



Training Calendar

Courses

TBD <i>Online</i>	Healthcare Modeling: Interventi per Migliorare Efficacia ed Efficienza delle Cure <i>Come creare valore per tutti gli attori del sistema salute attraverso progetti di partnership tra Industria e Payors</i> <i>Market Access</i>
TBD <i>Online</i>	Clinical Support Programs: Strategie e Progetti per Favorire le Attività degli HCP <i>Interventi a beneficio dei professionisti sanitari per ottimizzare il percorso di cura delle persone</i> <i>Market Access</i>
29 February and 01 March 2024 <i>Online</i>	Labelling Requirements for Medical Devices <i>Understanding the regulatory labelling requirements for medical devices in the context of the MDR 2017/745</i> <i>Medical Device / Regulatory</i>
06 and 07 March 2024 <i>Online</i>	Clinical Evaluation for Medical Devices <i>Understanding the clinical evaluation requirements for the MedTech industry in the context of MDR 2017/745</i> <i>Medical Device / Regulatory</i>
12 and 13 March 2024 <i>Online</i>	Medical Devices Periodic Safety Update Report (PSUR) <i>What's behind the MDCG guidelines</i> <i>Medical Device</i>
13 and 15 March 2024 <i>Online</i>	The Next Generation of Medical Affairs: ChatGPT and Large Language Models (LLMs) <i>Equip yourself with the skills and knowledge to succeed in the AI-powered medical affairs landscape</i> <i>Medical Affairs</i>



14, 19 e 21 marzo 2024 Online	La Statistica Medica per Non Statistici <i>I principi statistici dalla pianificazione della ricerca alla pubblicazione sulle riviste scientifiche</i> Medical Affairs / Clinical Research / Statistics and Data Management
20, 21 and 25 March 2024 Online	How to Write a Clinical Evaluation Plan and Report <i>Find your way around the clinical evaluation of a medical device from concept to market approval and beyond</i> Medical Device
20 March 2024 Online	“Attributability” in Pharmacovigilance: Still a Hot Topic <i>How to navigate the different algorithms and approaches used to evaluate the causality of an adverse event/reaction</i> Pharmacovigilance
26 marzo 2024 Online	Il Consenso quale Base Giuridica per il Trattamento Dati nella Ricerca Clinico-Scientifica <i>Deep dive sul consenso, legittimo interesse e riutilizzo dei dati</i> Medical Affairs / Regulatory / Clinical Research
26 e 27 marzo 2024 Online	Annex 1 e Contamination Control Strategy <i>Approcci e applicazioni tecnico-operative per la conformità al nuovo standard</i> GMP (Good Manufacturing Practices)
03 e 05 aprile 2024 Online	Le Sfide della Produzione di un Investigational Medicinal Product (IMP) <i>Come declinare le richieste delle Good Manufacturing Practices (GMP) nel processo di produzione e controllo del farmaco sperimentale</i> GMP (Good Manufacturing Practices)
03, 04 and 05 April 2024 Online	Clinical Study Protocols - Structure & Content Medical Affairs / Medical Writing
09 e 11 aprile 2024 Online	Il Responsabile del Servizio Scientifico <i>Consapevolezza di un ruolo articolato</i> Medical Affairs



