

EUROPEAN
PHARMACO
VIGILANCE
CONGRESS

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#EUPV2020

ONLINE

November 26 - 27

European Pharmacovigilance Congress 2020

PEC PHARMA
EDUCATION
CENTER



ABOUT

The European Pharmacovigilance Congress organized by Pharma Education Center is the international meeting venue of experts from marketing authorization holders, competent authorities, international pharmacovigilance organizations and European expert patient organizations. The Congress, now at its 4th edition, is a unique opportunity to get insight from different PV stakeholders, to learn more about new requirements, new emerging challenges in Pharmacovigilance and to discuss possible strategies and solutions to address them.

WHY ATTEND? WHY ATTEND?

- Learn about the latest PV trends and updates
- Share experiences and solutions at a leading european event
- Forge new collaborations with key decision makers
- Networking through the virtual platform with Pharma Companies delegates and Speakers

WHO SHOULD ATTEND?

Pharma, Biotech and Medical Devices Industries
Pharmacovigilance Associations
Regulatory Bodies
CROs and CMOs
Clinical Research Sites
Data Management Companies
Software Development Companies
University Faculties
PV Consultant Societies

What can you expect at the EUPV2020?

- Oral talks & keynote presentations
- Round Tables involving speakers and participants
- Poster session in the exhibition area. Selected posters abstract will be published by Sage.
- Exhibition area for Emerging Technologies and PV solutions
- Publication by Sage of the congress abstracts

EUPV CONGRESS is more than a traditional conference, it is a great opportunity to develop your professional skills and get in touch with renowned experts

FEATURED TOPICS

UPDATES FROM INTERNATIONAL PHARMACOVIGILANCE ORGANIZATIONS

SIGNAL DETECTION & EVALUATION

DIGITAL & TECHNOLOGY INNOVATION IN PV: OPPORTUNITIES FOR PATIENTS

DRUG INDUCED LIVER INJURY (DILI)

CLINICAL TRIALS REGULATION

COMUNICATION IN DRUG SAFETY

PHARMACOEPIDEMOLOGY & RISK MANAGEMENT

EUDRAVIGILANCE/EVDAS UPDATES

PV QUALITY SYSTEM

LOCAL VS GLOBAL PV REGULATIONS

#EUPV2020

AGENDA 26th

- 09:10 **Welcome and Opening remarks by the Chairperson of the European Pharmacovigilance Congress**
Marco Sardella - Chairperson of the EUPV congress
Chief Pharmacovigilance Officer & EU QPPV ADIENNE Pharma & Biotech
- 09:20 **A word from the Editor-in-Chief of Therapeutic Advances in Drug Safety - Eu PV Congress Media Partner**
Arduino Mangoni
Professor and Head, Dep. of Clinical Pharmacology, School of Medicine, Flinders University, Australia

1. UPDATES FROM INTERNATIONAL PHARMACOVIGILANCE ORGANIZATIONS

*Chairperson: **Marco Sardella** - Chairperson of the EUPV congress*

- 09:30 **Updates from the CIOMS**
Hervé Le Louët - (tbc)
President of CIOMS, member of PRAC
- 09:50 **Updates from ISoP**
Jan Petracek
ISoP Advisory Board at International Society of Pharmacovigilance
- 10:10 **Updates from the Uppsala Monitoring Centre**
Daniele Sartori
*Pharmacovigilance Scientist- Uppsala Monitoring Centre
DPhil student in Evidence-Based Medicine, University of Oxford*
- 10:30 **Updates from PIPA**
Sarah Hall
HonFPIPA, Managing Director MIPSOL
- 10:50 **Q&A**
- 11:00 **Coffee Break e networking**

2. SIGNAL DETECTION & EVALUATION

*Chairperson: **Fabio De Gregorio** - V.P., Head of Drug Safety Europe, UK QPPV at Shionogi Europe*

