

Our Services

We offer the following services:

- Study and Site Management
- Clinical Trials Monitoring and Auditing
- Clinical Trials Project Management
- Investigator Site Feasibility and Selection
- Regulatory affairs
- Clinical Research Training
- Health Systems and Outcomes Research
- Contract Laboratory services management
- Pharmacovigilance services
- Strategic Consulting services

Our Experience

Our team has extensive experience in infectious diseases, non-communicable diseases Projects Management in resource limited settings in Sub Saharan Africa and Asia countries (India and Cambodia).

We offer Phase I - IV capabilities as full service project based or Functional Service Resourcing model.

Our Team

Our team is comprised of :

- Clinical Trials Monitors and Auditors,
- Public Health and Epidemiologists
- Projects and Data Managers,
- Laboratory Scientists,
- Physicians/Medical Doctors,
- Contracts and Grants Managers
- Statisticians,
- Social Scientists
- Qualified Persons

About CLINWIN RESEARCH

About us

We are mid-sized Contract Research Organization (CRO), providing outsourced clinical development and Consulting services to biopharmaceutical companies, Academic Medical Centres, Investigator sites, not-for profit research organizations and Government.

Our Vision

To deliver quality and efficient services on time and cost to our clients.

Our Mission

To partner with biopharmaceutical, academia and CRO clients to support the successful outcome of there projects.

Our Values

We advance our clients' assigned projects through integrity, teamwork, quality and accountability

Our Principles

Efficiency, Quality and Ethics

Our Contacts

ClinWin Research Services

Lenwood Apartments, Lenana Road
P O Box 3289

Nairobi 00200 Kenya

Phone:+254(0)721 515 009

+254(0)790089440

info@clinwinresearch.com

clinwin.research@gmail.com

www.clinwinresearch.com



Efficiency .Quality. Ethics

Clinical Trials Monitoring and Site Management



- Feasibilities and site selection
- Study start up and contracts negotiations
- Regulatory and ethical submission support
- Site Initiation, Interim Monitoring
- Medical Monitoring
- Site Operations
- Patient Recruitment
- Data Collection and Management

Projects Management

- Projects feasibilities
- Risk assessment and Management
- Regulatory and ethical submission
- Project team management
- Resources planning and management
- Milestones tracking and reporting
- Budgets and Contracts management
- Project Communication Planning
- Quality management
- Projects Monitoring and Evaluation

Strategic Consulting

We offer our clients innovative and cost efficient solutions to make informed business decisions. These include:

- Product development advisory services
- Protocol design and feasibility
- Operational efficiency and effectiveness
- Patient recruitment and retention services
- Risk and Compliance Operations
- Productivity and Performance evaluations
- Contracts Negotiation and Legal Affairs
- Clinical Resourcing and Recruitment
- Document Translation service
- Market research and access

Health Systems Research

- Health Services effectiveness
- Social Science Research,
- Health Policy analysis,
- Health Services delivery,
- Health care organizational development
- Surveys and program evaluation



Clinical Research Training



We offer face to face and on-line courses to investigator site and Sponsor staff. The courses can be tailored as per client needs on all aspects of clinical research enterprise. This include:

- Introduction to Drug Development
- Clinical Research Principles and Practice
- Research Project planning and Management
- Good Clinical Practice (ICH-GCP) for beginners and refresher courses
- Good Clinical Laboratory Practice (GCLP)
- Internal quality assurance
- Clinical Research Ethics
- Infectious Diseases
- Data Management
- Regulatory affairs
- Clinical Research Methodology
- Projects Management