University of Nicosia, Cyprus

Course Code	Course Title	ECTS
BIOL-473	Special Topics IV: Clinical	4
	Research	
Department	Semester	Prerequisites
Life and Health	Spring/Fall	BIOL-371 Drug Action and
Sciences		Toxicology
Type of Course	Field	Language of Instruction
Elective	Biology, Medicine	English
Level of Course	Year of Study	Lecturer
1 st Cycle	4^{th}	Dr. Theodoros Kyprianou
Mode of Delivery	Work Placement	Co-requisites
Face-to-face	N/A	None

Objectives of the Course:

The course aims to provide an introduction to the processes for drug approval through clinical research. The main objectives of the course are to:

- Make students aware of the clinical research process and research terminology used.
- Discuss the basic study designs in health and biomedical setting.
- Review the four phases of interventional trials and the requirements for drug approval.
- Make students aware of regulations, responsibilities and ethical issues related to clinical trials.

Learning Outcomes:

After completion of the course students are expected to:

- 1. Explain the steps required for a new drug to be approved for human use.
- 2. Identify the four major clinical phases in the drug development process and explain the rational for each.
- 3. Describe the basic concepts of clinical pharmacology requirements in drug testing.
- 4. Name the sources for finding the rules and regulations for new drug development.
- 5. Describe the roles of various research and regulatory organizations involved in the clinical research process and the approval of new drugs.
- 6. Discuss the real and perceived conflicts and ethical issues in drug testing.

Course Contents:

- 1. Drug development introduction.
- 2. Pre-Clinical drug development.
- 3. Principles of Clinical Pharmacology
- 4. Clinical trial designs; advantages and disadvantages

- 5. Clinical phases
- 6. Drug safety and monitoring Boards
- 7. Clinical trials reporting; meta analyses studies
- 8. Dangers of Biomedical research
- 9. Biotechnology in Drug Development.
- 10. Clinical Research ethics

Learning Activities and Teaching Methods:

Lectures, Case studies discussions, Literature reviews.

Assessment Methods:

Individual assignments, Cooperative presentations; Final Exam

Required Textbooks/Reading:

Authors	Title	Publisher	Year	ISBN
1. Grahame- Smith, DG.	Clinical Pharmacology and drug therapy.	Oxford textbook	2000	ISBN – 0192616757
2. Adam Cohen and John Posner	A guide to clinical drug research	Kluwer Academic Publishers	1995	ISBN-: 0792361717

Recommended Textbooks/Reading:

Authors	Title	Publisher	Year	ISBN
1. Atkinson AJ	Principles of Clinical	Academic	2007	ISBN-
Jr et al.	Pharmacology	Press	2 nd Ed.	0123694175
2. Mark A.	Pharmacogenomics:	Wiley,	2006	ISBN:
Rothstein	social, ethical, and clinical dimensions			0471737798
	chilical difficultions			ICDM
3. Kenneth P.	Brody's human	Elsvier Mosby	2005, 4 th	ISBN: 032303286
Minneman, Lynn	pharmacology:		ed.	032303200
Wecker, Joseph	molecular to clinical			
Larner, Theodrore				
M. Brody				

http://www.fda.gov/fdac/special/newdrug/ndd_toc.html

http://www.fda.gov/oc/initiatives/criticalpath