- 11:20 **Signal detection and dissemination to members of the WHO Programme for International Drug Monitoring**
Daniele Sartori
*Pharmacovigilance Scientist- Uppsala Monitoring Centre
DPhil student in Evidence-Based Medicine, University of Oxford*
- 11:40 **Translating Pre- Marketing Adverse Drug Reactions and Signal Evaluation To The Routine Management of The Older Patient: Challenges and Opportunities**
Arduino Mangoni
Professor and Head, Dep. Of Clinical Pharmacology, School of Medicine, Flinders University, Australia
- 12:00 **Post marketing Signal detection - case studies**
Glyn Belcher
CEO of PV Consultancy Ltd

- 12:20 **Round table & Q&A Time**
Moderator: **F. De Gregorio**, *Shionogi Europe*
Participants: **G. Belcher**, *PV Consultancy Ltd*
M. Ciuca, *Area Head - Global Clinical Safety and Pharmacovigilance at CSL Behring*
A. Mangoni, *Flinders University, Australia*
D. Sartori, *Uppsala Monitoring Centre*
D. Stenver, *Indep. PV Adviser Founder of Unique Advice*

12:50 **Lunch & Networking**

3. DIGITAL & TECHNOLOGY INNOVATION IN PV: OPPORTUNITIES FOR PATIENTS

Chairperson: Valentina Mancini, *Director PV, EU QPPV - Shionogi Europe*

- 13:50 **Intelligent automation in Pharmacovigilance**
Juergen Schmider
President Drug and Device Vigilance Consulting LLC - Arisglobal

- 14:10 **Round table & Q&A Time**
Real World Evidence in PV, with digital tools for self-reporting by patients
Moderator: **V. Mancini**, *Shionogi Europe*
Participants: **S. Cazzaniga**, *Medical Affaire Excellence Head Janssen-Cilag Italy*
S. Hall, *HonFPIPA, Managing Director MIPSOL*
P. Kruger, *Expert patient Eupati*
J.Schmider, *Arisglobal*
B. Van Leeuwen, *Deputy-QPPV - Astellas Pharma Europe*

4. DRUG INDUCED LIVER INJURY (DILI)

*Chairperson: **Furlan Giovanni**, Safety Risk Lead, Director - Pfizer*

- 14:40 **Drug Induced liver injury: from pre-clinical to post-marketing studies**
Marco Tuccori
Unit of Adverse Drug Reactions Monitoring, University Hospital of Pisa
- 15:00 **Hervé Le Louët - (tbc)**
President of CIOMS, member of PRAC
- 15:20 **Nimesulide Case Study: an example of Benefit-Risk Assessment and EU Referral Procedures**
Mario Bertazzoli
Director, Group Head of Drug Safety and Reference Physician to EU QPPV at Helsinn Healthcare SA
- 15:40 **Round table & Q&A Time**
Moderator: **G. Furlan**, Pfizer
Participants: **M. Bertazzoli**, Helsinn Healthcare SA
H. Le Louët, CIOMS
M. Tuccori, University Hospital of Pisa
- 16:00 **Coffee break & networking**

5. CLINICAL TRIALS REGULATION

*Chairperson: **Mircea Ciuca**, Global Therapeutic Area Head - Global Clinical Safety and Pharmacovigilance at CSL Behring*

- 16:20 **CTFG Q&A RSI and CTR Q&A documents: what will change in terms of safety in clinical trials when CTR is in place?**
Elena Prokofyeva
*Head of drug safety Unit, DG PRE Authorization/Division R&D (Human)
Federal agency for medicines and health products - Bruxelles*
- 16:40 **Round table & Q&A Time**
Moderator: **M. Ciuca**, CSL Behring
Participants: **E. Di Martino**, Scientific Director PHARMA D&S
M. Forstner, Senior Director, Pharmacovigilance & Patient Safety - PRA Health Science
E. Prokofyeva, Head of drug safety Unit - AFMPS
A. Traversa, Independent PV Expert
- 17:10 **Marco Sardella - Chairperson's closing remarks**
Chairperson of the EUPV congress

AGENDA 27th

- 09:00 **Welcome and Opening remarks by the Chairperson of the European Pharmacovigilance Congress**
Marco Sardella - Chairperson of the EUPV congress
Chief Pharmacovigilance Officer & EU QPPV ADIENNE Pharma & Biotech
- 09:10 **A word from the Editor-in-Chief of Therapeutic Advances in Drug Safety**
Arduino Mangoni
Professor and Head, Dep. of Clinical Pharmacology, School of Medicine, Flinders University, Australia

