Design Study

Learn About Feasibility and Regulatory Requirements

Cohort Finding

Evaluate Study Feasibility

IRB Approval

Regulatory Support and Resources

Protocol Development

Consent Development

FDA Device (IDE) Submission

FDA Drug or Biologics (IND) Submission

Scientific Review

Explore Funding Opportunities & Budget Preparation

Identify Funding Opportunities

Proposal Preparation and Submission

Data Management Planning

Set Up Research Budgets

Overview of Coverage Analysis

Billing Codes and Research Pricing Information

CTSI Integrating Special Populations Program (ISP)

Seek Study Design Assistance & Resources

Request a CTSI Ethics Consultation

Research Data Management Best Practices (Consultation)

Review Project for HIPAA, Privacy & IT Security Requirements

Biostatistics - Information and Consultation

Study Design - Information and Consultation

Clinical Data Related to Research - Electronic Health Record