- A. The Medical Dictionary for Regulatory Activities (MedDRA) for coding medical history and adverse events
- B. WHO Drug dictionary for coding drugs
- C. EudraVigilance Medicinal Product Dictionary (EVMPD
- IV. Discuss the development, maintenance and use of the safety database
  - A. Key components of the safety database
  - B. Differentiate the clinical studies database and the safety database
  - C. Data reconciliation between systems
  - D. Discuss the setup of different systems
  - E. MSSO: Standardized MedDRA Queries
  - F. Monitoring of "Medically Important" predefined events
  - G. Trend Analysis
- V. Understand the functions needed for a compliant product safety/pharmacovigilance department
  - A. ICSR Processing
  - B. Aggregate Reporting
  - C. Signal Detection
  - D. Risk Management
  - E. Quality Management
  - F. Training
  - G. Compliance
  - H. QPPV Office (EU)

The textbook and other instructional material will be determined by the instructor.