6. COMMUNICATION IN DRUG SAFETY

Chairperson: Michael Forstner, Senior Director, Pharmacovigilance & Patient Safety- PRA Health Science

- 09:20 **Labelling**
Barbara De Bernardi
Vice President, EU QPPV - Pfizer
- 09:40 **Paradoxical effects of communicating information on adverse drug reactions**
Giovanni Furlan,
Safety Risk Lead Director - Pfizer
- 10:00 **Q&A**

7. PHARMACOEPIDEMIOLOGY & RISK MANAGEMENT

Chairperson: Doris Stenver, Indep. PV Adviser, Founder of Unique Advice

- 10:10 **RMP weaknesses and their evolution - Effectiveness of Risk minimization actions**
Jan Petracek
ISoP Advisory Board at International Society of Pharmacovigilance

10:30 **Systematic Lifecycle Benefit-Risk Management and Decision Making**
Michael Forstner
Senior Director, Pharmacovigilance & Patient Safety - PRA Health Science

10:50 **Coffee Break e networking**

11:15 **Round table & Q&A Time**
Moderator: **D. Stenver**, *Unique Advice*
Participants: **G. Belcher**, *PV Consultancy Ltd*
F. De Gregorio, *V.P. Head of Drug Safety Europe, UK QPPV - Shionogi Europe*
M. Forstner, *Senior Director, PV & Patient Safety PRA Health Science*
J. Petracek, *Board Member ISoP*
P. Rotunno, *Pharmacovigilance Consultant*

8. EUDRAVIGILANCE/EVDAS UPDATES

*Chairperson: **Francoise Sillan**, VP Head of Therapeutic area oncology - endocrinology Global patient safety, Ipsen*

11:45 **EVDAS-ISO IDMP- E2B R3**
Calin Lungu
MD, MRQA, BCPM, Eudravigilance and XEVMPD Trainer - EMA, CEO - DDCS

12:15 **Round table & Q&A Time**
Moderator: **F Sillan**, *Ipsen*
Participants: **L. Boga**, *QPPV - Dompé Pharmaceutical*
M. D'Antuono, *Corporate PV Director, EU QPPV- Italfarmaco*
C. Lungu, *CEO DDCS*
J.A. Ayala Ortiz, *PVpharm CEO*

12:45 **Lunch & networking**

9. PV QUALITY SYSTEM

*Chairperson: **Fernanda Ferrazin**, Independent Pharmaceutical Consultant, EuPV congress Scientific Board - PEC*

13:50 **PV System Inspection Readiness**
Raj Bhogal
Sr. Director, R&D Audits & Inspections - Jazz Pharmaceuticals

14:10 **Round table & Q&A Time**
Moderator: **Fernanda Ferrazin**, *Scientific Board - PEC*
Participants: **R. Bhogal, Sr. Director**, *R&D Audits & Inspections*
Jazz Pharmaceuticals
GN. Castiglione, *EU QPPV & Director Global*
Pharmacovigilance - Chiesi
I. Grisoni, *Sr. Director, EEA QPPV - Jazz Pharmaceuticals*
D. Marcozzi, *Head of R&D QA - Fidia Farmaceutici*

10. LOCAL VS GLOBAL PV REGULATIONS

*Chairperson: **Valentina Mancini**, Director PV, EU QPPV - Shionogi Europe*

14:40 **Is the limelight of PV regulation moving back to national?
How to maintain global PV compliance with different/conflicting national regulations?**
Bert. Van Leeuwen
Deputy-QPPV - Astellas Pharma Europe

15:00 **Getting the most from Patient Support Programmes—from a Safety Perspective**
Phillip Eichorn
Senior director, Pfizer Pharmaceuticals

15:20 **Management of global PSMF: compliance with local requirements**
Margherita D'Antuono
Corporate PV Director, EU QPPV - Italfarmaco

