

# Biostatistics for Clinical Trials Data Analysis Training Course

Professional Development Training Course Outline

|          |                            |               |                         |
|----------|----------------------------|---------------|-------------------------|
| Category | Research and Data Analysis | Duration      | 5 Days                  |
| Base Fee | \$1,500 USD                | Delivery Mode | Online / On-site Hybrid |

## Course Introduction

Clinical trials often deal with sensitive topics such as mental health, reproductive health, infectious diseases, substance use, and stigmatized conditions. Designing research and conducting data analysis in such trials requires a specialized understanding of biostatistics, ethical considerations, data privacy, and regulatory compliance. Biostatistics for Clinical Trials Data Analysis Training Course equips participants with cutting-edge statistical tools and methodologies tailored for sensitive research contexts, ensuring scientific integrity, participant safety, and actionable insights.

With increasing demand for ethical, evidence-based approaches in public health and biomedical research, this course integrates advanced biostatistical techniques, real-world clinical data analysis, and robust frameworks for handling sensitive variables. Participants will explore best practices, case studies, and data handling protocols that align with Good Clinical Practice (GCP), ICH guidelines, and data protection laws such as GDPR. This training is ideal for researchers, data analysts, clinicians, and regulatory professionals committed to responsible and rigorous research in high-stakes environments.

## Programme Curriculum

### Biostatistics for Clinical Trials Data Analysis Training Course

#### Introduction

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### Course Objectives

1. Understand biostatistical principles used in sensitive clinical trial research.
2. Analyze longitudinal clinical data using modern statistical modeling techniques.
3. Apply ethical frameworks in the design of studies involving vulnerable populations.
4. Conduct risk-adjusted statistical analysis for stigmatized health conditions.
5. Ensure data confidentiality and anonymization in sensitive health datasets.
6. Implement robust sampling strategies in low-consent environments.
7. Utilize advanced statistical software (e.g., R, SAS, STATA) for sensitive data.
8. Interpret missing data mechanisms in clinical trials with delicate subject matter.
9. Address bias and variability in multi-site sensitive research settings.
10. Design inclusive statistical protocols for underrepresented groups.
11. Evaluate data quality metrics in ethically challenging research.
12. Report findings with transparency, reproducibility, and cultural sensitivity.
13. Integrate machine learning tools for predictive analytics in sensitive topic trials.

### Target Audiences

1. Clinical Researchers
2. Biostatisticians
3. Public Health Officers
4. Medical Data Analysts
5. Ethics Review Committee Members
6. Healthcare Policy Makers
7. Medical Journal Reviewers
8. Regulatory Affairs Professionals

**Course Duration:** 5 days

### Course Modules

#### Module 1: Foundations of Biostatistics for Sensitive Research

- Key statistical concepts and distributions
- Ethical challenges in sensitive-topic trials
- Introduction to statistical software (R, SAS)
- Understanding informed consent in vulnerable populations
- Handling small sample sizes and variability
- **Case Study:** Biostatistical analysis in HIV stigma trials

#### Module 2: Designing Ethical and Inclusive Clinical Trials

- Trial design for high-risk populations
- Bias control in sensitive-topic studies
- Stratified and cluster sampling techniques
- Managing regulatory requirements (ICH-GCP)
- Cultural competency in protocol design

- **Case Study:** Trial design for reproductive health interventions

### **Module 3: Managing and Analyzing Missing Data**

- Identifying missing data patterns (MCAR, MAR, MNAR)
- Imputation techniques for sensitive datasets
- Use of multiple imputation in R/SAS
- Evaluating sensitivity analyses
- Protecting against data manipulation
- **Case Study:** Substance use trial with high attrition rates

### **Module 4: Data Privacy and Anonymization Techniques**

- Legal frameworks (GDPR, HIPAA)
- Methods of de-identification and re-identification risks
- Privacy-preserving data analysis techniques
- Safe data sharing and access controls
- Metadata management in sensitive trials
- **Case Study:** Mental health dataset privacy audit

### **Module 5: Advanced Statistical Models in Sensitive Research**

- Mixed-effects models for longitudinal data
- Generalized estimating equations (GEE)
- Bayesian approaches for small sample sizes
- Hierarchical modeling for multi-level data
- Survival analysis for time-to-event outcomes
- **Case Study:** Multi-center PTSD clinical trial

### **Module 6: Machine Learning & Predictive Analytics in Clinical Trials**

- Introduction to ML in health research
- Risk prediction models for vulnerable groups
- Handling algorithmic bias and fairness
- Feature selection from sensitive variables
- Integration of ML in trial monitoring
- **Case Study:** ML-based prediction of relapse in addiction treatment

### **Module 7: Communicating Results with Sensitivity**

- Creating culturally appropriate visualizations
- Reporting adverse events in stigmatized contexts
- Journal writing for sensitive-topic trials
- Effective data storytelling techniques
- Addressing audience bias and interpretation risks
- **Case Study:** Reporting gender-based violence data in journals

### **Module 8: Case Reviews and Capstone Project**

- Recap of sensitive trial principles
- Real-world data set group analysis
- Presentations and peer reviews
- Cross-cultural insights from global case studies
- Final assessment and feedback session
- **Case Study:** Capstone project on sensitive disease modeling

### **Training Methodology**

- Interactive expert-led sessions
- Real-time data lab simulations
- Group-based problem-solving activities
- Peer-reviewed project presentations
- Practical hands-on with real datasets

### Register as a group from 3 participants for a Discount

Send us an email: [info@fineskilltrainingcenter.org](mailto:info@fineskilltrainingcenter.org) or call +254769199797

### Certification

Upon successful completion of this training, participants will be issued with a globally- recognized certificate.

### Tailor-Made Course

We also offer tailor-made courses based on your needs.

### Key Notes

- The participant must be conversant with English.
- Upon completion of training the participant will be issued with an Authorized Training Certificate
- Course duration is flexible and the contents can be modified to fit any number of days.
- The course fee includes facilitation training materials, 2 coffee breaks, buffet lunch and A Certificate upon successful completion of Training.
- One-year post-training support Consultation and Coaching provided after the course.
- Payment should be done at least a week before commence of the training, to Fineskill Training Center account, as indicated in the invoice so as to enable us prepare better for you.

## Upcoming Scheduled Sessions

| Start Date  | End Date    | Location | Price   |
|-------------|-------------|----------|---------|
| 21 Sep 2026 | 25 Sep 2026 | Online   | \$1,000 |
| 21 Sep 2026 | 25 Sep 2026 | Physical | \$1,500 |
| 21 Sep 2026 | 25 Sep 2026 | Physical | \$0     |
| 21 Sep 2026 | 25 Sep 2026 | Physical | \$0     |
| 21 Sep 2026 | 25 Sep 2026 | Physical | \$0     |
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| 21 Sep 2026 | 25 Sep 2026 | Physical | \$0   |
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Ready to enroll or request a customized program? Book online or contact us:

Register Online Now

Email: [info@fineskilltrainingcenter.com](mailto:info@fineskilltrainingcenter.com) | Website: [www.fineskilltrainingcenter.com](http://www.fineskilltrainingcenter.com)