"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK



AGENDA AT A GLANCE

**Key Speakers Include** 



NICHOLAS CALL Special Agent FDA



SHAANTANU DONDE
Senior Director, Medical Portfolio Development
(Developed Markets), Upjohn, Division of Pfizer



SUSAN WELSH Chief Safety Officer CSL Behring



SCOTT CHANDLER

Vice President & Global Head Licensing and Early Development (LEAD) Safety, Roche



**JOHN SOLOMON** 

Head of Pharmacovigilance - UK & Ireland Sanofi



MICHAEL BEAN

Senior Director, Regulatory Compliance R&D Johnson & Johnson



TANJA PETERS

Senior Pharmacovigilance Expert, Deputy EU-QP-PV & Head PV Intelligence, Boehringer Ingelheim



**JABEEN AHMAD** 

Global PV Consultant Independent Consultancy



WIVINA DE WAELE

Director, Regional Safety Excellence EMEA.Global Drug Safety, Alexion



STEINAR MADSEN

Medical Director Norwegian Medicines Agency



RAJ BHOGAL

Head of Inspection, R&D Quality Takeda



PHILIP OLUWOLE

Associate Principal Surveillance Specialist Astrazeneca



MATE A. BALAZS

Country Head - Patient Safety - Hungary Novartis



**MIRCEA CIUCA** 

Global Therapeutic Area Head - Global Clinical Safety and Pharmacovigilance, CSL Behring



**SUMIT MUNJAL** 

Senior Medical Director, Head Europe PV Takeda Pharmaceuticals



JOHN POUSTIE

Medical Director & EU QPPV, Global PV Norgine



VALENTINA MANCINI

Director PV, EU QPPV Shionogi Europe



YUUNG YUUNG YAP

Senior International Regulatory Counsel, EU and International Regulatory Law, Pfizer



LUIZ LIMA

Senior Global Patient Safety Physician Neurology, Ipsen



FRANCK SCHWARTZ

QA Global Inspection, Intelligence Lead -Compliance and Regulatory Affairs Quality Novartis





"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK "I found it to be very well structured, focused on topics of interest for every PV professional. All the speakers were amazing and I look forward attending your future conferences"

PhV Manager, Bausch Health

AGENDA AT A GLANCE

## **Key Speakers Include**



KAREN CHENG Safety Risk Lead Pfizer



IVA SLAVCEVOVA
Deputy QP Pharmacovigilance/Global Patient
Safety, Baxter



ANDREA OLIVA Head of Pharmacovigilance Mylan



NATALIE SPRINGVELD Global Safety Leader Bayer



TEA BABIC Associate Director, Audits and Inspections,



KARE KEMP
Team leader (senior advisor), Signals and risk
management - PV/Safety
Danish Medicines Agency

Global Pharmacovigilance Compliance, Teva



ALINA TUDOR Associate Director, Senior PV Physician/ Deputy EU QPPV, Norgine



YVONNE NANCIU
Senior Manager PV & Medical Information,
Local QPPV, Abbvie



MARJAN DZEPAROSKI Head of Regulatory Affairs, Drug Safety & Intelectual Property, Bionika Pharmaceuticals



CHETAN SHATAPATHY
Principal Pharamcovigilance Physician Oncology R&D Unit, AstraZeneca



BARBARA DE BERNARDI Deputy EU QPPV and European Safety Office Head, Pfizer



SALVATORE GIORGIO CICIRELLO Senior Director Safety Science & PASS, Global Drug Safety & Risk Management Celgene



NICOLE BAKER Co-Founder BioLogit



MADDALENA LINO Terapheutic Area Safety Head Segirus



MARY LYNNE VAN POELGEEST President, World Federation for Incontinent Patients - (WFIP)



SANDY EISEN Chief Medical Officer Frontline Pharma Consulting



GEORGIA GAVRIILIDOU Counsel Sidley Austin

Plus many more COMING SOON.....

