

**Substantial Amendment Notification Form (Cf. Section 3.7.b of the *Detailed guidance on the request to the competent authorities for authorisation of a clinical trial on a medicinal product for human use, the notification of substantial amendments and the declaration of the end of the trial*<sup>1</sup>)**

**NOTIFICATION OF A SUBSTANTIAL AMENDMENT TO A CLINICAL TRIAL ON A MEDICINAL PRODUCT FOR HUMAN USE TO THE COMPETENT AUTHORITIES AND FOR OPINION OF THE ETHICS COMMITTEES IN THE EUROPEAN UNION**

*For official use:*

Date of receiving the request :	Grounds for non acceptance/ negative opinion : <input type="checkbox"/> Date :
Date of start of procedure:	Authorisation/ positive opinion : <input type="checkbox"/> Date :
Competent authority registration number of the trial: Ethics committee registration number of the trial :	Withdrawal of amendment application <input type="checkbox"/> Date :

*To be filled in by the applicant:*

This form is to be used both for a request to the Competent Authority for authorisation of a **substantial** amendment and to an Ethics Committee for its opinion on a **substantial** amendment. Please indicate the relevant purpose in Section A.

**A TYPE OF NOTIFICATION**

**A.1 Member State in which the substantial amendment is being submitted:**

**A.2 Notification for authorisation to the competent authority:**

**A.3 Notification for an opinion to the ethics committee:**

**B TRIAL IDENTIFICATION** (*When the amendment concerns more than one trial, repeat this form as necessary.*)

**B.1 Does the substantial amendment concern several trials involving the same IMP?**<sup>2</sup> yes  no

B.1.1 If yes repeat this section as necessary.

**B.2 Eudract number:**

**B.3 Full title of the trial :**

**B.4 Sponsor's protocol code number, version, and date:**

**C IDENTIFICATION OF THE SPONSOR RESPONSIBLE FOR THE REQUEST**

**C.1 Sponsor**

C.1.1 Organisation:

C.1.2 Name of person to contact:

C.1.3 Address :

C.1.4 Telephone number :

C.1.5 Fax number :

C.1.6 e-mail:

**C.2 Legal representative<sup>3</sup> of the sponsor in the European Union for the purpose of this trial (if different from the sponsor)**

C.2.1 Organisation:

C.2.2 Name of person to contact:

C.2.3 Address :

C.2.4 Telephone number :

C.2.5 Fax number :

C.2.6 e-mail:

**D APPLICANT IDENTIFICATION (please tick the appropriate box)**

<sup>1</sup> OJ, C82, 30.3.2010, p. 1; hereinafter referred to as 'detailed guidance CT-1'.

<sup>2</sup> Cf. Section 3.7. of the detailed guidance CT-1.

<sup>3</sup> As stated in Article 19 of Directive 2001/20/EC.

<b>D.1 Request for the competent authority</b>	
D.1.1 Sponsor	<input type="checkbox"/>
D.1.2 Legal representative of the sponsor	<input type="checkbox"/>
D.1.3 Person or organisation authorised by the sponsor to make the application.	<input type="checkbox"/>
D.1.4 Complete below:	
D.1.4.1 Organisation :	
D.1.4.2 Name of person to contact :	
D.1.4.3 Address :	
D.1.4.4 Telephone number :	
D.1.4.5 Fax number :	
D.1.4.6 E-mail	

<b>D.2 Request for the Ethics Committee</b>	
D.2.1 Sponsor	<input type="checkbox"/>
D.2.2 Legal representative of the sponsor	<input type="checkbox"/>
D.2.3 Person or organisation authorised by the sponsor to make the application.	<input type="checkbox"/>
D.2.4 Investigator in charge of the application if applicable <sup>4</sup> :	
• Co-ordinating investigator (for multicentre trial)	<input type="checkbox"/>
• Principal investigator (for single centre trial):	<input type="checkbox"/>
D.2.5 Complete below	
D.2.5.1 Organisation :	
D.2.5.2 Name :	
D.2.5.3 Address :	
D.2.5.4 Telephone number :	
D.2.5.5 Fax number :	
D.2.6 E-mail :	