10 and 11 April 2024 <i>Online</i>	<p>Medical Writing Course: Improve your Writing & Reviewing Skills <i>Writing, editing & proofreading tips for medical writers: a standardised process to make your message effective; review & ensure document quality</i></p> <p><i>Medical Affairs / Medical Writing / Clinical Research</i></p>
10 aprile 2024 <i>Online</i>	<p>I Radiofarmaci</p> <p><i>Medical Affairs / Regulatory / Clinical Research</i></p>
16 e 18 aprile 2024 <i>Online</i>	<p>Normativa della Ricerca Clinica tra Presente e Futuro <i>Pillole di regolatorio</i></p> <p><i>Clinical Research</i></p>
18 April 2024 <i>Online</i>	<p>Advertisement and Promotion Claims for Medical Devices <i>How to ensure regulatory compliance of advertisement and promotion claims and activities in the MedTech industry</i></p> <p><i>Medical Device / IVDs (In-Vitro Diagnostics) / Medical Affairs / Regulatory</i></p>
09 e 10 maggio 2024 <i>Online</i>	<p>Agile Project Management: Gestire l'Innovazione nel Contesto della Produzione Farmaceutica <i>I fondamentali di Agile Project Management per affrontare progetti innovativi in realtà di produzione farmaceutica e nel rispetto delle GMP</i></p> <p><i>GMP (Good Manufacturing Practices)</i></p>
23 April 2024 <i>Online</i>	<p>Select the Ideal PMCF Strategy for a Medical Device <i>An advanced training on post-market requirements under the MDR 2017/745</i></p> <p><i>Medical Device</i></p>
23 aprile 2024 <i>Online</i>	<p>Progettare un Patient Support Program (PSP): Configurazione del Processo e Strumenti</p> <p><i>Medical Affairs</i></p>
29 April 2024 <i>Online</i>	<p>From Good to Excellent: The Summary of Safety and Clinical Performance (SSCP) <i>An advanced training to improve your SSCP and write it in the most efficient way</i></p> <p><i>Medical Device</i></p>
07 e 09 maggio 2024 <i>Online</i>	<p>La Pubblicità del Farmaco <i>Dall'informazione scientifica alla pubblicità al pubblico</i></p> <p><i>Regulatory</i></p>



08 maggio 2024 <i>Online</i>	Progettare e Comunicare in Medical Affairs <i>Medical Affairs</i>
13, 15 and 16 May 2024 <i>Online</i>	How to Write a Clinical Evaluation Plan and Report <i>Find your way around the clinical evaluation of a medical device from concept to market approval and beyond</i> <i>Medical Device</i>
14, 16 e 21 maggio 2024 <i>Online</i>	Patient Advocacy ed Engagement nell'Azienda Farmaceutica <i>Ruolo e mission in un approccio inter-funzionale</i> <i>Medical Affairs</i>
15 e 16 maggio 2024 <i>Online</i>	La Sorveglianza Post-Market dei Dispositivi Diagnostici in Vitro <i>Come implementare le corrette procedure per la gestione post-vendita</i> <i>IVDs (In-Vitro Diagnostics)</i>
16 and 17 May 2024 <i>Online</i>	Person Responsible for Regulatory Compliance (PRRC) - An MDR/IVDR Requirement <i>Responsibilities & challenges for medical device companies within MDR & IVDR</i> <i>Medical Device / IVDs (In-Vitro Diagnostics) / Regulatory</i>
21, 24 and 28 May 2024 <i>Online</i>	Pharmacovigilance and Safety in Clinical Trials under the Clinical Trial Regulation (EU) No 536/2014 <i>First experience with CTR/CTIS, regulatory expectations for safety documents, CTR assessment procedures and CTR Q&A document, typical issues and how to avoid it.</i> <i>Pharmacovigilance / Regulatory</i>
21 and 22 May 2024 <i>Online</i>	Pharmacovigilance Agreements <i>What pharmaceutical companies should consider when contracting with partners who may receive safety information relevant to their active substances</i> <i>Pharmacovigilance</i>
22 May 2024 <i>Online</i>	Beyond Slide Decks <i>How to use storytelling in your presentations</i> <i>Soft Skill</i>
22 maggio 2024 <i>Online</i>	Etichettatura degli Integratori Alimentari - Applichiamo la Normativa ad Esempi Reali <i>Regulatory</i>