"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK

"Conference was very informative & added much knowledge about Pharmacovigilance systems, ADE, process flow of reporting, searching data & mobile networking"

Asst. Manager Regulatory Affairs, Emcure Pharmaceuticals

## WHO ATTENDS? 21st Pharmacovigilance 2020 70% 70% 50+ 6+ 02 01 Pharma & Golden Speakers Networking Days **Decision Makers** Biotech Opportunity Hours

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"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK "Panel discussions are very interactive as well as address real world and practical issues"

Head - Medical Affairs, Wockhardt

## AGENDA AT A GLANCE

## **OUR HISTORY**

Virtue Insight (VI) started it's wonderful journey in 2009 and now after a decade in the industry, we are honored to organise our 21st event in Pharmacovigilance to be held in 2020 in UK.

Over our past events, we have gained huge trust of our industry partners through our ability of providing best connect within the pharma regulators, stakeholders and the patients. Keeping our promise through all our 20 PV events in 3 different regions (UK, USA and India), our upcoming event holds the same intensity with newer challenges in PV along with new techniques to ease the process.

Our events have grown tremendously over the years within the pharma market, which lead to return of all our major clients every year to showcase their facilities to our senior level participants that help many to enhance their skills of critical drug safety evaluation process. We have been constantly rebuilding our content, format and agenda topics to stand ahead of what market demands.

This year, our event keeps an immersed eye on discussion of critical topics in PV domain, which capture influence of emerging technologies like AI, IoT, Big Data. Not merely that, we have exciting surprise activities which will help you to interact more with your peers. This will be surely an exciting event wherein you could get chance to meet big industry gems. Let's gather to shape the industry with your magnificent ideas.

UK is waiting for you!!!

## **MAJOR FOCUS ON**

## ENSURING PATIENT SAFETY



### WHO WILL YOU MEET

CEO's, CTO's, CIO's, Presidents, VPs, Directors, Heads, Managers, Scientific Advisors, Consultants of:

Pharmacovigilance, Pharmacoepidemiology, Pharmacogenomics, Drug/Product Safety, Drug Development, Information and Clinical Data Management, Clinical Pharmacology, Clinical Safety, RMPs, PSURs, PADERs, PBRERs, Risk Management, Research & Development, Quality Assurance, Patient Safety, Signal Detection, Safety Surveillance, Outcomes Research, Data Analysis, Epidemiology, Medical Affairs, Regulatory Affairs & Compliance, Information technology, Sales and Marketing

"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK "Very good platform to meet other pharmacovigilance expertise and interact with them about the advances & opportunites in pharmacovigilance. Virtue Insights is really good at coordinating and organizing"

Safety Physician, Sciformix

## AGENDA AT A GLANCE

## DAY ONE - 26th February 2020

08:30 - Coffee and Registration - An opportunity to meet and to network with your conference colleagues.  09:20 - Chairperson's opening remarks  SUSAN WELSH Chief Safety Officer CSL Behring	<ul> <li>Outsourcing in Pharmacovigilance- Best Practices, Challenges and key consideration</li> <li>Pharmacy practice and its guidelines</li> <li>Future Drivers for Pharmacovigilance</li> <li>New ways to generate evidence including real world evidence</li> <li>Proper communication - Sponsor - Site - CRO &amp; Patients</li> <li>Best practices</li> </ul> Moderator:		
••••••			
MARKET TRENDS & WAY FORWARD	JABEEN AHMAD Global PV Consultant		
09:30 – Harmonisation & Effective PV systems	Independent Consultancy		
TANJA PETERS	Panellists:		
Senior Pharmacovigilance Expert, Deputy EU-QPPV & Head PV Intelligence, Boehringer Ingelheim	TANJA PETERS Senior Pharmacovigilance Expert, Deputy EU-QPPV & Head PV Intelligence, Boehringer Ingelheim		
IMPACT OF TECHNOLOGY	SUMIT MUNJAL		
10:00 - New technologies in Pharmacovigilance	Senior Medical Director, Head Europe PV Takeda Pharmaceuticals		
<ul> <li>Artificial intelligence/Machine learning in Pharmacovigilance</li> <li>Can PV keep up with the pace of innovation?</li> <li>Are stakeholders and PV systems ready to embrace AI?</li> </ul>	PHILIP OLUWOLE Associate Principal Surveillance Specialist Astrazeneca		
<ul><li>Information technology in pharmacovigilance</li><li>Decision process</li><li>Conclusions / Discussion</li></ul>	12:00 – Solution Provider Presentation		
	For sponsorship opportunities please contact info.uk@virtueinsight.com		
10:30 - Solution Provider Presentation	••••		
For sponsorship opportunities please contact info.uk@virtueinsight.com	12:20 - Networking luncheon		
•••••	•••••		
10:50 - Morning Coffee/Tea & Discussion	13:10 - Pharmacovigilance and signal management at the Danish Medicines Agency		
•••••	Pharmacovigilance in general at DMA		
11:20 - Keynote Panel Discussion: Optimising the PV ecosystem for betterment	<ul> <li>Signal detection, management and assessment</li> <li>Practical examples and outcomes</li> <li>Future trends in pharmacovigilance at DMA</li> </ul>		
<ul> <li>Discuss on the possible impacts of Brexit</li> </ul>			

