Introduction to the Principles and Practice of Clinical Research (IPPCR)

October 16, 2012 – March 26, 2013

All sessions will meet on Monday and Tuesday evenings from 5:00 p.m. to approximately 6:30 p.m. (Eastern Standard Time) in the Lipsett Amphitheater.

Introduction	
Tuesday, October 16 Session 1	Welcome (30 minutes) John I. Gallin, M.D. Director NIH Clinical Center
	History of Clinical Research: A Merging of Diverse Cultures (30 minutes) John I. Gallin, M.D. Director NIH Clinical Center
	Chapter: 1
	Module I: Study Design and Statistics
Monday, October 22 Session 2	Unit 1: Overview of Clinical Study Design (1.5 hours) Laura Lee Johnson, Ph.D. Statistician Office of Clinical and Regulatory Affairs, NCCAM Chapter: 19
Tuesday, October 23 Session 3	Unit 2: Design of Epidemiologic Studies (1.5 hours) Laura Lee Johnson, Ph.D. Statistician Office of Clinical and Regulatory Affairs, NCCAM Chapter: 18
Monday, October 29 Session 4	Rescheduled for November 19 Chapter:
Tuesday, October 30 Session 5	Unit 3: Study Participant Selection (1 hours) Catherine Stoney, Ph.D. Program Director Prevention and Population Sciences Program, NHLBI Chapter: 2, 13, 19, 26

Monday, November 5 Session 6	Unit 4: Measures (1 hour) David Lukenbaugh, Ph.D. Biostatistician Experimental Therapeutics and Pathophysiology Branch, NIMH Chapter: 25, 26
Tuesday, November 6 Session 7	Unit 5: Secondary Data/Meta Analysis (1.5 hours) Charles Natanson, M.D. Senior Investigator and Head Anesthesia Section Critical Care Medicine Department, CC
Thursday, November 8 Session 8	Chapter: 27 Breakout Session – (1 hour) Title – TBD Laura Lee Johnson, Ph.D. Statistician Office of Clinical and Regulatory Affairs, NCCAM
Monday, November 12	FEDERAL HOLIDAY
Tuesday, November 13 Session 9	Unit 6: Issues in Randomization (1.5 hours) Laura Lee Johnson, Ph.D. Statistician Office of Clinical and Regulatory Affairs, NCCAM Chapter: 20, 24
Monday, November 19	Unit 7: Efficient Clinical Studies (1.25 hours) John Powers, III Senior Medical Scientist, NCI Frederick Chapter:
Tuesday, November 20	RECESS
Monday, November 26 Session 10	Unit 8: Overview of Hypothesis Testing (1.5 hours) Laura Lee Johnson, Ph.D. Statistician Office of Clinical and Regulatory Affairs, NCCAM Chapter: 21, 24
Tuesday, November 27 Session 11	Unit 9: Sample Size and Power (1.5 hours) Laura Lee Johnson, Ph.D. , Statistician Office of Clinical and Regulatory Affairs, NCCAM Chapter: 22, 24
Thursday, November 29 Session 12	Breakout Session – (1 hour) Title – TBD Laura Lee Johnson, Ph.D. Statistician Office of Clinical and Regulatory Affairs, NCCAM
Monday, December 3	Unit 10: Designing and Testing Questionnaires (45 minutes)
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Session 13	Bill Riley, PhD. Health Scientist Administrator, NCI
	Chapter:
	Unit 11: Quality of Life (45 minutes) John Ware, Ph.D.
	Chief Science Officer
	John Ware Research Group, Inc.
	Chapter: 25
Tuesday, December 4 Session 14	Unit 12: Conceptual Approach to Survival Analysis (1.5 hours) Laura Lee Johnson, Ph.D. Statistician
	Office of Clinical and Regulatory Affairs, NCCAM
	Chapter: 23, 24
Thursday, December 6	Breakout Session – (1 hour)
Session 15	Title – TBD Laura Lee Johnson, Ph.D.
	Statistician
	Office of Clinical and Regulatory Affairs, NCCAM
Monday, December 10	Unit 13: Using Large Datasets for Population-Based Health Research
Session 16	(1 hour)
	Leighton Chan, M.D. Chief, Rehabilitation Medicine Department, CC
	Chief, Renabilitation Medicine Department, CC
	Chapter: 28
Tuesday, December 11	Unit 14: Summary (1.5 hours)
Session 17	Laura Lee Johnson, Ph.D. Statistician
	Office of Clinical and Regulatory Affairs, NCCAM
	Chapter:
Module	e II: Ethical, Legal, and Regulatory Considerations
Monday, December 17	Unit 1: Ethical Principles in Clinical Research (45 minutes)
Session 18	Christine Grady, R.N., Ph.D.
	Head, Section on Human Subjects Research Chief, Bioethics Department, CC
	Chapter: 2
	Unit 2: Research with Vulnerable Participants (45 minutes)
	David Wendler, Ph.D.
	Head, Unit on Vulnerable Populations Section on Human Subjects Research, Bioethics Department, CC
	Chapter: 2, 5
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Tuesday, December 18 Session 19	Unit 3: Inclusion of Women and Minorities in Clinical Trials (1 hour)Miriam Kelty, Ph.D.Special Volunteer
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	Former Associate Director, Extramural Activities, NIA
	Chapter: 13
	Unit 4: Clinical Research from the Patient's Perspective (30 minutes) Jerry Sachs, B.A. Manager Guest Services (Retired) Smithsonian Museum of Natural History
	Chapter: 17
Monday, December 24	RECESS
Tuesday, December 25	RECESS
Monday, December 31	RECESS
Tuesday, January 1	RECESS
Monday, January 7 Session 20	Unit 5: Legal Issues in Clinical Research (1 hour) Carrie Pottker-Fishel, J.D. Attorney Advisor Office of General Counsel, NIH Chapter: 11
Tuesday, January 8 Session 21	Unit 6: FDA Product Regulation (1.5 hours) Bette Goldman, R.N., M.P.H. Senior Advisor on Clinical Issues to the Associate Director for Review Management, Center for Biologics Evaluation Research, FDA
Monday, January 14	Chapter: 7 Unit 7: Institutional Review Boards
Session 22	Jerry Menikoff, M.D., J.D. Director, Office for Human Research Protections Office of Public Health and Science, DHHS Chapter: 5, 6
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Tuesday, January 15 Session 23	Breakout Session: Mock IRB (2 hours) Jerry Menikoff, M.D., J.D. Director, Office for Human Research Protections Office of Public Health and Science, DHHS
	Chapter: 5, 6
Modu	le III: Preparing and Monitoring Clinical Studies
Monday, January 21	RECESS
Tuesday, January 22 Session 24	Unit 1: Information Resources for Clinical Research (1.5 hours) Josh Duberman, M.L.I.S. Informationist/Research Librarian, NIH Library
Monday, January 28 Session 25	Unit 2: Protocol Development (1 hour) Wendy Weber, N.D., Ph.D., M.P.H.

