



**NDA Simulation Workshop: Pharmaceutical Development Decisions  
June 2024  
4000 Legato Road, Suite 1100, Fairfax, VA**

***The Panoramic Conference Room***

**Monday, June 3, 2024**  
All Times Listed EDT

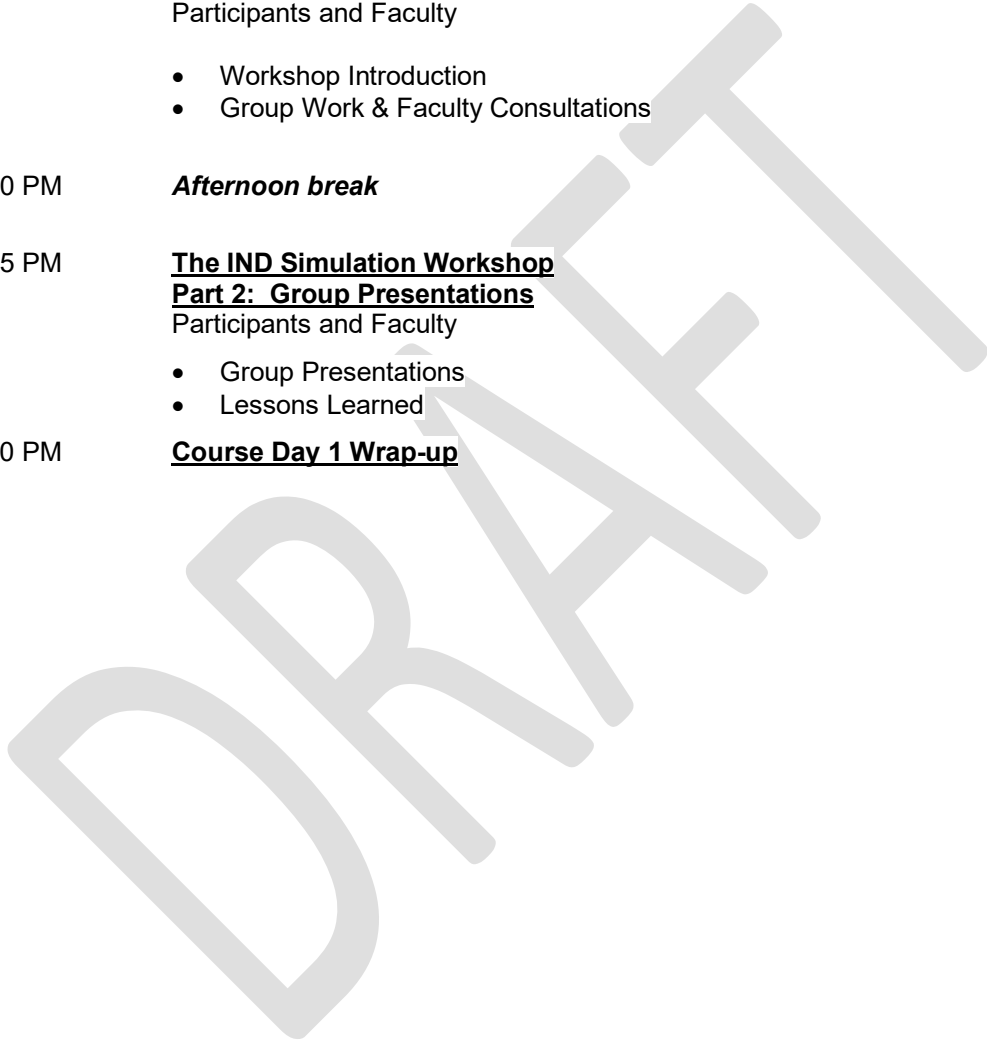
**Session**  
**No.**

8:00 – 8:30 AM	<b><u>Registration &amp; Continental Breakfast</u></b>	
8:30 – 8:45 AM	<b><u>PERI Welcome &amp; Course Introduction</u></b> <ul style="list-style-type: none"><li>• Course overview</li><li>• Faculty introductions</li><li>• Participant introductions</li></ul>	
8:45 – 10:00 AM	<b><u>Regulatory Overview</u></b> Allan J. Weinstein, MD <ul style="list-style-type: none"><li>• Regulations for new drug and biologic approval</li><li>• Overview of regulatory process</li><li>• Understanding the regulatory agencies' role in clinical development: US, EU, Japan</li><li>• Regulatory Update: changes in last 10 years</li><li>• Emerging Regulatory Landscape (PDUFA VII)</li></ul>	<b>1</b>
10:00 – 10:15 AM	<b><u>Morning Break</u></b>	
10:15 – 11:15 AM	<b><u>Chemistry, Manufacturing and Controls</u></b> Noel J. Cusack, PhD Independent Consultant, Nonclinical Development <ul style="list-style-type: none"><li>• Drug Substance (Active Pharmaceutical Ingredient)<ul style="list-style-type: none"><li>○ Small molecules</li></ul></li><li>• Drug Product (Formulated Drug Substance)<ul style="list-style-type: none"><li>○ Small molecules</li></ul></li><li>• Rationale for formulation Case studies</li></ul>	<b>2</b>
11:15 AM – 12:30 PM	<b><u>Key Concepts in Nonclinical Safety and Toxicology</u></b> Noel J. Cusack, PhD <ul style="list-style-type: none"><li>• Objectives of Nonclinical Safety for Clinical Trial Planning<ul style="list-style-type: none"><li>○ Small molecules</li></ul></li><li>• Nonclinical safety assessment (Risk)</li><li>• Dose selection for First-in-Human clinical trial</li><li>• Safety support during drug development</li><li>• Case studies</li></ul>	<b>3</b>
12:30 – 1:15 PM	<b><u>Lunch Break</u></b>	