KARE KEMP

management - PV/Safety Danish Medicines Agency

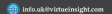
Team leader (senior advisor), Signals and risk

guidelines

Staying ahead in the race - Update on PV in EU, USA

& RoW - Current trends for PV, and new and future

Documentation (RMPs, PSURs, PADERs, PBRERs)



"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK "Very well organized and the sessions were so well placed. Got enough time for networking and well time managed"

Country Safety Lead, Pfizer Limited

## AGENDA

## DAY ONE - 26th February 2020

## **QUALITY - SAFETY - SIGNAL DETECTION**

## 13:40 - Panel Discussion - Quality, Safety & Signal Detection - Future of 2020

- Strategies for best practice in Signal Detection
- Exploring patient support and marketing research programs from a safety perspective
- How should we approach?
- Using technology to enhance interactive connection with patients
- Statistical signal detection as a routine pharmacovigilance practice
- Latest updates and hot topics

#### Moderator:

## **SUSAN WELSH** Chief Safety Officer

**CSL Behring** 

## **Panellists:**

#### **JOHN POUSTIE**

Medical Director & EU QPPV, Global Pharmacovigilance Norgine

## MIRCEA CIUCA

Global Therapeutic Area Head - Global Clinical Safety and Pharmacovigilance, CSL Behring

## YVONNE NANCIU

Senior Manager PV & Medical Information, Local QPPV Abbyie

## MADDALENA LINO

Terapheutic Area Safety Head Segirus

## 14:20 - Drug Safety contributions to First-In-Human (FIH) studies:

- Supporting translation of safety data from preclinical to clinical
- Evaluating possible safety concerns
- Identifying potential risks, designing appropriate risk minimization measures

## **MIRCEA CIUCA**

Global Therapeutic Area Head - Global Clinical Safety and Pharmacovigilance, CSL Behring

#### 14:50 - EU-RMP

- Changing paradigms for the list of safety concerns
- Transferring from previous templates to the R(2) template
- Reclassification of risks as not important for inclusions in the EU-RMP

### KAREN CHENG

Safety Risk Lead Pfizer

## 15:20 - Afternoon Tea/Coffee

15:40 - Emerging PV technologies and the future of the Drug Safety Professional: practical considerations for adoption of machine learning and NLP

- A framework model for leveraging PV innovation
- Development of cognitive services and the role of drug safety professional
- How the PV tech revolution will affect role of the drug safety professional
- How to pave the way to transformation and role evolution

## SALVATORE GIORGIO CICIRELLO

Senior Director Safety Science & PASS, Global Drug Safety & Risk Management, Celgene

#### 16:10 - Brexit Implications for the UK - Impacts on PV

- What would 'no deal' mean for medicine?
- Time to prepare now with not much of choice
- Solving stocked drugs issue
- Preparing for a smooth transition
- Pitfall and Learnings
- Innovation in PV

#### Moderator:

### SUSAN WELSH

Chief Safety Officer
CSL Behring

## Panellists:

#### CHETAN SHATAPATHY

Principal Pharamcovigilance Physician - Oncology R&D Unit, AstraZeneca





"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK "A great platform to understand the current practices & situation all across the industry, as well as individual approach of each company toward the goal of patient safety."