22 maggio 2024 <i>Online</i>	Patient Support Program (PSP) e Compliance: quali Normative Considerare? <i>Il codice deontologico Farmindustria e i requisiti di privacy e farmacovigilanza per un PSP</i> <i>Pharmacovigilance / Market Access / Medical Affairs / Regulatory</i>
23 e 24 maggio 2024 <i>Milano</i>	La Gestione dell'Advisory Board nei Progetti Life Science <i>Modelli e strumenti</i> <i>Medical Affairs / Soft Skill</i>
27 May 2024 <i>Online</i>	Oral Presentations <i>Skills to help you survive or even shine</i> <i>Soft Skill</i>
28 e 30 maggio 2024 <i>Online</i>	Il Responsabile del Servizio Scientifico <i>Consapevolezza di un ruolo articolato</i> <i>Medical Affairs</i>
04 e 06 giugno 2024 <i>Online</i>	General Data Protection Regulation (GDPR), Ricerca Scientifica e Norme Locali <i>L'impatto del GDPR sugli studi clinici a livello locale ed internazionale</i> <i>Clinical Research</i>
04, 06 e 11 giugno 2024 <i>Online</i>	Signal Detection e Signal Management <i>Ricerca, identificazione, interpretazione e gestione dei segnali di sicurezza</i> <i>Pharmacovigilance</i>
04 June 2024 <i>Online</i>	How to Become a Successfully Published Author - Practical steps to make the publication process as smooth and as successful as possible <i>Medical Device / Medical Affairs / Medical Writing / Clinical Research</i>
05 e 07 giugno 2024 <i>Online</i>	La Gestione del Case Processing in Farmacovigilanza <i>Gestione delle segnalazioni di sospette reazioni avverse e aspetti di qualità nel case processing</i> <i>Pharmacovigilance</i>
05 e 06 giugno 2024 <i>Milano</i>	Comunicare con l'Intelligenza Relazionale <i>La flessibilità nelle relazioni per comunicare, guidare e motivare colleghi e interlocutori, in presenza e da remoto</i> <i>Soft Skill</i>



06, 07, 12 e 14 giugno 2024 <i>Online</i>	Il Sistema di Qualità Applicato alla Farmacovigilanza <i>Approfondire il ruolo centrale svolto dall'Assicurazione della Qualità nelle attività di Farmacovigilanza</i> <i>Pharmacovigilance</i>
10 e 17 giugno 2024 <i>Online</i>	Hospital Meeting: come Pianificare e Condurre una Riunione di Successo <i>Corso teorico pratico di alta formazione diretto all'apprendimento delle basi di pianificazione e conduzione ottimale di un hospital meeting</i> <i>Medical Affairs</i>
10, 14 e 17 giugno 2024 <i>Online</i>	Computer System Validation (CSV) - GxP Process Owner and Quality Assurance: In or Out? <i>Il ruolo del QA e del Process Owner nella convalida dei sistemi computerizzati GxP</i> <i>Regulatory / Clinical Research</i>
11 giugno 2024 <i>Online</i>	Le Good Calibration Practices (GCaIP) negli Impianti Automatici di Produzione <i>Come fidarsi dei dati che produciamo?</i> <i>GMP (Good Manufacturing Practices)</i>
12 e 13 giugno 2024 <i>Online</i>	Il Patient Support Program (PSP) e la Transizione Digitale: Opportunità e Rischi <i>Medical Affairs</i>
14 and 21 June 2024 <i>Online</i>	The PSMF (Pharmacovigilance System Master File): from GVPs to Inspections <i>A practical, hands-on guide</i> <i>Pharmacovigilance</i>
18 and 20 June 2024 <i>Online</i>	Fundamentals of European Cosmetics Regulatory Affairs <i>Regulatory</i>
18 and 20 June 2024 <i>Online</i>	Pharmacovigilance System: Audit & Inspection Readiness <i>Pharmacovigilance</i>
18, 24 and 27 June 2024 <i>Online</i>	A Practical Guide to Innovative Trial Design <i>Clinical Research / Statistics and Data Management</i>



