"Please do take into account that this is a translation of the original French version validated in the Quality Management System (QMS) of Cliniques universitaires Saint-Luc through the software Ennov GED. Therefore in case of doubt, differences, inconsistency or discrepancy in this English version, the French version shall prevail.”

1 PROCEDURE’S OBJECT

The objective of this procedure is to determine the end of approval of a clinical experiment and to specify the modalities to inform the Ethics Committee, the institution, the investigator and the competent authorities.

2 PROCEDURE’S SCOPE

8. Comité d'Ethique Hospitalo-Facultaire (CEHF)

3 RESPONSIBILITIES AND AUTHORITIES

- The **CEHF administrative secretary** is in charge of encoding the date of end of experiment in the CEHF database. The information is provided by the sponsor of the experiment or the investigator at time of initial submission. If no amendment has been submitted requesting for prolongation of the experiment, the date of end of experiment will be the one mentioned in the initial submission package.

- The **sponsor** of an experiment is responsible to inform the LEC of the end of study, suspension, early termination or temporary halt.

- The **investigator** of an experiment is responsible to transmit the documents provided by the sponsor to the CEHF in case of end of study, suspension, early termination or temporary halt. He/she is responsible the record these events in the database Claire. He/is also responsible to ensure the participants’ safety.

- **Validity of the positive opinion**: the positive opinion of the LEC is applicable for the declared duration of the experiment, as defined at time of submission of the research project to the EC. The approval is therefore applicable from approval date to the presumed termination of the experiment as mentioned in Document 1. However, the renewal of the validity of the approval depends on the continuing review of the CEHF and of the analysis of the annual report, provided by the investigator each year after the date of the initial approval.
4 PROCEDURE’S REVISION

5 PROCEDURE’S DESCRIPTION

5.1 End of experiment

5.1.1 Information of the EC for each clinical research:

- The sponsor informs the EC (the competent EC for monocenter study, and the leading EC for multicenter study) by mail/email within 90 days after the end of the experiment.
  - The 90 days timing is shortened to 15 days when the end of experiment must be anticipated. Notifications will detail clearly the reasons of anticipated stop.
  - A report (http://www.ich.org/fileadmin/Public_Web_Site/ICH_Products/Guidelines/Efficacy/E3/E3_Guideline.pdf) of end of experiment must be sent within the year following the notification of end of experiment for non-paediatric clinical trials. For paediatric clinical trials, the timing is shortened to 6 months (2009/C168/02).

- The investigator informs by email the CEHF of the end of experiment at the CUSL as notified by the sponsor.
  - He/she records the center closure date in Claire
  - He creates the submission « déclaration de fin d’étude » or « déclaration de clôture de site ».

Documents to provide to the LEC for information of the (+ notification of the local EC) end of trial:

<table>
<thead>
<tr>
<th>Document</th>
<th>Paper copy (1)</th>
<th>Electronic copy</th>
<th>Provided by sponsor</th>
<th>Provided by the CUSL investigator</th>
<th>Signed by investigator</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acknowledgement of receipt</td>
<td>X</td>
<td></td>
<td>X if not provided by the sponsor</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Notification of end of trial form: document provided by the sponsor or CEHF-FORM-028</td>
<td>X</td>
<td>X</td>
<td>X if not provided by the sponsor</td>
<td>X</td>
<td></td>
</tr>
</tbody>
</table>

FOR DRUG EXPERIMENT : DOCUMENTS TO PROVIDE ADDITIONNALLY

- CT End of trial form
  X                              X

- Summary of experiment report (according to Eudralex Volume 10 : Detailed guidance on the application format and documentation to be submitted in an application for an Ethics Committee opinion)
  - X                              X

5.1.2 Information of the competent authorities for clinical trials with medicinal product only:

N°: CEHF-SOP-019 / REV003
Page 2 sur 6
• The sponsor informs the competent authorities of each country concerned by the trial by mail within 90 days after the end of the experiment. 
CT End of trial form [http://ec.europa.eu/health/files/eudralex/vol-10/declaration_end_trial_form.doc]

- The 90 days timing is shortened to 15 days when the end of experiment must be anticipated. Notifications will detail clearly the reasons of anticipated stop. 
  Document: « Declaration of the End of Trial Form » : Eudralex- Volume 10 Clinical Trials guidelines

- An end of experiment report (CT End of trial form [http://ec.europa.eu/health/files/eudralex/vol-10/declaration_end_trial_form.doc]) must be sent within the year following the notification of end of experiment for non-paediatric clinical trials. For paediatric clinical trials, the timing is shortened to 6 months (2009/C168/02).