Senior Executive, Lupin

AGENDA AT A GLANCI DAY ONE - 26th February 2020

VALENTINA MANCINI Director PV, EU QPPV Shionogi Europe
SANDY EISEN Chief Medical Officer Frontline Pharma Consulting
16:50 - Chairperson's closing remarks and end of conference
••••••
17:00 – 18:00 – Networking Drinks - Take your discussions further & build new relationships in a relaxed & informal setting
NETWORKING DRINKS
Meet with your industry peers for a relaxed drink at the end of day one
FOR DELEGATE REGISTRATIONS:-
Our potent conference agenda delivering the latest information and the world class leaders as speakers attract delegates to attend from around the world. We aim for our attendees to be equipped with knowledge of latest developments & enable them to network with the industry key personnel.
$Delegate\ Registration\ \ delegate.uk@virtuein sight.com$
•••••





"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK "Informative session focusing on new and grey areas of Pharmacovigilance patient care being the utmost priority on minds of all the pharma company new aspect discussion and light on the grey areas had open new arena for Pharmacovigilance thank you"

Drug Safety Associate, Cipla

## AGENDA AT A GLANCE

## DAY TWO - 27th February 2019

08:30 - Coffee and Registration - An opportunity to meet and to network with your conference colleagues.  09:20 - Chairperson opening remarks  SUSAN WELSH Chief Safety Officer	<ul> <li>A review of general issues and the specific challenges with patients</li> <li>A practical approach to reshaping patient safety</li> <li>Next generation pharmacovigilance for enhanced patient safety</li> <li>Moderator:</li> <li>SUSAN WELSH</li> </ul>
CSL Behring	Chief Safety Officer
	CSL Behring
DV FOR 2020	Panellists:
PV FOR 2020	
09:30 - Pharmacovigilance in 2020	SHAANTANU DONDE Senior Director, Medical Portfolio Development (Daysland Markets) Uniohn Division of Pfirer
<ul> <li>Future horizons and efficiencies in data acquisition, evaluation and risk management</li> <li>Future-proofing Safety Systems</li> <li>Where are we?</li> <li>Boldly Shaping the Future</li> </ul>	(Developed Markets), Upjohn, Division of Pfizer  MATE A. BALAZS  Country Head - Patient Safety - Hungary  Novartis
ALINA TUDOR Associate Director, Senior PV Physician/Deputy EU QPPV, Norgine	MARY LYNNE VAN POELGEEST President World Federation for Incontinent Patients - (WFIP)
•••••	••••••
10:00 - Overview of FDA OCI and our involvement in Clinical Fraud	11:50 - Communication between global and local affiliate during HA Pharmacovigilance inspection
NICHOLAS CALL Special Agent FDA	<ul> <li>Identify "best common practice" to be prepared for a PV Inspection: after receiving a communication by HA about a pharmacovigilance inspection, global and local function has to prepare and verify that everything will be ok during the inspection.</li> </ul>
•••••	<ul> <li>List of aspects that's important to remember</li> </ul>
10:30 - Solution Provider Presentation	How local and global communicate during a PV inspection: during an inspection to an affiliate it's very
For sponsorship opportunities please contact info.uk@virtueinsight.com	important the continuous updating from local to global, in order to be aware about any potential finding and to be supportive to the affiliate for any question; so, how
	this communication can be ensured?
10:50 - Morning Coffee/Tea & Discussion	ANDREA OLIVA
•••••	Head of Pharmacovigilance Mylan
PATIENT SAFETY	••••••
11:10 - Keynote Panel Discussion: Pharmacovigilance and Patient Safety	12:20 - Networking luncheon
<ul> <li>Driving patient centricity into your PV plans</li> <li>Pharmacovigilance as a tool for safety and monitoring</li> <li>Patient-Perspectives in Benefit-Risk Assessments</li> </ul>	••••••••••••

"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK "Very nice opportunity to share current challenges within its own organisation with other Pharmacovigilance agents and hear about future initiatives to make our contribution to PV, safety, more efficiently moving forward."

