The Center for Clinical Research (CCR) assists faculty in designing and completing studies to assess new treatments and to improve patient health outcomes and the delivery of health care.

The CCR is a comprehensive support system for your clinical investigation. Our faculty and staff can assist with clinical research design and methods, protocol development, grant submission, statistical analysis, data management and study coordination. We bring expertise in project management, study preparation, regulatory document preparation and submission, study coordination, monitoring and data management and analysis.

The CCR administrative office is located on the second floor of SIU School of Medicine's Administrative Offices Building, 201 E. Madison Street. We are centrally located with our affiliated teaching hospitals, HSHS St. John's Hospital and Memorial Medical Center, and with clinical studies being conducted across the SIUSOM campus.

Visit siumed.edu/ccr for more detailed information on our cores and services, or call 217-545-9700.
The central concept of the CCR is to support faculty and their research teams using a flexible offering of services; either by utilizing specific study services or optioning for full-service support.

STATISTICS & INFORMATICS
The Statistics & Informatics Division can support the clinical, basic science, educational and administrative research needs for your project. Our team provides statistical consulting for design, power analysis/sample size estimates, result analysis and interpretation.

We can help with database creation, data acquisition, management, storage and analysis of health system, governmental or other large databases for health outcomes research. Staff can also assist with entry, coding and de-identification to ensure compliance with regulatory guidelines and maintenance of patient confidentiality.

SCHOLARLY ACTIVITY SUPPORT
The Scholarly Activity Support Division focuses on the development of faculty-directed, investigator-initiated clinical research. Its services include clinical research education, training, and career development for clinical faculty, fellows, residents, medical students, and research teams.

Our teams can assist investigators with research design, protocol development and completion of Investigational New Drug/Investigational Device Exemption (IND/IDE) applications. The division also provides assistance with grant development, preparation and submission, in addition to clinical study registration on ClinicalTrials.gov, a federal requirement.

As the study winds down, Scholarly Activity Support can ramp up, helping with manuscript, abstract and poster/presentation development.

REGULATORY SUPPORT
CCR provides IRB regulatory support for clinical trials to Principal Investigators, Sub-Investigators, residents and faculty including the collection of feasibility questionnaires and initial study start-up materials, negotiation of study budgets, preparation of mandatory FDA regulated clinical research forms, submission of protocol updates and amendments, development of informed consent documents and maintenance of study specific site binders.

Clinical Trial Contracting
All investigators are required to contact the CCR to initiate clinical trial contract negotiations. The CCR manages the administration, processing and negotiation of confidential disclosure agreements (CDAs) and clinical trial agreements (CTAs) from pharmaceutical and biotech companies that sponsor clinical research conducted by SIU faculty at SIU and its affiliated hospitals.

Institutional Review Board
The Springfield Committee for Research Involving Human Subjects (SCRIHS) is the Institutional Review Board (IRB) for the School of Medicine, its affiliated hospitals and other community sites. All research involving collection of data from or about human subjects must be submitted to the IRB for review. Contact: scrihs@siumed.edu, 217-545-7602, or visit siumed.edu/adrfa/SCRIHS.html.

OPERATIONS
The Operations Division assists in planning, initiating and conducting clinical trials. Team members work closely with sponsors, investigators and the IRB to facilitate successful clinical trials that meet all regulatory requirements governing human subject research.

Operations personnel can provide the infrastructure for sponsored and investigator-initiated clinical trials, including assistance with patient screening and consenting, as well as budgeting, billing, and financial management. Division research nurses and coordinators are available to conduct study start-up procedures, assist in enrolling subjects and follow-up visits, and the close-out of each clinical trial.

DIRECTIONS
From the north or south via Interstate 55: Take exit 98-B (Clear Lake Avenue) and proceed west on Clear Lake, which becomes Jefferson Street. Turn right at Second Street and proceed north to the corner of Second and Madison Streets. Visitor parking is located on the southeast side of the building just off of Madison Street.