



GCP Refresher:
ICH Guideline E6(R3) (ICH-GCP)
CTR / CTIS
4. AMG Änderungsgesetz und Medizinforschungsgesetz (MFG)
Aktuelle Fragen zu GCP

Referentin: Dr. Dagmar Chase

ICH Guideline E6(R3) (ICH-GCP Principles and Annex I)

- History of ICH E6
- New Structure, Scope and Summary of Changes
- Principles of ICH GCP
- Investigators, in particular
 - Responsibilities
 - Protocol Deviations
 - Participant Medical Care and Safety Reporting
 - Informed Consent of Trial Participants
 - Investigational Product Management
 - Records (Source and Case Report Forms)
- Sponsors, in particular
 - Agreements
 - Sponsor Oversight
 - Risk-based Quality Management / Risk-based Monitoring
 - Safety Assessment and Reporting
 - Investigational Product
 - Data and Records
 - Reports
- Data Governance – Investigator and Sponsor
- Essential Records (TMF / eTMF)

EU Clinical Trials Regulation 536/2014 and CTIS

- New Concepts
- Submissions and Authorisation Procedure
- CTIS Roles and Permissions
- Safety Reporting
- Revised Transparency Rules

4. AMG Änderungsgesetz und Medizinforschungsgesetz (MFG)

Aktuelle Fragen zu GCP

- Protocol Waivers
- Unblinding