19 and 20 June 2024 <i>Online</i>	Combination Products under the EU Medical Devices Regulation (MDR) <i>A targeted training to understand the regulatory pathway for device drug and drug device combinations in the European Union (EU). Updates from the MDR 2017/745 and EMA guidance</i> <i>Medical Device / Regulatory</i>
25 e 27 giugno, 2 e 4 luglio 2024 <i>Online</i>	Selezione e Convalida di Una Soluzione Cloud in Ambito GxP <i>Rischi e opportunità nella scelta di una soluzione Cloud a supporto di processi GXP</i> <i>Clinical Research</i>
25 and 26 June 2024 <i>Online</i>	Reporting Requirements in Veterinary Pharmacovigilance <i>Animal Health / Pharmacovigilance</i>
26 giugno 2024 <i>Online</i>	Integratori Alimentari e Advertising: Applichiamo la Teoria alla Pratica <i>Regulatory</i>
28 giugno 2024 <i>Online</i>	Qualifica dei Vendor nella Ricerca Clinica - Approfondimenti <i>Esempi e aspetti pratici</i> <i>Clinical Research</i>
02 July 2024 <i>Online</i>	Statistical Process Control for Pharmaceutical Manufacturing <i>Tips for non-statisticians to navigate statistical methods and make data-driven decisions</i> <i>GMP (Good Manufacturing Practices)</i>
02 and 03 July 2024 <i>online</i>	Labelling Requirements for Medical Devices <i>Understanding the regulatory labelling requirements for medical devices in the context of the MDR 2017/745</i> <i>Medical Device</i>
16 and 17 July 2024 <i>Online</i>	Good Distribution Practices of Medical Devices <i>Ensuring Compliance</i> <i>Medical Device / Regulatory / Clinical Research</i>
16 September 2024 <i>Online</i>	Quality by Design <i>How to understand and control process and product variables to ensure quality from the very beginning</i> <i>GMP (Good Manufacturing Practices)</i>



<p>16, 19 and 23 September 2024</p> <p>Online</p>	<p>Pharmacovigilance and Safety in Clinical Trials under the Clinical Trial Regulation (EU) No 536/2014 <i>First experience with CTR/CTIS, regulatory expectations for safety documents, CTR assessment procedures and CTR Q&A document, typical issues and how to avoid it.</i></p> <p><i>Pharmacovigilance / Regulatory</i></p>
<p>17, 18 and 19 September 2024</p> <p>Online</p>	<p>Tips and Tricks to Improve your Technical/Scientific Writing <i>Learn the basic techniques to effectively write technical/scientific documents</i></p> <p><i>Medical Writing</i></p>
<p>17 and 18 September 2024</p> <p>Online</p>	<p>Introduction to Pharmacovigilance <i>A short but comprehensive guide to the basis of drug safety</i></p> <p><i>Pharmacovigilance</i></p>
<p>17 September 2024</p> <p>Online</p>	<p>Good Distribution Practices (GDP) for Veterinary Medicinal Products</p> <p><i>Animal Health / Regulatory</i></p>
<p>18 e 19 settembre 2024</p> <p>Online</p>	<p>EU GMP Annex 15 - La Qualifica e la Convalida nell'Industria Farmaceutica <i>Panorama Normativo, linee guida applicabili e approcci per la conformità di impianti, attrezzature e sistemi</i></p> <p><i>GMP (Good Manufacturing Practices)</i></p>
<p>18 September 2024</p> <p>Online</p>	<p>Pharmacovigilance Documents - Basic Concepts and Definitions for Pharmacovigilance Writing</p> <p><i>Pharmacovigilance / Medical Writing</i></p>
<p>01 and 02 October 2024</p> <p>Online</p>	<p>Process Validation <i>Strategies to implement a continuous process verification during the product life cycle</i></p> <p><i>GMP (Good Manufacturing Practices)</i></p>
<p>19 e 20 settembre 2024</p> <p>Online</p>	<p>La Conduzione di uno Studio Clinico con un Dispositivo Medico-Diagnostico in Vitro <i>Dai requisiti regolatori per la sottomissione alla gestione dello studio</i></p> <p><i>IVDs (In-Vitro Diagnostics) / Clinical Research</i></p>