**Monday, June 3, 2024 (continued)**  
 All Times Listed EDT

**Session**  
**No.**

1:15 – 2:00 PM	<p><b><u>The IND Document: Incorporating Preclinical Data</u></b>                  Noel J. Cusack, PhD</p> <ul style="list-style-type: none"> <li>• Structure of an Investigational New Drug (IND) Application</li> <li>• Incorporating Preclinical Data</li> </ul>	4
2:00 – 3:15 PM	<p><b><u>The IND Simulation Workshop</u></b>  <b><u>Part 1: Introduction &amp; Submission of IND</u></b>                  Participants and Faculty</p> <ul style="list-style-type: none"> <li>• Workshop Introduction</li> <li>• Group Work &amp; Faculty Consultations</li> </ul>	5
3:15 – 3:30 PM	<p><b><i>Afternoon break</i></b></p>	
3:30 – 4:45 PM	<p><b><u>The IND Simulation Workshop</u></b>  <b><u>Part 2: Group Presentations</u></b>                  Participants and Faculty</p> <ul style="list-style-type: none"> <li>• Group Presentations</li> <li>• Lessons Learned</li> </ul>	6
4:45 – 5:00 PM	<p><b><u>Course Day 1 Wrap-up</u></b></p>	



**Tuesday, June 4, 2024**  
**All Times Listed EDT**

8:00 – 8:30 AM	<b><u>Continental Breakfast</u></b>	
8:30 – 9:30 AM	<b><u>Clinical Development Plan Considerations – Part 1</u></b> Allan J. Weinstein, MD	<b>7</b>
	<ul style="list-style-type: none"> <li>• Goals of a clinical development program</li> <li>• Expectations of regulatory agencies</li> <li>• Phases and expectations of clinical research</li> <li>• Strengths and weaknesses of clinical Trials</li> <li>• Adaptive design</li> <li>• Bayesian statistics</li> <li>• Big Data</li> </ul>	
9:30 – 10:30 AM	<b><u>Clinical Development Plan Considerations – Part 2</u></b> Allan J. Weinstein, MD	<b>8</b>
	<ul style="list-style-type: none"> <li>• Study designs</li> <li>• Challenges of global development</li> <li>• The protocol</li> <li>• Epidemiology of disease</li> <li>• Good clinical practices</li> <li>• Ethical considerations</li> </ul>	
10:30 – 10:45 AM	<b><i>Morning Break</i></b>	
10:45 AM – 11:45 AM	<b><u>The NDA Document: Incorporating Clinical Data</u></b> Allan J. Weinstein, MD	<b>9</b>
	<ul style="list-style-type: none"> <li>• Structure and format of a New Drug Application (NDA)</li> <li>• Incorporating Clinical Data</li> </ul>	
11:45 AM – 1:00 PM	<b><u>The NDA Simulation Workshop</u></b> <b><u>Part 1: Introduction to NDA Simulation</u></b> Participants and Faculty	<b>10</b>
	<ul style="list-style-type: none"> <li>• Team building and role playing</li> <li>• FDA as interactive participants</li> <li>• Decision-making in the drug development process</li> <li>• The evolving product label</li> </ul>	
1:00 – 2:00 PM	<b><i>Lunch break</i></b>	
2:00 – 3:30 PM	<b><u>The NDA Simulation Workshop</u></b> <b><u>Part 2: NDA Assembly and FDA Interactions</u></b> Participants and Faculty	<b>11</b>
	<ul style="list-style-type: none"> <li>• The evolving product label</li> <li>• Consultant interactions</li> </ul>	
3:30 – 3:45 PM	<b><i>Refreshment Break</i></b>	

**Tuesday, June 4, 2024 (continued)**

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**No.**

3:45 – 4:45 PM

**The NDA Simulation Workshop**  
**Part 3: Label Submission to “FDA”**

Participants and Faculty

- Sponsor company NDA presentations to “FDA” panel
- Final product label
- Approval of indication
- Brand NAME
- Lessons Learned

12

4:45 – 5:00 PM

**Course wrap-up and adjournment**

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