## E SUBSTANTIAL AMENDMENT IDENTIFICATION

<b>E.1 Sponsor's substantial amendment code number, version, date for the clinical trial concerned:</b> ( )
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<b>E.2 Type of substantial amendment</b>	
E.2.1 Amendment to information in the CT application form	yes <input type="checkbox"/> no <input type="checkbox"/>
E.2.2 Amendment to the protocol	yes <input type="checkbox"/> no <input type="checkbox"/>
E.2.3 Amendment to other documents appended to the initial application form	yes <input type="checkbox"/> no <input type="checkbox"/>
E.2.3.1 If yes specify:	
E.2.4 Amendment to other documents or information:	yes <input type="checkbox"/> no <input type="checkbox"/>
E.2.4.1 If yes specify:	
E.2.5 This amendment concerns mainly urgent safety measures already implemented <sup>5</sup>	yes <input type="checkbox"/> no <input type="checkbox"/>
E.2.6 This amendment is to notify a temporary halt of the trial <sup>6</sup>	yes <input type="checkbox"/> no <input type="checkbox"/>
E.2.7 This amendment is to request the restart of the trial <sup>7</sup>	yes <input type="checkbox"/> no <input type="checkbox"/>

<sup>4</sup> According to national legislation.

<sup>5</sup> Cf. Section 3.9. of the detailed guidance CT-1.

<sup>6</sup> Cf. Section 3.10. of the detailed guidance CT-1.

<sup>7</sup> Cf. Section 3.10. of the detailed guidance CT-1.

<b>E.3</b>	<b>Reasons for the substantial amendment:</b>	
E.3.1	Changes in safety or integrity of trial subjects	yes <input type="checkbox"/> no <input type="checkbox"/>
E.3.2	Changes in interpretation of scientific documents/value of the trial	yes <input type="checkbox"/> no <input type="checkbox"/>
E.3.3	Changes in quality of IMP(s)	yes <input type="checkbox"/> no <input type="checkbox"/>
E.3.4	Changes in conduct or management of the trial	yes <input type="checkbox"/> no <input type="checkbox"/>
E.3.5	Change or addition of principal investigator(s), co-ordinating investigator	yes <input type="checkbox"/> no <input type="checkbox"/>
E.3.6	Change/addition of site(s)	yes <input type="checkbox"/> no <input type="checkbox"/>
E.3.7	Other change	yes <input type="checkbox"/> no <input type="checkbox"/>
E.3.7.1	If yes, specify:	
E.3.8	Other case	yes <input type="checkbox"/> no <input type="checkbox"/>
E.3.8.1	If yes, specify	

<b>E.4</b>	<b>Information on temporary halt of trial<sup>8</sup></b>
E.4.1	Date of temporary halt (YYYY/MM/DD)
E.4.2	Recruitment has been stopped <span style="float: right;">yes <input type="checkbox"/> no <input type="checkbox"/></span>
E.4.3	Treatment has been stopped <span style="float: right;">yes <input type="checkbox"/> no <input type="checkbox"/></span>
E.4.4	Number of patients still receiving treatment at time of the temporary halt in the MS concerned by the amendment ( )
E.4.5	Briefly describe (free text): <ul style="list-style-type: none"> <li>• Justification for a temporary halt of the trial</li> <li>• The proposed management of patients receiving treatment at time of the halt (<i>free text</i>).</li> </ul> The consequences of the temporary halt for the evaluation of the results and for overall risk benefit assessment of the investigational medicinal product ( <i>free text</i> ).

**F DESCRIPTION OF EACH SUBSTANTIAL AMENDMENT<sup>9</sup> (*free text*):**

Previous and new wording in track change modus	New wording	Comments/explanation/reasons for substantial amendment