20 settembre 2024 <i>Online</i>	ICH Q9 Quality Risk Management <i>Strumenti per la gestione del rischio nell'industria farmaceutica</i> <i>GMP (Good Manufacturing Practices)</i>
23 and 26 September 2024 <i>Online</i>	Searching the Medical Literature <i>Best resources and tips for finding medical information</i> <i>Medical Device / Pharmacovigilance / Market Access / Medical Affairs / Medical Writing / Clinical Research</i>
23 e 24 settembre 2024 <i>Online</i>	La Biocompatibilità e la Caratterizzazione Chimica: Dispositivi Medici Sicuri <i>Requisiti normativi, gestione del rischio e valutazione dei dati</i> <i>Medical Device / Regulatory</i>
24, 26 settembre e 01 ottobre 2024 <i>Online</i>	Patient Support Program (PSP) e Patient Solution: Compiere una Scelta Strategica a Supporto del Paziente <i>Market Access / Medical Affairs / Regulatory</i>
24 e 26 settembre 2024 <i>Online</i>	La Ricerca della Letteratura Scientifica: dal Quesito ai Risultati <i>Principali funzionalità delle banche dati biomediche e degli strumenti di gestione delle ricerche bibliografiche</i> <i>Medical Device / Pharmacovigilance / Medical Affairs / Medical Writing / Clinical Research</i>
25 e 26 settembre 2024 <i>Online</i>	ISO 14155/2020 - Come Svolgere uno Studio Clinico con Dispositivi Medici <i>Implementazione, Ottemperanza e Chiavi per il Successo</i> <i>Medical Device</i>
25 September 2024 <i>Online</i>	Pharmacovigilance Documents - Focus on Signal Management and Development Safety Update Reports (DSUR) <i>Pharmacovigilance / Medical Writing</i>
STEP 2 : 10 and 17 October 2024 <i>Online</i>	A Systematic Approach to Real World Evidence (RWE) Generation in Life Sciences - A 2 Step Intensive Course <i>How to manage the main methodological, regulatory and operational challenges in using Real World Data (RWD) for regulatory and strategic purposes</i> <i>Evidence Generation / Medical Affairs / Clinical Research</i>



26 e 27 settembre 2024 <i>Online</i>	Buone Pratiche per la Gestione dei Documenti Cartacei e dei Dati Elettronici in Ambito GxP <i>Strumenti per la conformità alle Good Documentation Practices (GDP) e ai requisiti di Data Integrity</i> <i>Pharmacovigilance / GMP (Good Manufacturing Practices) / Clinical Research</i>
27 settembre 2024 <i>Online</i>	Codice Deontologico di Farmindustria e le Linee Guida per la Certificazione delle Attività Scientifiche: come applicarle in azienda <i>Medical Affairs / Regulatory</i>
15 October 2024 <i>Online</i>	Introduction to Aseptic Process Simulation (APS) <i>Fundamentals to comprehend Aseptic Process Simulation</i> <i>GMP (Good Manufacturing Practices)</i>
09 ottobre 2024 <i>Online</i>	Integratori Alimentari ed Advertising <i>Caratteristiche di una Corretta Comunicazione Commerciale e Vincoli Normativi</i> <i>Regulatory</i>
01 e 03 ottobre 2024 <i>Online</i>	Cosmetici - Aspetti Tecnico-Regolatori e Panorama Normativo <i>Regulatory</i>
02 October 2024 <i>Online</i>	Pharmacovigilance Documents - Focus on the Risk Management Plan (RMP) <i>Pharmacovigilance / Medical Writing</i>
02 October 2024 <i>online</i>	Electronic Submissions and Data Management in Regulatory Affairs <i>From eCTD to xEVMPD and IDMP, and how to manage your regulatory information fit for purpose</i> <i>Regulatory</i>
21 and 28 October 2024 <i>Online</i>	Biologics and Biosimilars Manufacturing <i>A practical guide to demystify and address the most common challenges</i> <i>GMP (Good Manufacturing Practices)</i>
08 e 10 ottobre 2024 <i>Online</i>	Gestione della Proprietà Industriale - Il Brevetto per Tutelare le Invenzioni <i>Regulatory</i>



