

# Annual Progress Report to Research Ethics Committee

## Clinical Trial of an Investigational Medicinal Product (CTIMP)

**To be completed and submitted by the Chief Investigator or sponsor. Please email this report to the REC. For questions with Yes/No options please indicate answer in bold type.**

1. **Details of the Chief Investigator**

|  |  |
| --- | --- |
| Name: |  |
| Address: |  |
| Telephone: |  |
| E-mail: |  |

### Details of Study

|  |  |
| --- | --- |
| Full title of study: |  |
| Name of REC: |  |
| IRAS ID: |  |
| REC reference number: |  |
| Date of favourable ethical opinion: |  |
| Sponsor: |  |
| EudraCT Number: |  |

1. **Commencement and termination dates**

|  |  |
| --- | --- |
| 3.1 Has the study started in the UK? | Yes / No |
| 3.2 If yes, what was the actual start date in the UK? |  |
| If no, what are the reasons for the study not commencing in the UK?What is the expected start date?Please note, if the study will not start within 24 months of the REC Favourable Opinion date the REC may review its’ opinion.  |  |
| Has the study finished?If yes, complete and submit “Declaration of end of trial” form at Annex 3 to ENTR/CT1, available on the [Gov.UK Website](https://www.gov.uk/guidance/clinical-trials-for-medicines-manage-your-authorisation-report-safety-issues#end-of-trial) | Yes / No |
| If no, what is the expected completion date?If you expect the study to overrun the planned completion date, what are the reasons for this?  |  |
| If you do not expect the study to be completed, give reason(s) |  |

1. **Registration**

|  |  |
| --- | --- |
| Has the trial been registered? | Yes / No |
| If yes, please provide the name of the registry and registration number? | Registration number:  |
| If no: | 1. What is the reason for non-registration?
2. What are your intentions for registration?
 |

1. **Site Information**

|  |  |
| --- | --- |
| Number of UK research sites proposed in original application:Number of UK research sites recruited to date: |  |
| Do you plan to increase the total number of UK sites proposed for the study?Please note, the addition of any new sites not listed in the original application should be submitted as an amendment. | Yes / No |

1. **Recruitment of Participants**

|  |  |
| --- | --- |
| \*Number of participants recruited: | Proposed in original application:Actual number recruited to date: |
| \*Number of participants completing trial: | Actual number completed to date: |

##### \*Number of withdrawals from trial date due to:

|  |  |
| --- | --- |
| (a) withdrawal of consent  |  |
| (b) loss to follow-up |  |
| (c) death (where not the primary outcome) |  |
| Total study withdrawals: |  |

##### \*Number of treatment failures to date (prior to reaching primary outcome) due to:

|  |  |
| --- | --- |
| (a) adverse events |  |
| (b) lack of efficacy |  |
| Total treatment failures: |  |

\* In the case of international trials, please provide separate figures for UK and non-UK participants.

|  |  |
| --- | --- |
| Have there been any serious difficulties in recruiting participants ? | Yes / No |
| If yes, give details: |  |
| Do you plan to increase the planned recruitment of participants into the study?Please note, any increase in planned recruitment or changes to the recruitment methodology should be notified to the REC as a substantial amendment for ethical review. | Yes / No |

1. **Safety Reports**

|  |  |
| --- | --- |
| Have there been any Suspected Unexpected Serious Adverse Reactions (SUSARs) in this trial in the UK and have they been notified to the Committee? | Yes / No |
| Has the Development Safety Update Report (DSUR) been submitted?Sponsors are required to submit a DSUR within one year of the Development International Birth Date (DIBD – the date of first authorisation of a clinical trial in any country worldwide) and provide annual DSUR submissions until all open clinical studies have ended (the final clinical study is completed and its study report has been submitted). | Yes / No / Not yet due |

1. **Amendments**

|  |  |
| --- | --- |
| Have any substantial amendments been made to the trial during the year? | Yes / No |
| If yes, please give the date and amendment number for each substantial amendment made. |  |

1. **Serious breaches of the protocol or Good Clinical Practice**

|  |  |
| --- | --- |
| Have any serious breaches of the protocol or GCP occurred in relation to this trial during the year?Under the Clinical Trials Regulations, all serious breaches must be notified to the MHRA GCP inspectors within 7 days of the matter coming to the sponsor’s attention. | Yes / No |
| If yes, please give the date of each notification to the MHRA.Please provide the REC with a copy of each notification for information (unless previously notified). |  |

1. **Other issues**

|  |  |
| --- | --- |
| Are there any other developments in the trial that you wish to report to the Committee? | Yes / No |

1. **Declaration**

|  |  |
| --- | --- |
| \*Signature/Electronic authorisation of Chief Investigator or Sponsor representative: \*Please print name below and insert electronic signature, if possible |  |
| Print name: |  |
| Date of submission: |  |