



## EU Clinical Trial Regulation 536/2014 (CTR)

### Clinical Trial Information System (CTIS)

## 4. AMG Änderungsgesetz und Medizinforschungsgesetz (MFG)

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### CTR and CTIS

- Actual Status
- New Concepts
  - Co-sponsoring
  - Low-intervention clinical trials
- Submission and Authorization Procedures
  - Responsibilities
  - Timelines
- CTIS
  - Overview and Interface to other EMA Data Sources (OMS, xEVMPD)
  - Sponsor Functionality (Workspace)
  - User and Access Management (Roles and Permissions)
  - Organization Centric vs Trial Centric Approach
- Application Dossier (Part I, Part II)
  - Language Requirements
  - Differences between Member States
  - Patient Facing Documents
- Substantial Modification / Non-substantial Modifications / Subsequent Addition of a MS concerned
- Safety Reporting: What is new?
- Revised EU Transparency Rules
- How to protect Personal Data and Commercially Confidential Information when using CTIS?

## 4. AMG Änderungsgesetz und Medizinforschungsgesetz (MFG)

- Verfahren zur Genehmigung einer klinischen Prüfung
- Ethik-Kommissionen in Deutschland
- Verfahrensordnung und Geschäftsverteilungspläne
- Weitere Änderungen in der klinischen Forschung durch das Medizinforschungsgesetz (MFG)