Spring 2015

BIOS 9132 - Advanced Clinical Trial Methodology

Karl E. Peace
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Syllabus: BIOS 9132 – Advanced Clinical Trial Methodology  
Jiann-Ping Hsu College of Public Health  
Georgia Southern University  
Hendricks Hall, PO Box 8015  
Statesboro, GA 30460

**Instructor:** Karl E. Peace  
**Office:** 1005 Hendricks Hall  
**Phone:** 912-478-7905  
**E-Mail Address:** [kepeace@georgiasouthern.edu](mailto:kepeace@georgiasouthern.edu)  
**Office Hours:** Wednesday– 3:00 PM – 5:00 PM  
Other times by appointment:  
Students are encouraged to make frequent use of email contact where each question will be responded via return email  
**Class Meets:** Wednesday – 5:00 AM-to-7:45 PM

**Prerequisites:** Clinical Trial Methodology, Statistical Issues in Drug Research and Development, or by permission of instructor.

**Catalog Description:** Students are introduced to regulatory, scientific, statistical and practical aspects of methods inherent in design, monitoring and analyzing clinical trials. Clinical trials in many areas of drug development are presented, discussed and critiqued.

**Required Textbook:** No textbook required. The course is presented using power points developed by the professor. Students are provided copies of the power points on a flash drive.

**Secondary Texts:**


**Course Objectives:** At the end of this course, students will be able to:

1. Explain the requirements for good protocol development for biomedical research clinical trials and develop the statistical analysis section of such protocols
2. Describe methodological alternatives to commonly used statistical methods used in biomedical research clinical trials when analysis assumptions are not met, and describe prerequisites for validity of inference from clinical trials
3. Interpret results of statistical analyses of data collected from biomedical clinical trials
4. Develop written and oral presentations based on statistical analyses of biomedical research clinical trials, for both biomedical research professionals and educated lay audiences
5. Identify key federal regulations ‘governing’ the conduct of clinical trials
6. Discuss the Ethics of Clinical Trial Research
7. Describe the components of Population Nonlinear, Mixed Effects Modeling
8. Design, analyze and interpret results of bioequivalence, cancer and non-inferiority clinical trials
9. Describe the issues in group sequential clinical trials and in subset analyses
10. Describe ‘intention to treat’ and its impact on inference when data are missing and methods of imputing missing data

**Instructional Methods:** Class meetings will be a combination of lecture and class discussion. Approximately half of the class meetings will be facilitated via Adobe Connect in real time (blended format) and the remainder physically in the classroom. Homework assignments – including a research project that requires class participation and the final examination constitute the basis of student evaluation. Students are expected to make use of ample office hours to discuss concepts or difficulties they may have.

**Daily Study Log:** Students are required to keep a daily computerized study log. The study log should have a column for the date, a column to identify topic of study, a column to identify the time of beginning study, a column to identify the ending time of study, and a column to identify the amount of time spent in studying the topic.

**Grading:** Weighting of assignments for purposes of grading will be as follows:
The following point scale will be utilized in grading:

- 90% - 100% A
- 80% - 90% B
- 70% - 80% C
- 60% - 70% D

There are times when extraordinary circumstances occur (e.g., serious illness, death in the family, etc.). In such circumstances, and/or if you need additional time to satisfactorily complete any course requirement, please consult with the instructor within a reasonable amount of time.

**Academic Misconduct:** As a student registered at this University, it is expected that you will adhere to only the strictest standards of conduct. Your continued enrollment in this course is an implied contract between you and the instructor on this issue; from this point forward, it is assumed that you will conduct yourself appropriately.

Academic integrity relates to the appropriate use of intellectual property. The syllabus, lecture notes, and all materials presented and/or distributed during this course are protected by copyright law. Students are authorized to take notes in class, but that authorization extends only to making one set of notes for personal (and no other) use. As such, students are not authorized to sell, license, commercially publish, distribute, transmit, display, or record notes in or from class without the express written permission of the instructor.

**Plagiarism:** Plagiarism includes (but is not limited to): A. Directly quoting the words of others without using quotation marks or indented format to identify them. B. Using published or unpublished sources of information without identifying them. C. Paraphrasing material or ideas without identifying the source. D. Unacknowledged use of materials prepared by another person or agency engaged in the selling of term papers or other academic material.

**Attendance Policy:** Federal regulations require attendance be verified prior to distribution of financial aid allotments. Students are expected to attend all classes, whether taking for credit or auditing (Instructor will permit missing 1-2 classes for valid reasons).

**One Final Note:** The contents of this syllabus are as complete and accurate as possible. The instructor reserves the right to make any changes necessary to the syllabus and course material. The instructor will make every effort to inform students of changes as they occur. It is the responsibility of the student to know what changes have been made in order to successfully complete the requirements of the course.

**Advanced Clinical Trial Methodology Content to be Covered During the Semester:**
Module I: An Overview of the Regulation of Pharmaceuticals

Module II: An Overview of the Processes of Discovery, Basic Research, Clinical Development and Manufacturing in Pharmaceutical Development

Module III: Biostatistical Aspects of Clinical Drug Development
   A. The Components of a Protocol
   B. Statistical Analysis Section of a Clinical Trial Protocol
   C. The Statistical Analysis Plan

Module IV: Ethics of Clinical Trial Research

Module V: Validity of Statistical Inference

Module VI: Bioavailability and Bioequivalence

Module VII: Biosimilarity

Module VIII: Population Nonlinear, Mixed Effects Modeling of Primary Efficacy Endpoint in Enrichment Trials of Alzheimer’s disease

Module IX: Biostatistical Aspects of the Design of Cancer Trials

Module X: Biostatistical Aspects of the Analysis and Interpretation of Cancer Clinical Trials

Module XI: Interim Analyses: p-Value and Power Computations in Group Sequential Trials

Module XII: Design and Analysis of Non-inferiority Clinical trials

Module XIII: Statistical Analysis of Dose Response Trials

Module XIV: Safety Assessment in Clinical Trials

Module XV: Subgroup Analyses in Clinical Trials from a Causal Inference Viewpoint

Module XVI: Statistical Paradigms and Methodologies for Clinical Development

Module XVII: Intention-to-Treat and Inferential Impact of Missing Data

Module XVIII: Methods for Imputing Missing Data

Module XIX: Overview of Meta-Analyses
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