

## Our Services

We offer outsourced services in:

- 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 Experts 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

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**ClinWin** Research Services

Efficiency. Quality. Ethics



**Clinical Trials Monitoring and Consulting**



**Biopharmaceutical**

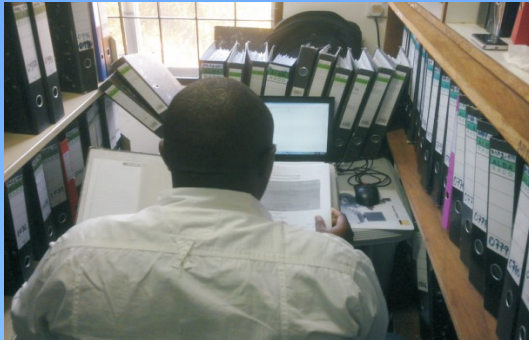


**Technology**



**Data and Analytics**

## 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

## 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:

- Entry and Advanced Level Clinical Trials Monitoring Course
- 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