

3 May 2024 EMA/174397/2024 European Medicines Agency

# CTIS newsflash - 3 May 2024

#### Introduction

This regular CTIS newsflash provides key updates on CTIS and links to useful reference materials.

The next issue will be circulated on 17 May 2024.

Previous issues of the CTIS Newsflash are available on the EMA website.

## Reminder: Transition ongoing trials from CTD to the Clinical Trials Regulation

Sponsors are advised to transition any trials that are expected to continue after 30 January 2025 from the Clinical Trials Directive (CTD) to the Clinical Trials Regulation (CTR).

Sponsors should take into account the time necessary for completion of the Member State(s) evaluation procedure, which can take up to 3 months. Members States have agreed on an expedited procedure for transitioning trials to the CTR which will be applied whenever possible.

Further resources to support sponsors transitioning trials are available on the CTIS website.

## Reminder: Launch of revised CTIS transparency rules on 18 June 2024

The <u>revised CTIS transparency rules</u> will become applicable on 18 June 2024, with the launch of an updated version of the <u>CTIS public portal</u>. Sponsors are advised to adapt their business processes accordingly, and can refer to the <u>quick quide for users</u> for an overview of the changes.

For all clinical trial applications submitted on or after 18 June 2024:

- it will no longer be possible to defer the publication of data and documents;
- data and documents will be published according to the established timelines for the trial category, population age and trial phase;
- publication of documents will be focused on key documents of interest.

Data on all clinical trial applications submitted before 18 June 2024 will be made publicly available in line with the principles and timelines defined in the revised transparency rules. Please note that existing CTIS documents of these trials will not be published. Documents included in subsequent



applications of these trials submitted after 18 June 2024 will be published in line with the revised rules<sup>1</sup>: more details are available in the <u>quick guide for users</u>.

In the interim period until 18 June 2024, sponsors may already follow the principles of the revised CTIS transparency rules, as defined in section 4 of the <u>ACT EU Q&A</u>.

More information and resources are available on the <u>ACT EU website</u> news item and on the <u>CTIS</u> website.

## **Recently published**

- The latest <u>KPI report with data from March 2024</u> on the implementation of the Clinical Trials Regulation has been published on the ACT EU website.
- Revised <u>"Recommendations related to contraception and pregnancy testing in clinical trials"</u> have been published on the CTCG website.



## Save the date: Upcoming events and trainings

The next <u>CTIS walk-in clinic</u> will take place on 15 May 2024 at 16:00 – 17:00 CEST. Participants can submit their questions via <u>Slido</u> until 8 May 2024, with the code #clinic245.

Sponsors can already register to the upcoming CTIS user training on <u>10-13 June</u> 2024, 09:00-13:30 CEST. For more information on previous training sessions, including supporting materials, see: <u>Clinical Trials Information System: training and support | European Medicines Agency (europa.eu).</u>

# New CTIS feature: Updating sponsor information via a Non-Substantial Modification

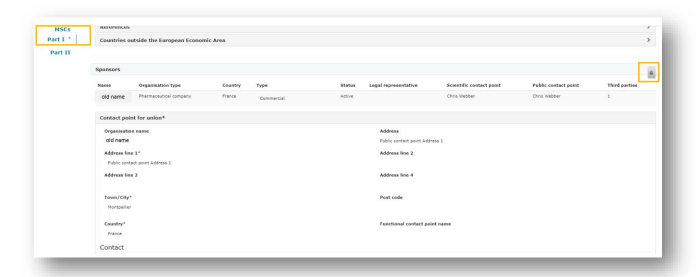
A new feature now allows sponsor users to update the details of the sponsor of their trial.

Users can submit a non-Substantial Modification (non-SM) to update the sponsor details recorded on an application form, such as the sponsor's name (without change of the legal entity), contact details, or address; see column 81.9NSM in <a href="Annex IV of the Q&A document on CTR">Annex IV of the Q&A document on CTR</a>.

