

Virtual Training Agenda

Introduction to Regulation of Drugs and Biologics in the United States: Pre-Course Part One

Introduction to Regulation of Drugs and Biologics in the United States: Pre-Course Part Two

*Pre-course work can be done through the DIA Learning Center at any time prior to the virtual event.

Regulatory Affairs: Part 1: The IND – Investigational New Drug

9:00 – 9:30 AM	Welcome and Introduction
9:30 – 9:45 AM	 Drug Development: Pre-Course Work Review* Introduction to Drug Development Regulation Highlights Key Definitions Drug Development Highlights Regulatory Strategy
9:45 – 10:00 AM	Session 1: The IND – A General Introduction • What is an IND • When is an IND Required/Not Required • Types of INDs
10:00 – 10:45 AM	Session 2: The IND in Detail – Modules 1, 2, and 5 IND Item 1: Form FDA 1571 IND Item 2: Table of Contents IND Item 3: Introductory Statement IND Item 4: General Investigational Plan IND Item 5: Investigator's Brochure IND Item 6: Protocols
10:45 – 11:00 AM	Break
11:00 – 11:45 AM	Session 3: IND in Detail – Modules 3 and 1 • IND Item 7: Chemistry, Manufacturing, and Controls
11:45 AM – 12:45 PM	Session 4: The IND in Detail – Modules 4, 5 and 2 IND Item 8: Nonclinical Pharmacology and Toxicology IND Item 9: Previous Human Experience IND Item 10: Additional Information
12:45 – 1:30 PM	Lunch Break



Regulatory Affairs: Part 2: The IND Amendments

1:30 – 1:50 PM	 Session 5: The IND in Detail – Additional Topics Additional Requirements for Biologics and Biotechnology-Derived Products Submission of an Initial IND
1:50 – 2:30 PM	Session 6: FDA's Actions on the Original IND and Future Amendments • FDA's Review of an IND • Clinical Holds: Basis for Imposition and Process for Removal
2:30 – 2:45 PM	Break
2:45 – 4:00 PM	 Session 7: IND Amendments and Maintenance Amendments to the IND Application Annual Reports/DSUR Noncommercial INDs Exploratory INDs, Sponsor-Investigator INDs, Expanded Access IND Administrative Actions and Sponsor Activities
4:00 – 5:15 PM	Session 8: IND Amendments Workshop • In this workshop, learners will break into teams to determine the type and content of IND amendments needed to support changes to a protocol and associated development activities in their hypothetical company
5:15 – 6:15 PM	Virtual Reception
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Regulatory Affairs: Part 3: Special Topics

9:00 – 9:15 AM	Welcome and Review of Day 1		
9:15 – 10:45 AM	Session 9: Special Topics for Clinical Research		
	Changes to the Investigational DrugFinancial Disclosure by Clinical InvestigatorsSpecial Protocol Assessment		

10:45 - 11:00 AM Break



11:00 AM – 12:30 PM Session 10: Special Regulatory Considerations for Development

- Expedited Programs for Serious Conditions
- Background and Subpart E
- Concepts for Expedited Programs
 - Serious Conditions/Available Therapy/Unmet Medical Need
- Expedited Programs
 - Fast Track
 - Breakthrough Therapy (BTD)
 - o Regenerative Medicine Advanced Therapy (RMAT)
 - Qualified Infectious Disease Product (QIDP)
 - Accelerated Approval
 - o Priority Review
- Other Programs
 - The Animal Rule
 - Limited Population Pathway for Antibacterial and Antifungal Drugs (LPAD)
 - Orphan Products
 - Priority Review Vouchers
- Overview of Special Development Programs

12:30 – 1:15 PM Lunch Break

Regulatory Affairs: Part 4: Adverse Events and Quality Assurance

1:15 – 2:15 PM Session 11: Reporting Adverse Events (AEs) During Clinical Trials

- Definitions of Terms
- IND Safety Reports
- IND Annual Reports Safety Information
- Discontinuation of Studies for Safety Reasons

2:15 – 2:20 PM Break

2:20 – 4:05 PM Session 12: Adverse Event (AE) Reporting Workshop

 In this workshop, learners will break into teams to discuss IND Safety Reporting based on safety information received from clinical trials and other sources in their hypothetical company

4:05 – 4:15 PM Break



4:15 – 4:45 PM Session 13: Quality Assurance in Drug Development (GxPs)

- Good Clinical Practices
 - Sponsor Responsibilities
 - Investigator Responsibilities
 - Institutional Review Boards
 - Informed Consent
- Good Laboratory Practices
- Good Manufacturing Practices

4:45 – 5:00 PM Question & Answer

Regulatory Affairs: Part 5: The NDA - New Drug Application

9:00 – 9:15 AM Welcome and Review of Day 2

9:15 – 10:30 AM Session 14: The NDA: Planning, Content, Types of NDAs/BLAs, and Exclusivity