09 and 10 October 2024 <i>Online</i>	Medical Writing Course: Improve your Writing & Reviewing Skills <i>Writing, editing & proofreading tips for medical writers: a standardised process to make your message effective; review & ensure document quality</i> <i>Medical Affairs / Medical Writing / Clinical Research</i>
09 October and 16 October 2024 <i>Online</i>	Pharmacovigilance Documents - Focus on the Periodic Safety Update Report (PSUR) <i>Pharmacovigilance / Medical Writing</i>
10 and 11 October 2024 <i>Online</i>	State of the Art Section for Medical Devices - Unpacking the Tips and Tricks of a Complex Document <i>Medical Device</i>
10 and 11 October 2024 <i>Online</i>	US FDA 101 - Fundamentals of Pre-market Submissions to CDRH <i>Understanding US FDA pre-market submission pathways from an EU perspective - and why engaging the Agency through Pre-Submission Meetings is a valuable tool for Sponsors</i> <i>Medical Device / Regulatory</i>
15 and 16 October 2024 <i>Online</i>	Medical Devices Periodic Safety Update Report (PSUR) <i>What's behind the MDCG guidelines</i> <i>Medical Device</i>
15 and 17 October 2024 <i>Online</i>	The Basics of Regulatory Affairs for Cosmetic Products in US and Canada <i>Regulatory</i>
15 e 23 ottobre 2024 <i>Online</i>	Off-Label, Uso Compassionevole e Accesso Precoce (Early Access) <i>Le normative e le loro applicazioni</i> <i>Medical Affairs / Regulatory / Clinical Research</i>
17 October 2024 <i>Online</i>	How to Create Effective Visuals for Better Communicating your Science <i>Pharmacovigilance / Medical Affairs / Medical Writing / Soft Skill</i>
17 October 2024 <i>Online</i>	Human Error: the True Root Cause of a Deviation? <i>Tips and strategies to understand and manage why human errors occur and to minimize them</i> <i>GMP (Good Manufacturing Practices)</i>



21 e 24 ottobre 2024 <i>Online</i>	Terapie Digitali (DTx): a che Punto Siamo? <i>L'innovazione digitale per il futuro delle terapie</i> <i>Medical Affairs / Clinical Research</i>
22 October 2024 <i>Online</i>	What You Need to Know about Medical Device Software and Never Dared to Ask <i>More than just apps - An introduction to Medical Device Software</i> <i>Medical Device</i>
22 and 23 October 2024 <i>Online</i>	Veterinary Pharmacovigilance System <i>Animal Health / Pharmacovigilance</i>
22 e 29 ottobre 2024 <i>Online</i>	Terapie Geniche e Terapie Cellulari <i>Dalla Ricerca all'Impiego nel Real World</i> <i>Regulatory</i>
23 and 24 October 2024 <i>Online</i>	Clinical Study Reports - a 360° Perspective <i>Planning and Authoring CSRs in Accordance with Public Disclosure Requirements</i> <i>Medical Writing / Clinical Research</i>
23 October 2024 <i>Online</i>	Pharmacovigilance Documents - Focus on Addendum to the Clinical Overview (AddCO) and Referrals <i>Pharmacovigilance / Medical Writing</i>
28 and 29 October 2024 <i>Online</i>	Initiating the Development of Artificial Intelligence (AI) Medical Devices <i>Developing AI medical devices considering the latest industry expectations</i> <i>Medical Device</i>
29 October 2024 <i>Online</i>	Writing Science for Lay Audiences <i>Ways to better get your message across</i> <i>Medical Device / Medical Affairs / Medical Writing / Clinical Research</i>
29 and 31 October 2024 <i>Online</i>	Cleaning Validation <i>GMP requirements and technical methods to ensure clean production systems and small materials</i> <i>GMP (Good Manufacturing Practices)</i>