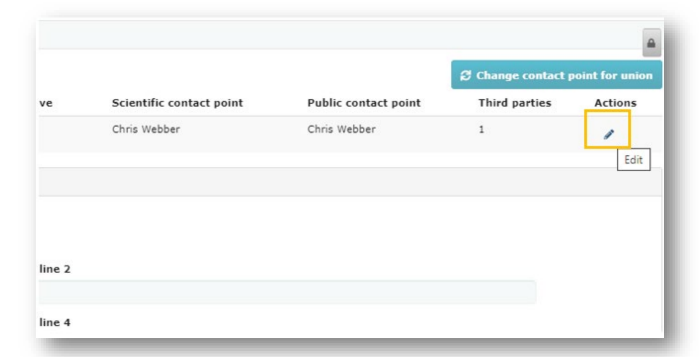
Before recording the changes in CTIS, sponsor users need to update the OMS data. Users should submit a change request to update the organisation details <u>in OMS</u> (see document E - OMS Change Requests in <u>OMS document repository</u>), wait until their change request is validated and approved by the OMS team, and then apply the changes in CTIS by following the instructions below.

Sponsor details are captured in Part I; therefore, sponsor users need to submit a non-SM that affects Part I (Part I only or Part I & II). After creating the non-SM draft (Module 10), users can click on the Part I page and use the padlock of the 'Sponsors' section to edit the previously recorded details.

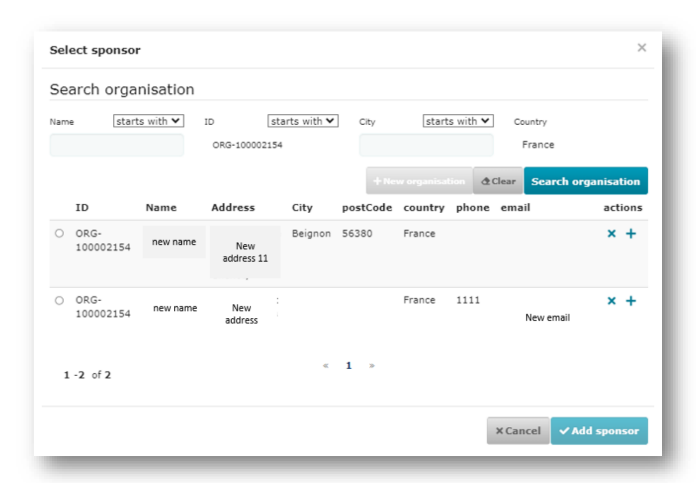
<sup>&</sup>lt;sup>1</sup> This applies to documents of all types of applications, with the exception of part I documents of Non Substantial Modifications and Additional Member State applications.



After selecting the padlock, users should see an edit button (right side).



After selecting the edit button, a pop-up window appears which allows users to search for organisations. The fields 'organisation ID' and 'country' are already populated and cannot be edited; changes to these fields imply a change of legal entity. Users can use the non-mandatory fields 'Name' or 'City' to search for the organisation. If the OMS change request has been approved, the new entry with the updated sponsor details will appear in the search results.



Users need to select the correct result (using the radio button on the left), to activate the 'Add sponsor button'. After clicking the 'Add sponsor' button, the updated sponsor details will overwrite the obsolete ones. Users may save the draft and submit it.

After the change of the sponsor details on CTIS, any new application, Annual Assessment Report (ASR), Ad hoc assessment or Inspection that is created will reflect the updated sponsor details. Previous applications (or ASRs or Ad-hoc assessments) submitted before the change of sponsor information in CTIS will keep the previous sponsor details and will not be impacted by the change.