Associate Director, Pharmacovigilance Operations, INCYTE Biosciences International

## AGENDA AT A GLANCE

## DAY TWO - 27th February 2019

### **RISK MANAGEMENT & PLANNING**

# 13:20 - Panel Discussion - PV - Risk Management and Planning

- Risk management in the lifecycle of a drug
- How effective is your risk management?
- Challenges and overcoming problems in Pharmaceutical product life cycle management
- Implementation and maintenance of RMP's Overcoming its challenges
- Risk management in different jurisdictions
- Benefit/Risk ratio: the common denominator
- New approaches for managing benefit-risk
- Research and development improvement

#### Moderator:

## SUSAN WELSH Chief Safety Officer

**CSL Behring** 

#### **Panellists:**

### **JOHN SOLOMON**

Head of Pharmacovigilance - UK & Ireland Sanofi

## BARBARA DE BERNARDI

Deputy EU QPPV and European Safety Office Head Pfizer

## IVA SLAVCEVOVA

Deputy QP Pharmacovigilance/Global Patient Safety Baxter

## NICOLE BAKER

Co-Founder BioLogit

## LUIZ LIMA

Senior Global Patient Safety Physician Neurology, Ipsen

## 14:00 - Patient-centric Safety: Innovative approaches and novel methodologies

- Emerging role of genetics in understanding drivers of toxicity
- · Translating biology into clinical decision-making
- Applying advanced technologies to perform novel safety analyses

#### **SCOTT CHANDLER**

Vice President & Global Head Licensing and Early Development (LEAD) Safety, Roche

## 14:30 - Why does pharmacovigilance sometimes fail and where could the fault lie?

- Risk blindness industry or drug authorities?
- It's not my fault but whom to blame?
- Hard to detect adverse reactions
- · Do we learn from previous experiences?

### **STEINAR MADSEN**

**Medical Director** 

**Norwegian Medicines Agency** 

## 15:10 - Afternoon Tea/Coffee

### **DATA COLLECTION - MANAGEMENT**

## 15:30 - Panel Discussion - PV Audit & Inspections -Knowing what is to be done

- Data Quality Management and Analysis
- PV Inspection readiness: What to expect? How ready can we be?
- Risk based selection criteria for auditing
- Methodologies, scope and oversight
- Preparing and managing safety data exchange agreements
- Relationship to other GxPs

## Moderator:

### **SUSAN WELSH**

Chief Safety Officer CSL Behring

#### **Panellists:**

## **WIVINA DE WAELE**

Director, Regional Safety Excellence EMEA.Global Drug Safety, Alexion

## FRANCK SCHWARTZ

QA Global Inspection, Intelligence Lead - Compliance and Regulatory Affairs Quality, Novartis

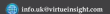
#### NATALIE SPRINGVELD

Global Safety Leader Bayer

### **TEA BABIC**

Associate Director, Audits and Inspections, Global Pharmacovigilance Compliance, Teva





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Safety Physician, Sciformix

AGENDA

## DAY TWO - 27th February 2019

## **REGULATION OVERVIEW & UPDATE**

## 16:10 - Panel Discussion: PV - Regulatory Updates

- Key current changes and their impact on current PV
- Impact of Brexit Regulatory aspect
- Pharmacovigilance and the role of regulatory affairs: How to achieve compliance across the business
- Future Legislation: Pharmacovigilance Industry Vision
- PV System Legislation Updates
- Current PV practices in the EU & US
- Enhancing communication between regulators, regional authorities and patients

### Moderator:

SUSAN WELSH Chief Safety Officer CSL Behring

Panellists:

### **MICHAEL BEAN**

Senior Director, Regulatory Compliance R&D Johnson & Johnson

#### YUUNG YUUNG YAP

Senior International Regulatory Counsel, EU and International Regulatory Law, Pfizer

## **RAJ BHOGAL**

Head of Inspection, R&D Quality Takeda

## **GEORGIA GAVRIILIDOU**

Counsel Sidley Austin

## MARJAN DZEPAROSKI

Head of Regulatory Affairs, Drug Safety & Intelectual Property, Bionika Pharmaceuticals

16:50 - 17:00 - Chairperson's closing remarks and end of the conference

### FOR SPONSORSHIP OPPORTUNITIES:-

Sponsorship or exhibition is the best way to speed network with decision makers. The world leader speakers in our conferences attract niche delegates from all over the world. This would be a wonderful opportunity to reach the right audience and save money and time on all your other advertising gimmicks. To give you an advertising edge we constantly update the industry pioneers via emails/news letter about the event and advertise the event via different forms of media.