30 October and 06 November 2024 <i>Online</i>	Advanced Therapy Medicinal Product (ATMP): a Roadmap from Classification to Regulation and Manufacturing <i>GMP (Good Manufacturing Practices) / Regulatory</i>
30 October 2024 <i>Online</i>	Veterinary Marketing Authorisation Application (MAA) in the EU <i>Clinical and Safety documentation</i> <i>Animal Health / Regulatory</i>
31 ottobre 2024 <i>Online</i>	La Ricerca della Letteratura Scientifica: come Sfruttare le Risorse Gratuite in Rete <i>Medical Device / Pharmacovigilance / Medical Affairs / Medical Writing / Clinical Research</i>
04 novembre 2024 <i>Online</i>	La Compliance ai Requisiti GMP: Ruolo e Funzioni dell'Assicurazione Qualità <i>Standard GMP e processi di qualità: teoria ed esempi pratici</i> <i>GMP (Good Manufacturing Practices)</i>
05 and 07 November 2024 <i>Online</i>	Marketing Authorization Application in EU, US and UK <i>Regulatory framework for a strategic plan until the entry in the market</i> <i>Regulatory</i>
05 and 07 November 2024 <i>Online</i>	Beyond PubMed <i>Additional approaches and sources for cost-effective literature monitoring</i> <i>Pharmacovigilance / Medical Affairs / Clinical Research</i>
05, 07, 12 e 14 novembre 2024 <i>Online</i>	Safety Management e Farmacovigilanza <i>Aspetti normativi, clinici e metodologici della farmacovigilanza, gestione della safety nello sviluppo clinico del farmaco</i> <i>Pharmacovigilance / Clinical Research</i>
06, 08 e 13 novembre 2024 <i>Online</i>	Integratori Alimentari - Aspetti Tecnico-Regolatori e Panorama Normativo <i>Regulatory</i>
06, 13 and 20 November 2024 <i>Online</i>	All You Need to Know to Understand Statistic if You are not a Statistician <i>Statistical principles from research planning to publication in scientific journals</i> <i>Medical Affairs / Medical Writing / Clinical Research</i>



11, 15 e 18 novembre 2024 <i>Online</i>	Audit to Computer Systems <i>Be ready!</i> <i>Clinical Research</i>
12 e 14 novembre 2024 <i>Online</i>	La Vigilanza Post Market per i Dispositivi Medici secondo MDR e FDA <i>Dalla gestione del reclamo alla sua notifica come incidente grave alle Autorità Competenti in Europa o come evento avverso in USA. La gestione delle field action in Europa.</i> <i>Medical Device / Regulatory</i>
12 and 14 November 2024 <i>Online</i>	Sterilization Validation <i>How to be sure of achieving sterility in pharmaceutical processes</i> <i>GMP (Good Manufacturing Practices)</i>
13 e 15 novembre 2024 <i>Online</i>	Condurre uno Studio Clinico Adempiendo alla Normativa GDPR: un Ponte fra Good Clinical Practices (GCP) e General Data Protection Regulation (GDPR) <i>Clinical Research</i>
19 and 21 November 2024 <i>Online</i>	Mastering Benefit Risk Assessment in Pharmacovigilance <i>Pharmacovigilance</i>
19, 21, 26 and 28 November 2024 <i>Online</i>	Computer System Validation (CSV) and Assurance (CSA) <i>Insights into CSV fundamentals and different approaches for CSV and CSA</i> <i>GMP (Good Manufacturing Practices)</i>
19 and 21 November 2024 <i>Online</i>	Pharmacovigilance Quality Management System (QMS) <i>Pharmacovigilance</i>
20 and 25 November 2024 <i>Online</i>	Protocol Writing and Communication of Real World Evidence <i>International methodological standards for writing and publishing observational study protocols</i> <i>Evidence Generation / Medical Affairs / Medical Writing / Clinical Research</i>
20 November 2024 <i>Online</i>	Decoding Electronic Product Information (ePI) <i>An in-depth exploration from paper to ePI</i> <i>Regulatory</i>