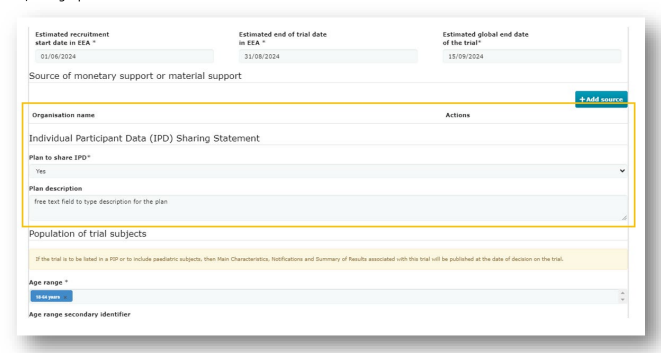
Training materials are being updated to provide more information on this new feature. Users will be notified of the publication of revised training materials via the regular communication channels, including this newsflash.

## **New CTIS feature: Field on Individual Participant Data**

As required in the <u>WHO Trial Registration Data Set</u>, the field 'Individual Participant Data (IPD) Sharing Statement' collects information on how this data will be made available to other researchers. Sponsor users in CTIS can now record in a structured way how IPD will be shared in the future.

Two new fields have been added in the Part I section of the application form in CTIS, above the fields for the population of trial subjects. The first new field, 'Plan to share IPD', is mandatory. Users need to select a response from a drop-down list of pre-defined values (Yes/No/Undefined). We recommend that sponsor users select "Yes" or "No", to meet the <u>requirements of the International Committee of Medical Journals Editors (ICJME)</u>.

The second field, 'Plan description', is optional. It allows users to describe the plan in detail, in free text, using up to 1000 characters.



#### **System improvements**

In addition to the two new features described above, the CTIS release on 30 April 2024 introduced several other improvements:

- In Non-Substantial Modifications for Part I only, users can now update the number of subjects
  without this action leading to a duplication of the Member State Concerned (MSC). This also applies
  when creating a subsequent Substantial Modification for Part I only.
- In Non-Substantial Modifications (non-SMs) for Part I & II or Part II only, if the user chooses not to submit Part II for one or more of the MSC, then a draft non-SM Part II is created (but not submitted) for these MSC.
- Member State users in the authority workspace can now download the temporary halt notification from the 'Notifications' tab.
- When there are multiple documents in content labelling of Investigational Medicinal Products, users
  can perform changes in the context of a Request for Information (any phase) or submission of any
  type of application, without impacting the links to products.
- Users with the combined roles Q-IMPD (Investigational Medicinal Product Dossier Quality)
   Preparer and CT Admin can now see all the corresponding Requests for Information and their considerations, quality and non-quality.

More information on the latest system improvements is available in the published <u>release notes</u> as well as in the Lists of known issues and proposed workarounds for <u>sponsors</u> and for <u>Member State users</u>.

EMA continues to closely monitor user feedback, working in collaboration with our stakeholders to deliver further system improvements and enhance the user experience.

## **Current operational experience with CTIS**

This section on weekly CTIS metrics provides key data and trends.

The data presented below refer to the period from 23 to 29 April 2024.

#### **CTA Submissions**



#### CTAs with a Decision



The data presented below refer to the period from 16 to 22 April 2024.

# **CTA Submissions**



#### CTAs with a Decision



## **Requesting access to the CTIS Training Environment**

Sponsors can express their interest in gaining access to the CTIS Training Environment by filling in the ongoing <u>survey</u>. The training environment is a simulation of CTIS used in production and allows users to get familiar with system functionalities.

#### **More information**

Are you a sponsor user starting out with CTIS? Please consult the 'Sponsor quick guide: Getting started with CTIS' or refer to the CTIS training material, including the new version of the 'CTIS Handbook for clinical trial sponsors'. The handbook provides useful information on how sponsors can navigate CTIS to create and submit clinical trial information to the member states of the European Union as required by the Clinical Trial Regulation (EU) No 536/2014.

Resources to support sponsors transitioning trials from the Clinical Trials Directive (CTD) to the CTR are available on the <u>CTIS website</u>.