

# **GCP | ENHEDERNE**

God klinisk forskning ✓✓✓

**The Danish GCP Units guide  
to transition of trials from EudraCT to CTIS**

## Content

1	What is a Transitional trial .....	3
2	Create and submit a Transitional Trial.....	4
2.1	How to create a Transitional trial.....	4
2.2	Content of new cover letter .....	4
2.3	Documents required under the CTR.....	5
2.4	New requirement for archiving data in 25 years.....	6
2.5	How to populate a Transitional trial.....	6
2.6	How to submit a Transitional trial .....	6
2.7	Transition of multinational trials .....	7
3	Respond to Request for Information (RFI) for a Transitional trial.....	7
3.1	How to respond to RFI.....	7
4	Timelines.....	8
5	How to submit notifications for a Transitional trial.....	8
6	Transparency requirements under CTR.....	8
7	Changes log.....	9

## 1 What is a Transitional trial

Trials authorised under the Clinical Trials Directive (CTD) 2001/20/EC must be transitioned to CTIS as transitional trials. All clinical trials authorised under CTD must be transitioned not later than 30 January 2025. When the trial is transitioned to the CTIS database, it is authorised under the EU Clinical Trials Regulation (CTR) No 536/2014 and all the Regulation requirements will apply.

**Further readings on transitional trials:**

**EMA training module 23:**  
[EMA Quick Guides for sponsors – please refer to module 23](#)

[European Commission Guidance for the transition of clinical trials from the Clinical Trials Directive to the Clinical Trials Regulation](#)

**Info box:**

If a clinical trial does not fully comply with the CTR, the sponsor shall request a substantial amendment under the CTD before switching to the CTR, specifying its intention to align the trial with the CTR. When the substantial amendment is accepted, sponsor can transition the trial to CTIS and thereafter follow the CTR.

Trials for which a request for a substantial amendment is under assessment cannot be transitioned until the procedure is completed

**EMA recommends transition of ongoing trials not later than September 2024:**

**Figure 5.1.2.** Schematic representation of the Clinical Trial Regulation transition period from 31 January 2023.



**Info box:**

Please notify the local Ethics Committee (e.g. Danish VEK) if the trial is transferred to CTIS before 31 January 2025.

## 2 Create and submit a Transitional Trial

### 2.1 How to create a Transitional trial

Go to the CTIS 'Clinical trials' tab and create a trial by selecting the 'New trial' button at the bottom-right corner.

A pop-up window opens, and users can populate all the required fields:

ID	Name	Address	City	postCode	country	phone	email
ORG-100012585	Test Organisation	Dionysiana Street 12	Athens	111 42	Greece	0200000000	info@testorganisation.com
ORG-100012564	Test Organisation	Santiago Calle 10	Madrid	28001	Spain	0200110000	info@testorganisation-spain.com

- 'Full title',
- Organisation details (search for the sponsor organisation address)
- When fields are populated, select the checkbox '**Transitional trial**'. *If users do not select the checkbox, they need to cancel the CTA (clinical trial application) and create a new one.*

Click on the '**Create**' button to create a draft Transitional trial.

**Info box:**

It must be indicated that it is a Transitional application at the moment of creating the draft Initial application in CTIS; if users do not select the checkbox, they need to cancel and create a new application.

### 2.2 Content of new cover letter

A new cover letter must be uploaded for the transitional trial (the old cover letter from the original application is not valid).

The following text must be included in the cover letter:

- Declaration, that the clinical trial is in line with the requirements for transitioning from the CTD to the CTR, and with the authorisation given under the CTD
- Declaration that all documents which need approval and are transitioned have been approved by the Member State concerned (MSC) prior to transition
- The name of the Ethics Committee who has given a positive opinion on the clinical trial under CTD
- The EudraCT number
- Change in the archiving period to 25 years
- For multinational trials only, the [cover letter template](#) must be used

### **2.3 Documents required under the CTR**

Documents that need to be uploaded in CTIS are e.g.

- Cover letter
- Protocol with track changes – if protocol is updated (e.g with archiving period)
- Compliance with GDPR
- "Synopsis of the protocol"
- Subject information sheet
- Informed consent form (ICF)
- Documents related to the Investigational Medicinal Product (IMP) e.g. SmPC, IMPD (if applicable)
- Documents related to non-investigational medicinal products (i.e. auxiliary medicinal products under the CTR, if applicable)

Some of the documents listed in Annex I of the CTR are new as they were not required under the CTD.

*It is not a requirement to submit the form "Suitability of the clinical trial sites facilities" (one per site) with the transition application. This is only required if new sites are added as substantial modification AFTER transition is complete).*

The sponsor may submit additional documentation in addition to what is required above for the transitioning application, if these documents were assessed and authorised under the CTD.

Users can upload a blank document to continue the creation of the transitional trial application if a document was not a requirement under the CTD, but mandatory to be submitted for the CTA in CTIS.

The document must clarify that this aspect was assessed by National Competent Authority (NCA) and/or Research Ethics Committee (REC) and therefore is covered by the conclusion of the assessment under the CTD.

Documents that has been replaced by a blank document will need to be submitted as part of the first substantial modification application after the transition authorisation.

## 2.4 New requirement for archiving data in 25 years

In CTR the requirement for archiving is 25 years for all documents in TMF and CRF. Source data from the medical records should be archived according to national law.

The 25-year archiving period must be updated in the protocol when submitting the application for transition.

