



# **Project Management in Clinical Trials**

## **Working with CROs**

### **Sponsor Oversight**

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#### **Project Management – Overview**

- **Definitions**
- **Hard Factors – Soft Factors**
- **Quality**
- **Time Management**
  - **Work break down structure**
  - **Critical path**
- **Budget / Resource Planning**
- **Project Controlling (The PDCA Cycle)**

#### **Risk-based Quality Management in Clinical Trials including Project Plan and Risk Mitigation Plan**

#### **Sponsor Oversight**

#### **Working with CROs (see next page)**



# Project Management in Clinical Trials

## Working with CROs

### Sponsor Oversight

(Fortsetzung)

## Working with CROs

### General Aspects

- Regulatory Framework
- Typical CRO and Clinical Trial Team Structure
- CRO Selection Process and Contracts: What Inspectors Expect

### Trial Start-up

- Kick-off Meeting
- Preparation of Trial Documents
- Site Selection and Site Approval Process
- Training of the Trial Team (Sponsor / CRO)
- Setting up Trial Files

### Trial Conduct

- CRO Oversight and Key Performance Indicators
- How to Handle Out of Scope Costs?
- Sponsor-CRO Relationship
  - What can go wrong?
  - How to manage a crisis?
  - Developing a good relationship

### Trial Completion

- Report Writing
- Returning Trial Material
- Archiving Duties
- How to Measure CRO Performance at Trial End?
- Lessons Learned