

BASIC TRAINING

Introduction of CTIS and user management

- Introduction to the Clinical Trial Regulation No 536/2014
 - Purpose of CTR
 - CTIS development/transition period and implementation calendar
 - Differences CT Directive vs CT Regulation
- CTIS overview
 - Key benefits
 - Workspaces + interactions with other systems
 - Main common functionalities
 - Publication aspects: transparency and GDPR
- CTIS User access management
 - Access/Registration
 - User roles hierarchy:
 - Default
 - Administrator Roles
 - Business Roles
 - User roles permissions
 - User management approaches:
 - Organisation Centric
 - Trial Centric
- Conclusions / Recommendations

COMPLETE TRAINING

CTIS overview and practical case for Clinical Trial Application submission

- Submission of an initial Clinical Trial Application in CTIS
 - Content and structure of an initial CTA
 - Form section
 - Part I
 - Part II
- · Creation, submission and withdrawal of a CTA
- Search, view and download information on clinical trials and clinical trial applications
- Update of an initial CTA
 - Substantial modifications
 - Addition of a new Member State Concerned
 - Requests for Information (RFI)
 - Non-substantial modifications
 - Notifications required during the life-cycle of a clinical trial
 - Trial and recruitment periods
 - Other notifications
- Submission of trial results and layperson summary
- Transparency publication of clinical trial information
 - General principles
 - o Deferral rules: definition and management in CTIS
- · Work-planning and management tools
 - Notices and alerts
 - o RFIs
 - Timetable