The change in archiving period is a Non Substantial Modification (NSM) but the change in the protocol must be mentioned in the transition cover letter.

## 2.5 How to populate a Transitional trial

Once the draft Transitional trial application has been created, users can start populating CTIS with the required fields of the sections: **Form**, **MSCs**, **Part I** and **Part II**.

In the Form section, in the Transition trial sub-section, users can **include the relevant EudraCT number** for that respective Transitional trial. This sub-section is only available when a user has marked the 'Transitional trial' checkbox when creating the CTA. Click on the '+ Add EudraCT Trial' button, then search the respective trial via the search bar.

The screenshot displays the 'Form' section of a Transitional trial application. The 'Form details' section is expanded to show 'Initial Application details' and 'Transition Trial'. The 'Transition Trial' section is highlighted with an orange box, showing a dropdown menu with 'Transition Trial' selected and a text field for 'EUDRA CT number'. Below this, a modal window titled 'EudraCT Trial Search' is shown, featuring a search bar for 'EUDRA CT number', 'CLEAR' and 'SEARCH' buttons, and a 'Search result' section. The 'Add EudraCT Trial' button is also highlighted in orange.

Thereafter users must populate the rest of the CTA sections.

In the section 'MSCs', only those sites (and MSC) where the clinical trial is still ongoing need to be mentioned.

## 2.6 How to submit a Transitional trial

Check that all information has been populated correctly by clicking on the 'Check' button. The system will highlight the sections where mandatory data or documents are missing.

The draft Transitional trial cannot be submitted if any mandatory fields or documents are not filled in or uploaded.

## 2.7 Transition of multinational trials

It is required to harmonise all documents common to all Member States concerned (i.e. documents from Part I). Sponsors are only allowed to upload **one version of the protocol for each trial**.

**Info box:**

For multinational trials only, the [cover letter template](#) must be used

*Adopted by the Clinical Trial Coordination Group (CTCG) under the Heads of Medicines Agency (HMA)*

Sponsors will be required to have a harmonised or consolidated protocol approved under Directive 2001/20/EC, prior to transitioning. Differences in a consolidated version of the protocol are limited to the Member State-specific requirements.

*A declaration of the harmonized/consolidated protocol must be included in the cover letter for multinational trials (please see cover letter template in info box).*

**For more information for transition of multinational trials:**

[CTCG Best Practice Guide Vs 2.0 Sep 2023 for sponsors of multinational clinical trials with different Part I documents approved in different MSs](#)

## 3 Respond to Request for Information (RFI) for a Transitional trial

### 3.1 How to respond to RFI

Sponsors can access the 'Evaluation' section about information that needs to be clarified, called RFI. For more information on how to respond to RFIs, sponsor users can refer to the materials of Module 11.

**EMA training module 11 for responding to RFI:**

[EMA Quick Guides for sponsors – please refer to module 11](#)

There will be no re-assessment of the submitted dossier during the transitioning application. The documentation of which the ongoing trial was authorised is already available within the Member State(s).

## 4 Timelines

Member States Concerned (MSCs) may choose to authorise the trials within or in **less than 60 days**. In case of Requests for Information (RFIs) the trial will be authorised within a **maximum period of 106 days**.

**Info box:**

Please be aware to check “Notices & alerts” in CTIS regularly, if a RFI is received.

When the trial is authorised under the CTR in CTIS, all requirements of the Regulation will apply (e.g. obligations of notification via CTIS, safety reporting rules, archiving requirements, transparency requirements, procedures for submission of substantial modifications and summary of results).

## 5 How to submit notifications for a Transitional trial

Notifications can be submitted when the trial has been authorised. The sponsor may submit a 'Start trial' notification with a trial start date in the past and prior to the authorisation date documented in CTIS, as the application was already authorised under the CTD.

**EMA training module 5 for submission of notifications:**

[EMA Quick Guides for sponsors – please refer to module 5](#)

In connection with transition to CTIS, the system will automatically send a mail saying that the trial is “on hold”, hence right after authorisation in CTIS the trial start dates (start of recruitment and start of inclusion) needs to be mentioned for all MSCs where the trial is ongoing.

## 6 Transparency requirements under CTR

Documents submitted by the sponsor in the transition application fall under the transparency requirements of the CTR and have to be made publicly available.

When transitioning a minimum dossier (see section 2.3), the sponsor must prepare redacted versions for publication of:

- the protocol,
- subject information sheet(s)
- informed consent form(s).

Name the document “(for publication)” in the file name.

The other documents in the dossier can be submitted as the non-redacted documents already approved by the Member State(s). Instead of redacted versions for other parts of the application dossier, a document referring to the previous approvals under the Directive from health authority and/or ethics committee, can be uploaded in CTIS.



Users can click on the 'Add' button (+) that appears in the document section after uploading the first version of the document intended for publication (only available in CTIS sections where the documents to be uploaded are published).



At the time of the next substantial modification application, redacted versions must also be submitted for publication for the rest of the dossier.

Any new notification after the trial has been transitioned, will fully fall under the transparency rules of the CTR (including deferrals) for making certain documents publicly available.

## 7 Changes log

Version 1.0

Version 1.1: Updated with recommendation of transition not later than Sep. 2024 and that sponsor must notify local EC when trial is transitioned to CTIS.

Version 1.2: Added requirement for updating the protocol with the archiving period 25 years when submitting the transition.

Version 1.3: Updated according to European Commission guidance for transition of clinical trials