- Getting from the IND to the NDA
- NDA Data Sources and Specific Populations
- Types of NDAs
- · BLAs: Biologics and Biosimilars
- Combination Products and OTC Drugs
- Patent Term Restoration and Exclusivity

10:30 - 10:45 AM Break

10:45 AM- 12:00 PM Session 15: The NDA in CTD Format: Modules 1-5

- The Common Technical Document (CTD) Format Overview
- Module 1
- Module 3
- Module 4
- Module 5
- Module 2
- Safety Update Reports (CTD Module 5)

12:00 – 12:45 PM Lunch Break

Regulatory Affairs: Part 6: Interactions with FDA

12:45 – 1:45 PM Session 16: NDA Submission, FDA Review and Action on Applications

- Electronic Submissions
- FDA Review of Applications and Actions on Applications
- Amendments to an Unapproved Application
- Reasons Applications are not Approved
- PDUFA



1:45 – 2:15 PM	Session 17: The FDA and Risk Management The foundation of risk management Pre-Marketing Risk Assessment Post-Marketing Risk Assessment Risk Evaluation and Mitigation Strategies (REMS)
2:15 – 2:35 PM	Break
2:35 – 3:50 PM	 Session 18: Interactions with FDA – Part 1 Communications with FDA Formal Meetings with FDA Time Course of Events in Requesting and Preparing for a Meeting Objectives and Conduct of Specific Meetings
3:50 – 3:55 PM	Stretch Break
3:55 – 5:15 PM	Session 19: Interactions with FDA – Part 2 Principles for Communicating with FDA Meeting Etiquette Resolving Issues or Disputes with FDA Summary on Interacting with FDA Advisory Committee Meetings Advisory Committee Meeting Video
5:15 – 5:30 PM	Questions and Answers / Preparation for Mock FDA Meeting

Regulatory Affairs: Part 7: Mock FDA Meeting

9:00 – 9:15 AM Welcome and Review of Day 3

9:15 AM – 12:15 PM Session 20: Mock FDA Meeting (There will be a 15-minute break during this session)

Learners will break into FDA or company teams and conduct a pre-NDA meeting.

12:15 - 1:00 PM Lunch Break



Regulatory Affairs: Part 8: Post-NDA

1:00 - 2:30 PM Session 21: Regulatory Compliance and FDA Inspections: What to Expect After Submitting the NDA

- GLP Inspections
- GCP Inspections
- Inspection Outcomes: Additional Considerations in GCP Inspections
- GMP Inspections
- Inspection Outcomes (GLPs, GCPs, GMPs)
- FDA Enforcement Actions
- Application Integrity Policy (AIP)
- Inspection Video

2:30 – 2:45 PM Break

2:45 – 4:00 PM Session 22: Post-NDA Approval Regulatory Requirements

- Post-NDA Approval Obligations
- Post-Marketing Requirements and Commitments
- Supplements and Other Changes to an Approved Application
- Post-Marketing Safety Reporting
- NDA Annual Reports
- Other Post-Marketing Reports

4:00 – 4:15 PM Questions and Answers

Regulatory Affairs: Part 9: Regulatory Requirements and Post-Approval Workshop

9:00 – 9:15 AM Welcome and Review of Day 4

9:15 – 10:30 AM Session 23: Requirements for Prescription Drug Labeling

- Definitions
- Labeling Requirements of Immediate Containers and Cartons
- Content and Format of Prescribing Information (PI)
- Patient Labeling
- Structured Product Labeling (SPL)

10:30 - 10:45 AM Break

10:45 – 11:30 AM Session 24: Requirements for Prescription Drug Advertising and Promotional Labeling

- Definitions
- Statutory Basis for Promotional Regulations
- Required Elements for Advertisements and Promotional Labeling
- Reminder Advertisements/Labeling
- Direct-to-Consumer (DTC) Advertising
- Social Media
- Disease Awareness/Help-Seeking
- Launch of Promotional Pieces
- Post-Marketing Submission of Advertising



- Pre-Approval Promotion and Dissemination of Off-Label Information
- FDA Enforcement Actions

11.00 11.40 / W Working Edition Dica	11:30 - 11:45 Al	√ Working	Lunch	Break
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11:45 AM – 1:15 PM Session 25: Post-Approval Workshop

In this workshop, learners will break into teams to review planned changes to a
hypothetical approved product and determine the type and content of FDA submissions
needed to implement these changes.

1:15 – 2:00 PM Session 26: Review of Resources for Regulatory Intelligence

2:00 – 3:00 PM Session 27: Regulatory Intelligence Workshop

In this workshop, learners will break into and use publicly available resources to address
potential development, lifecycle management and general regulatory questions in their
hypothetical company.

3:00 – 3:15 PM Course Wrap-Up

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