15:40 **Round table & Q&A Time**
Moderator: **V. Mancini**, *Shionogi Europe*
Participants: **J.A. Ayala Ortiz**, *PVpharm CEO*
M. D'Antuono, *Italfarmaco*
P. Eichorn, *Pfizer Pharmaceuticals*
I. Grisoni, *Sr. Director, EEA QPPV - Jazz Pharmaceuticals*
S. Hall, *HonFPIPA, Managing Director MIPSOL*
B. Van Leewen, *Deputy QPPV - Astellas Pharma Europe*

16:10 **End Closure Remarks**
Marco Sardella
Chairperson of the EUPV congress

16:20 **End of Congress**

SPEAKERS



José Alberto Ayala Ortiz
PVpharm CEO, EU QPPV, PV
Consultant, LCPV Services, Spain



Glyn Belcher
CEO of PV Consultancy Ltd



Mario Bertazzoli
M.D., Director, Group Head of
Drug Safety and Reference Physi-
cian to EU QPPV at Helsinn Heal-
thcare SA



Raj Bhogal
Sr. Director, R&D Audits &
Inspections - Jazz Pharmaceuticals



Laura Boga
QPPV - Dompé Pharmaceutical Spa



Gian Nicola Castiglione
M.D., EU QPPV & Director Global
PV - Chiesi Farmaceutici



Sara Cazzaniga
Medical Affairs Excellence Head
at Janssen-Cilag Italy



Mircea Ciuca
Global Therapeutic Area Head in
Global Clinical Safety and PV at
CSL Behring - Scientific Board
EUPV congress



Margherita D'Antuono
PharmD, PhD, Corporate PV Direc-
tor, EU QPPV, Italfarmaco



Barbara De Bernardi
Vice President, EU QPPV Deputy,
Pfizer



Fabio De Gregorio
M.D., Vice President, Head of
Drug Safety Europe,
Shionogi Europe



Elisabetta Di Martino
Scientific Director PHARMA D&S



Philip Eichorn
M.D., Senior Director Worldwide
Safety & Regulatory - Pfizer Phar-
maceuticals



Fernanda Ferrazin
Former Head of Pharmacovigilance
Dept. - Italian Medicines Agency -
Scientific Board PEC (Pharma Edu-
cation Center)



Michael Von Forstner
PhD, Senior Director, Pharmacovigi-
lance & Patient Safety - PRA Health
Science



Giovanni Furlan
Pharm.D., Safety Risk Lead,
Director, Pfizer -Scientific Board
EUPV congress



Sarah Hall
PhD, HonFPIPA, Managing Direc-
tor MIPSOL



Ilaria Grisoni
Sr. Director, EEA QPPV, Jazz
Pharmaceutical



Paola Kruger
Expert Patient, EUPATI (European
Patient's Academy for Therapeutic
Innovation)



Hervé Le Louët
M.D., PhD, President of CIOMS,
member of PRAC (tbc)



Calin A. Lungu
MD, MRQA, BCPM, Eudravigilance
and XEVMPD Trainer - EMA,
CEO - DDCS



Valentina Mancini
Director PV, EU QPPV, Shionogi
Europe - Scientific Board EUPV
congress



Arduino Mangoni
MBBS, MD (Hons), PhD, Professor
and Head, Department of Clinical
Pharmacology, School of Medicine,
Flinders University, Australia



Daniela Marcozzi
Head of R&D QA Company
Representative for Competent
Health Authorities - Fidia Farma-
ceutici S.p.A.



Jan Petracek
MD. PhD, ISoP Advisory Board at
International Society of Pharmaco-
vigilance



Elena Prokofyeva
MD, MPH, PhD, Head of Drug
Safety Unit - AFMPS



Patrizia Rotunno
PV Consultant, Scientific Board
EUPV congress



Marco Sardella
Chief Pharmacovigilance Officer ,
EU QPPV ADIENNE Pharma &
Biotech, Chairperson for EU PV
Congress



Daniele Sartori
MSc Pharm, Pharmacovigilance
Scientist- Uppsala Monitoring
Centre



Schmider Juergen Schmider
M.D., Ph. D., President Drug and
Device Vigilance Consulting LLC -
Arisglobal