**G CHANGE OF CLINICAL TRIAL SITE(S)/INVESTIGATOR(S) IN THE MEMBER STATE CONCERNED BY THIS AMENDMENT**

<b>G.1</b>	<b>Type of change</b>
G.1.1	<b>Addition of a new site</b>
G.1.1.1	<b>Principal investigator</b> (provide details below)
G.1.1.1.1	Given name
G.1.1.1.2	Middle name (if applicable)
G.1.1.1.3	Family name
G.1.1.1.4	Qualifications (MD.....)
G.1.1.1.5	Professional address
G.1.2	<b>Removal of an existing site</b>
G.1.2.1	<b>Principal investigator</b> (provide details below)
G.1.2.1.1	Given name
G.1.2.1.2	Middle name (if applicable)
G.1.2.1.3	Family name
G.1.2.1.4	Qualifications (MD.....)
G.1.2.1.5	Professional address
G.1.3	<b>Change of co-ordinating investigator</b> (provide details below of the new coordinating investigator)
G.1.3.1	Given name
G.1.3.2	Middle name
G.1.3.3	Family name

<sup>8</sup> Cf. Section 3.10. of the detailed guidance CT-1.

<sup>9</sup> Cf. Section 3.7.c. of the detailed guidance CT-1. The sponsor may submit this documentation on a separate sheet.

- G.1.3.4 Qualification (MD.....)
- G.1.3.5 Professional address
- G.1.3.6 Indicate the name of the previous co-ordinating investigator:
- G.1.4 **Change of principal investigator at an existing site** (provide details below of the new principal investigator)
- G.1.4.1 Given name
- G.1.4.2 Middle name
- G.1.4.3 Family name
- G.1.4.4 Qualifications (MD.....)
- G.1.4.5 Professional address
- G.1.4.6 Indicate the name of the previous principal investigator:

## H CHANGE OF INSTRUCTIONS TO CA FOR FEEDBACK TO SPONSOR

### H.1 Change of e-mail contact for feedback on application\*

H.2 Change to request to receive an .xml copy of CTA data  yes  no

H.2.1 Do you want a .xml file copy of the CTA form data saved on EudraCT?  yes  no

H.2.1.1 If yes provide the e-mail address(es) to which it should be sent (up to 5 addresses):

H.2.2 Do you want to receive this via password protected link(s)<sup>10</sup>?  yes  no

If you answer no to question H.2.2 the .xml file will be transmitted by less secure e-mail link(s)

H.2.3 Do you want to stop messages to an email for which they were previously requested?  yes  no

H.2.3.1 If yes provide the e-mail address(es) to which feedback should no longer be sent:

(\*This will only come into effect from the time at which the request is processed in EudraCT).

## I LIST OF THE DOCUMENTS APPENDED TO THE NOTIFICATION FORM (cf. Section 3.7 of detailed guidance CT-1)

*Please submit only relevant documents and/or when applicable make clear references to the ones already submitted. Make clear references to any changes of separate pages and submit old and new texts. Tick the appropriate box(es).*

- I.1 Cover letter
- I.2 Extract from the amended document in accordance with Section 3.7.c. of detailed guidance CT-1 (if not contained in Part F of this form)
- I.3 Entire new version of the document<sup>11</sup>
- I.4 Supporting information
- I.5 Revised .xml file and copy of initial application form with amended data highlighted
- I.6 Comments on any novel aspect of the amendment if any :

## J SIGNATURE OF THE APPLICANT IN THE MEMBER STATE

- J.1 I hereby confirm that/ confirm on behalf of the sponsor that (delete which is not applicable)
- The above information given on this request is correct;
  - The trial will be conducted according to the protocol, national regulation and the principles of good clinical practice; and
  - It is reasonable for the proposed amendment to be undertaken.

J.2 APPLICANT OF THE REQUEST FOR THE COMPETENT AUTHORITY (as stated in section D.1):

<sup>10</sup> This requires a EudraLink account. (See <https://eudract.ema.europa.eu/> for details)

<sup>11</sup> Cf. Section 3.7.c. of the detailed guidance CT-1.

J.2.1 Signature <sup>12</sup>:

J.2.2 Print name :

J.2.3 Date :

**J.3 APPLICANT OF THE REQUEST FOR THE ETHICS COMMITTEE** (as stated in section D.2):

J.3.1 Signature <sup>13</sup>:

J.3.2 Print name:

J.3.3 Date :

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<sup>12</sup> On an application to the Competent Authority only, the applicant to the Competent Authority needs to sign.

<sup>13</sup> On an application to the Ethics Committee only, the applicant to the Ethics Committee needs to sign.