	Program Officer Division of Extramural Research, NCCAM Chapter: 29, 32
	Unit 3: Protocol Mechanics and Tools (30 minutes) Philip Lightfoot, B.S., B.A. Systems Analysis Department of Clinical Research Informatics, CC
	Chapter: 32
Tuesday, January 29 Session 26	Unit 4: Development of Manuals of Operating Procedures (1.5 hours) Wendy Weber, N.D., Ph.D., M.P.H. Program Director Division of Extramural Research, NCCAM
	Chapter: 29
Monday, February 4 Session 27	Unit 5: Pharmaceutical Development: Management of Projects (1 hour) Christopher Breder, M.D., Ph.D. Medical Officer Center for Drug Evaluation and Research, FDA
	Chapter: 7, 26, 37, 43
Tuesday, February 5 Session 28	Unit 6: Evaluation of a Protocol Budget (1.5 hours) Phyllis Klein, R.N., CCRC, BSN Director, Regulatory Support and Compliance Washington University in St. Louis
	Chapter: 33
Monday, February 11 Session 29	Unit 7: NIH Peer Review Process (1 hour) TBD
	Chapter: 36
Tuesday, February 12 Session 30	Unit 8: Design of Case Report Forms (1 hour) Diane St. Germain, R.N., M.S., C.R.N.P. Nurse Consultant Division of Cancer Prevention, NCI
M 1 F1 10	Chapter: 33, 37
Monday, February 18	FEDERAL HOLIDAY
Tuesday, February 19 Session 31	Unit 9: Data Management in Clinical Trials (1 hour) Diane St. Germain, R.N., M.S., C.R.N.P. Nurse Consultant Division of Cancer Prevention, NCI
	Chapter: 8
Monday, February 25 Session 32	Unit 10: Data & Non-Data Aspects of Quality Control in Clinical Studies (1 hour) Elizabeth Ness, RN, MSN Staff Development NCI/CCR
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	Chapter:
Tuesday, February 26 Session 33	Unit 11: Data and Safety Monitoring Committees (1 hour) Pamela Shaw, Ph.D. n Mathematical Statistician Biostatistics Research Branch, NIAID
	Chapter: 9
Monday, March 4 Session 34	Unit 12: Clinical Trial Registration and Results Reporting (1 hour) Deborah Zarin, M.D. Assistant Director for Clinical Research Projects Lister Hill National Medical Center for Biomedical Communications, NIH
	Chapter: 15
	Module IV: Miscellaneous Topics
Tuesday, March 5 Session 35	Unit 1: Technology Transfer (1.5 hours) Bruce Goldstein, J.D. Unit Coordinator Technology Transfer Branch, NCI
	Chapter: 30, 31
Monday, March 11 Session 36	Unit 2: Scientific Conduct (1 hour) James L. Gulley, M.D., Ph.D., F.A.C.P. Director Clinical Trials Group, Center for Cancer Research, NCI
	Chapter: 4, 12
Tuesday, March 12 Session 37	Unit 3: The Clinical Researcher and the Media (45 minutes) John Burklow, M.S. Associate Director for Communications Office of Communications and Public Liaison, NIH Chapter: 16
Monday, March 18	Unit 4: Health Disparities Research (1 hour)
Session 38	Irene Dankwa-Mullan, M.D., M.P. H. Acting Director Office of Innovation and Program Coordination, NIMHD
	Chapter: 46
Tuesday, March 19 Session 39	Unit 5: Dissemination and Implementation Research (1 hour) Catherine Stoney, Ph.D. Program Director Prevention and Population Sciences Program, NHLBI
	Chapter:
	Unit 6: Community-Based Participatory Research (30 minutes) Francisco Sy, M.D., Dr PH Director

	Division of Extramural Activities and Science Programs, NIMHD Chapter: 46
Monday, March 25 Session 40	Unit 7: Team Science Howard Gadlin, Ph.D. NIH Ombudsman Director, Center for Cooperative Resolution, NIH Michelle Bennett, Ph.D. Deputy Scientific Director Division of Intramural Research, NHLBI