

# Good Clinical Practice (GCP) Training for Investigators and Study Teams

## PROGRAMME

### Day 1:

---

#### Registration Day 1

**08:15** Registration Opens

---

#### General | Welcome

**08:45** Introduction of participants, course program and structure

---

#### Lecture 1 | Introduction to Good Clinical Practice

**09:00** Historical Background, Research Ethics, Drug Regulations, Introduction to ICH E6 (R2) GCP Guideline, Investigators responsibilities

**10:15** Coffee Break

---

#### Lecture 1 | Introduction to Good Clinical Practice (continuation)

**10:30** Historical Background, Research Ethics, Drug Regulations, Introduction to ICH E6 (R2) GCP Guideline, Investigators responsibilities

---

#### Lecture 2 | Basic Statistical Concepts and Study Design

**11:00** Defining the research question

**12:00** Lunch Break

---

#### Exercise 1 | Protocol Optimization

**13:00** How would you improve the example protocols?

---

#### Lecture 3 | Source Records and Data Management

**13:45** From Case Report Forms to Final Analysis

**14:45** Coffee Break

---

#### Exercise 2 | Critical review of Source Documents and CRF Design

**15:00** Principles of ALCOAC; Improving Data Quality from the Start

---

#### Lecture 4 | Study Documents and Archiving

**16:00** Essential Documents, Trial Master File, Retention

---

**17:00** End of Day 1

# Good Clinical Practice (GCP) Training for Investigators and Study Teams

## PROGRAMME

### Day 2:

---

#### Lecture 5 | Focus on Informed Consent and Safety Reporting

09:00 Investigator Responsibilities

10:15 Coffee Break

---

#### Exercise 3 | Critical Review of an Informed Consent Form

10:30 Analyze and discuss potential problems and ethical issues with the example informed consents

---

#### Lecture 6 | Quality Management

11:30 SOPs, Training, Monitoring, Auditing; Common Audit and Inspection Findings

12:30 Lunch Break

---

#### Lecture 7 | Research Misconduct and Fraud

13:30 Definition and Consequences; Case Examples

---

#### Lecture 8 | Legal Background, Ethics Committees & Submission processes

14:15 Introduction to HFG, ethics landscape Switzerland, BASEC, Swissmedic

---

#### Summary |

15:30 13 Key Principles of ICH GCP; A look at addendum to ICH E6 (R2)

---

16:00 End of Day 2 / End of Course

### Information:

**Additional time:** Prior to the course, participants will be expected to have read the “**ICH Harmonised Guideline – Integrated Addendum to ICH E6(R1): Guideline for Good Clinical Practice E6(R2)**” and the “**Declaration of Helsinki - Ethical Principles for Medical Research Involving Human Subjects**” (*Version October 2013, Fortaleza*) (appr. 3 hours).

**After the course, participants will need to pass a competency assessment** (multiple choice consisting of 20 questions) **with a minimum passing score of 70%**. Following successful completion, a Certificate of Training will be provided.