25 November 2024 <i>Online</i>	GMP Inspections: How to Be Ready? <i>Practical Guide to Preparing for and Managing a Regulatory Inspection of a Pharmaceutical Manufacturing Facility</i> <i>GMP (Good Manufacturing Practices)</i>
26 and 27 November 2024 <i>Online</i>	Medical Reading <i>The critical evaluation of scientific publications</i> <i>Medical Device / Medical Affairs / Clinical Research</i>
27 November 2024 <i>Online</i>	Veterinary Marketing Authorisation Variation (MAV) in the EU <i>Animal Health / Regulatory</i>
28 November 2024 <i>Online</i>	The “Global” Qualified Person Responsible for Pharmacovigilance (QPPV) Workshop <i>Evolution of the role, challenges and opportunities in a global environment</i> <i>Pharmacovigilance</i>
03 December 2024 <i>Online</i>	Veterinary Pharmacovigilance Quality Management <i>Animal Health / Pharmacovigilance</i>
10 and 12 December 2024 <i>Online</i>	Introduction to Veterinary Pharmacovigilance <i>Animal Health / Pharmacovigilance</i>
13 dicembre 2024 <i>Online</i>	ICH GCP (R3) <i>Innovazione, Flessibilità e Qualità nella Ricerca Clinica</i> <i>Clinical Research</i>

Percorsi Formativi

Dal 12 marzo al 18 giugno 2024 <i>Online</i>	Percorso Formativo GMP Project Management <i>Il ruolo del Project Manager nella gestione di progetti nei siti produttivi farmaceutici secondo la ISO 21502</i> <i>GMP (Good Manufacturing Practices) / Soft Skill</i>
From 18 September to 23 October 2024 <i>Online</i>	Pharmacovigilance Documents in the Life Cycle of a Medicinal Product <i>From Papers and Patients to Health Authorities</i> <i>Pharmacovigilance / Medical Writing</i>



<p>dal 27 settembre al 23 novembre 2024</p> <p>Online</p>	<p>Clinical Quality Assurance: un Ruolo Chiave nella Ricerca Clinica <i>Percorso di alta specializzazione per lo sviluppo delle competenze peculiari del Clinical Quality Assurance (QA)</i></p> <p><i>Clinical Research</i></p>
<p>Dal 04 ottobre al 13 dicembre 2024</p> <p>6 week end (online) e 1 project work (online)</p>	<p>Medical Affairs: Prepararsi alle Sfide Future di un Ruolo Complesso <i>Corso di alta specializzazione per lo sviluppo delle competenze tecnico-scientifiche peculiari del Medical Advisor (MA) e del Medical Science Liaison (MSL)</i></p> <p><i>Medical Affairs</i></p>

Conferences

<p>17 September 2024</p> <p>Frankfurt am Main</p>	<p>German Pharmacovigilance Day <i>Pharmacovigilance: The Many Facets of one EU Framework</i></p>
<p>07 and 08 October 2024</p> <p>Copenhagen</p>	<p>MedDev Day <i>Medical Devices Regulation Update: Exploring the Thriving Regulatory Landscape</i></p>
<p>23 Ottobre 2024</p> <p>Bergamo</p>	<p>Italian Pharmacovigilance Day <i>La Farmacovigilanza al Servizio del Paziente: Nuove Sfide, Esempi di Successo e Strategie di Eccellenza</i></p>
<p>13 November 2024</p> <p>Copenhagen</p>	<p>Nordic Pharmacovigilance Day <i>Harmonizing Perspectives to Support Implementation of Common Objectives in Pharmacovigilance: Insights from the Nordic Region</i></p>

