | ilaria.butta@lsacademy.com



△ Adis Pharmacovigilance

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Two ways to register

Online

please register here

https://internationalpharmacovigilanceday2019.lsacademyevents.it/orders/new

In this case, you can choose to pay by credit card or by bank transfer

By email o fax

please fill the registration form below for each attendee and send it by **email**: ilaria.butta@lsacademy.com or by **fax**: +39(0)35.4501262 In this case, you can pay by bank transfer.

- € 1090,00 Early Bird fee extended until May 31st, 2019
- **€ 1250,00** Ordinary fee (after May 31st, 2019)
- € 890,00* Patronage members fee (Association ______)
- € 630,00* Academy, Public Administration, Freelance
- € 300,00* Asphalion reserved fee

The fee includes: seat at the conference, copy of presentations of Speakers who allow the distribution, informative literature for the day, welcome coffee, networking lunch, coffee break, organisational office assistance, certificate of attendance. *For Italian companies: + 22% VAT

For any additional information, please contact:

Ilaria Butta | Ilaria.butta@LSacademy.com Phone: +39 (0)35.4123594

Payment by bank transfer

The full amount must be paid on registration to EasyB S.r.l by bank transfer. If you pay by bank transfer, please attach proof of payment to the registration form.

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Date _

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