Françoise Dumas Sillan
M.D., VP Head of Therapeutic area
oncology - endocrinology Global
patient safety, Ipsen



Doris Irene Stenver
D.M., MPA, Indep. PV Adviser
Founder of Unique Advice, former
Chief Medical Officer Danish
Medicines Agency and former
member of PRAC



Alessandra Traversa
Independent Pharmacovigilance
Expert



Marco Tuccori
Drug Safety Manager at University
Hospital of Pisa



Bert Van Leeuwen
M.D., Deputy-QPPV of Astellas
Pharma Europe

CONGRESS MEDIA PARTNER



Published by SAGE, Therapeutic Advances in Drug Safety (Impact Factor: 3.463) is an international peer-reviewed Open Access journal, delivering the highest quality original research articles, reviews, and scholarly comment on pioneering efforts and innovative studies pertaining to the safe use of drugs in patients. The journal has a strong clinical and pharmacological focus and is aimed at an international audience of clinicians and researchers in drug safety, providing an online forum for rapid dissemination of recent research and perspectives in this area.

As the official Media Partner of the 4th edition of the European Pharmacovigilance Congress, Therapeutic Advances in Drug

Safety will be publishing an **online abstract supplement** which will be free to access online.

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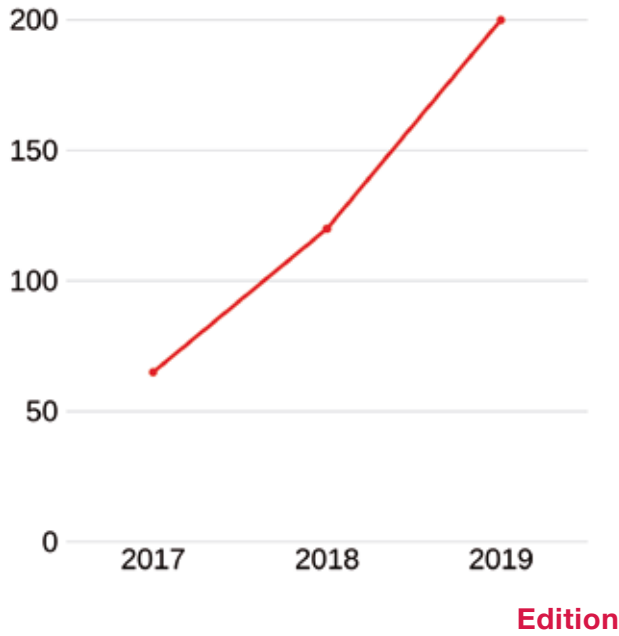
The European Pharmacovigilance Congress 2020 provides an excellent opportunity to get in touch with PV professionals and decision makers through a powerful virtual exhibition area.

To sponsor the event, contact us at: info@pharmaeducationcenter.it



ATTENDANCE

Number of participants



Last edition

200
Participants

78
Pharma
companies

34
Speakers



ONLINE CONGRESS



HOW TO REGISTER

ENTRY FEE

650 € early bird before September 30th

750 € early bird before October 31th

950 € full price

VAT not included

TERMS OF PAYMENT

Advance payment is required with respect to the event date by bank transfer to BANCO BPM Spa - Florence (Italy),
IBAN: IT90U0503402815000000001400,
Bic/SWIFT: BAPPIT21G74 made payable to Pharma Education Center S.r.l., Via dei Pratoni 16, 50018 Scandicci (FI), VAT number 02173670486, specifying event title along with name and surname of the participant enrolling. Attendance to the event will be allowed upon payment received.

Invoice of payment will be issued after the second half of the month after the course.

For further information and/or further assistance please contact (+39) 055 7224179 or email: amministrazione@pharmaeducationcenter.it

COURSE/EVENT CANCELLATION

If the minimum number of participants is not reached, Pharma Education Center reserves the right to cancel or schedule the event for another date.

Formal communication will be given within 5 days before the event date.

In this case Pharma Education Center will refund the registration fee in full and without additional charges. Alternatively, the participant can request a spendable coupon for participating in another PEC event scheduled in the current year.

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HOW TO REGISTER

Please, fill the form on the web site
<https://www.eupharmacovigilance.com/>

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