



R 4,100 excl Vat

Discounts apply to groups



eLearning -Online



Average of 24 hours

The course is designed to provide a practical context to help clinical research professionals learn more about conducting and coordinating clinical trials. Staff who are new to the research environment will also find the course beneficial, as it provides a comprehensive overview of the role of a clinical research coordinator and various site activities.

The course has 4 modules including a "Test your knowledge" at the end of each module:

- Introduction
- Module 1: Introduction to Clinical Research
- Module 2: The role of the Clinical Research Coordinator during the pre-study phase
- Module 3: The role of the clinical research coordinator during the study conduct phase
- Module 4: The role of the clinical research coordinator during the study conduct phase



At the end of the course learners will have a good understanding of;

- The multifaceted workings of a clinical trial in a research environment
- The responsibilities as clinical research coordinator in accordance with GCP
- Practices that ensure effective, efficient and ethical study conduct.



At the end of this course, learners will be expected to complete a final assessment:

- There are 50 multiple choice questions
- The pass mark is 70%
- Learners have (2) attempts
- A certificate will be issued on passing the assessment
- CPD Points will be issued on completion



Clinical Research Clinicians, Nurses and Professionals.



Coaching Services