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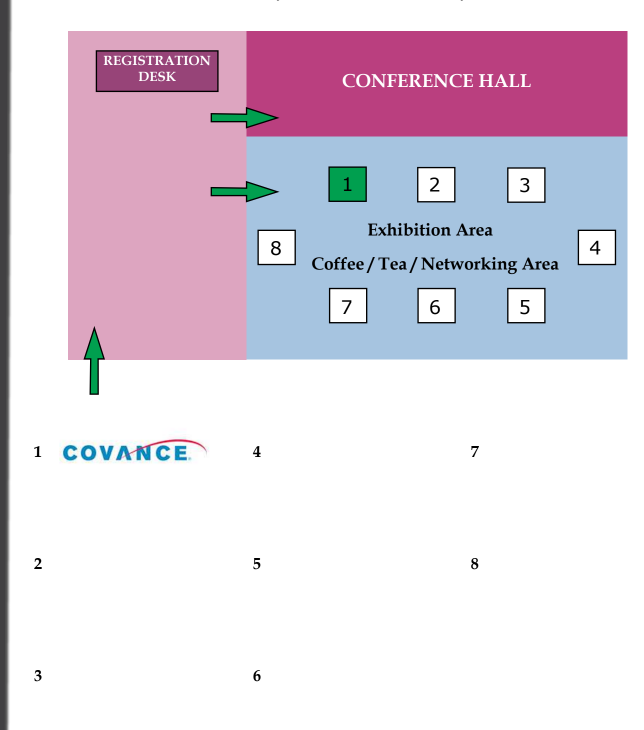
"Latest developments in pharmacovigilance, drug safety and RMP"  $\,$ 

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK "This conference was very good for the pharmacovigilance professionals as well as business people. Organising this event and the event management was nicely done by Virtue Insight"

IT Administrator, Oviya Med Safe Pvt. Ltd

AGENDA AT A GLANCE

**FLOOR PLAN -** Book your stalls now before they run out !!!



**Note :-** The floorplan is subject to change at the discretion of the organisers.





"Latest developments in pharmacovigilance, drug safety and RMP"

26th & 27th February 2020 Pestana Chelsea Bridge Hotel London, UK

"The conference was interesting and was a good platform for networking. The audience and the panelists were from varying backgrounds giving an insight to various challenges being faced by the Indian industry"

Manager - BD, ELC Research

#### **REGISTER ONLINE:**

Link: https://www.virtueinsight.com/pharma/21st-Pharmacovigilance-2020/products/

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## 3 for 2 Offer

#### TERMS AND CONDITIONS:

Payment terms: Virtue Insight requires the full amount to be paid before the conference. We may refuse entry to delegates who have not paid their invoice in full.

Cancellations: Delegates and vendors are subject to the following charges and refunds upon withdrawal or cancellation between 2-3 month's prior 75% cancellation fee/ 25% refund. Less than 2 months prior to the event Full cancellation fee / No refund.

Administration Fee: If you cancel your participation (once confirmed) and haven't paid the attendance fee you will be liable to pay an administration fee of £200

Substitutions/Name Change: If you are unable to attend you may nominate, in writing, another delegate to take your place at any time prior to the start of the event. This can be done at no extra cost.

Presentation: If you cannot attend the conference, you can still purchase the presentations for £500.

Indemnity: Virtue Insight reserves the right to make alterations to the conference/executive briefing content, timing, speakers or venue without notice. The event may be postponed or cancelled due to unforeseen events beyond the control of Virtue Insight. If such a situation arises, we will reschedule the event.

Fee: The conference fee includes lunch, refreshments and conference papers provided on the day. This fee does not include travel or hotel accommodation.

#### **VENUE**

Pestana Chelsea Bridge Hotel

Address: 354 Queenstown Rd, London SW8 4AE, UK Phone: +44 20 7062 8000



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