

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* <ul style="list-style-type: none">• 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 <ul style="list-style-type: none">• 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 <ul style="list-style-type: none">• 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 <ul style="list-style-type: none">• IND Item 7: Chemistry, Manufacturing, and Controls
11:45 AM – 12:45 PM	Session 4: The IND in Detail – Modules 4, 5 and 2 <ul style="list-style-type: none">• 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: The IND, NDA, and Post-Marketing

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**

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**
- Adequate and Well-Controlled Trials
 - Adaptive Study Designs
 - Comparative Effectiveness Research (CER)
 - Patient Reported Outcomes
 - Surrogate Endpoints
 - Qualification of Drug Development Tool
 - Foreign Clinical Trials
 - Changes to the Investigational Drug
 - Financial Disclosure by Clinical Investigators
 - Special Protocol Assessment
- 10:45 – 11:00 AM **Break**

Regulatory Affairs: The IND, NDA, and Post-Marketing

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)
 - Regenerative Medicine Advanced Therapy (RMAT)
 - Qualified Infectious Disease Product (QIDP)
 - Accelerated Approval
 - 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

Regulatory Affairs: The IND, NDA, and Post-Marketing

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

Regulatory Affairs: The IND, NDA, and Post-Marketing

- 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: The IND, NDA, and Post-Marketing

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

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

11:30 – 11:45 AM Working Lunch Break

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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