

- B. The supervisor will arrange for a quiet place to provide feedback to the student on an individual basis, ensuring privacy and confidentiality.
- C. The supervisor will identify one project that the student can observe for at least two sessions, so that the student can adequately accomplish their fieldwork assignments.
- D. The supervisor will provide the opportunity for the student to observe and participate in a MPSP research project.
- E. Pharmaceutical company or contract research organization experiences may include, but not be limited to data entry, case processing, writing case narratives, preparing safety reports, etc.

STUDENT'S RESPONSIBILITIES:

- A. Students are responsible for confirming their fieldwork with the clinical site supervisor at least one week prior to the scheduled time to determine hours, dress code, materials needed, location of the site facility and directions to the initial meeting place.
- B. During the first session, students should review their individual objectives and assignments of the fieldwork experience with their supervisor.
- C. The student will identify the specific type MPSP project in place at the specific fieldwork site and identify the roles of specific disciplines involved in the coordination of clinical research projects.
- D. The student will receive information from the site supervisor regarding:
 - i. The pharmaceutical drug or medical device that is the subject of the safety report
 - ii. The known effects of the pharmaceutical drug or medical device, both beneficial and adverse
 - iii. The phase of development in which the product is currently being researched
 - iv. The regulatory status of the product
 - v. The protocol design and objectives of the study
 - vi. Members and roles of the MPSP team
- E. The student will observe the supervisor while interacting in meetings, collecting relevant data, and performing study-required tasks.
- F. During the clinical research industry site rotation, the student will observe and participate in, as appropriate, planning meetings, medical monitor site visits, case report processing and review, and other administrative responsibilities related to the MPSP project.