- FORM for Belgium : to be sent by registered mail :
  Agence Fédérale des Médicaments et des Produits de Santé
  Responsable de la Division Recherche et Développement
  Bâtiment Eurostation, 8ème étage
  Place Victor Horta 40, boîte 40
  B-1060 Bruxelles

5.2 Suspension or interdiction of experiment (Law 7 May 2004)
See 108-AAHRPP-SOP-040 SOP for the declaration of non-compliance, deviation, violation, unexpected events occurring during an experiment (GCP 4.12.3)

5.3 Early termination of experiment decided by the sponsor

5.3.1 Information of the EC in case of early termination of any experiment

The sponsor:

- Informs the LEC by regular mail or e-mail within 15 days following the termination.
- Details clearly the reasons of early termination
- Communicates the specific measures taken to guarantee the safety of participants, if applicable.
- Uses the standard European document (« Declaration of the End of Trial Form »): Eudralex- Volume 10 Clinical Trials guidelines) for clinical trials with IMP.
- Uses his own template for any other study type

5.3.2 Information of the competent authorities in case of early termination of a clinical trial with IMP:

The sponsor:

N°: CEHF-SOP-019 / REV003
Page 3 sur 6
• Informs the competent authorities of each country concerned by the trial by registered mail within 15 days after the early termination of the experiment.
• Details clearly the reasons of early termination
• Communicates the specific measures taken to guarantee the safety of participants, if applicable.
• Uses the standard European document (« Declaration of the End of Trial Form »: Eudralex- Volume 10 Clinical Trials guidelines) for clinical trials with IMP.
• Informs the investigator who will inform the institution by encoding the information in the database Claire (GCP 4.12.2)

5.4 Premature termination or suspension of a trial (GCP 4.12.1)
The investigator who terminates or suspends a trial without prior agreement of a sponsor:
• Informs directly the sponsor and provides him with a detailed written explanation of the termination or suspension
• Informs directly the Ethics Committee and provides them with a detailed written explanation of the termination or suspension
• Informs the institution in encoding the termination/suspension of the trial in the database Claire

5.5 Temporary halt, as decided by the sponsor

• The sponsor informs the EC by mail, within 15 calendar days after the temporary halt of a study for safety reasons.
• The sponsor informs directly the principal investigator who will inform the institution in encoding the halt in the database Claire (GCP 4.12.2)

5.5.1 Information of the EC in case of temporary halt of any prospective experiment for safety reasons
The sponsor:

• Informs the LEC by regular mail or e-mail within 15 days following the temporary halt.
• Details clearly the reasons of temporary halt
• Communicates the specific measures taken to guarantee the safety of participants, if applicable.

5.5.2 Information of the competent authorities in case of temporary halt of a clinical trial with IMP:
The sponsor:

• Informs the competent authorities of each country concerned by the trial by registered mail within 15 days after temporary halt of the experiment.
• Details clearly the reasons of temporary halt
Communicates the specific measures taken to guarantee the safety of participants, if applicable.

5.5.3 Any temporary halt decided by the sponsor for any non-safety reason will be included in a substantial amendment and submitted to the Ethics Committee and to the Competent Authorities according to the procedure 061-CEHF-SOP-015.

5.5 Restart of experiment

The sponsor submits a request of restart, formulated as a substantial amendment with the form Annex 2 "Notification of a substantial amendment".

The grounds related to safety justifying the restart of the experiment must be detailed.

The experiment will restart only after approval of the EC (LEC if multicentre study) and if competent authorities (FAMHP) didn’t raise any motivated objections.

If the sponsor decides to not restart the experiment, he must inform the EC and competent authorities within 15 days, as described in section early termination.

6 DEFINITIONS AND ABBREVIATIONS

No definition of end of trial exists. It must be defined in each protocol of clinical research.

CEHF : Comité d’Ethique Hospitalo-Facultaire Saint-Luc - UCL
EC : Ethics Committee
FAMHP : Federal Agency for medicines and health products
IMP: Investigational Medicinal Product
LEC : leading EC
NLEC : non leading EC

7 REFERENCE DOCUMENTS

108 - AAHRPP-SOP-040 SOP for the declaration of non compliance, deviation, violation, unexpected events occurring during an experiment
189 – CEHF-FORM-028 End of experiment notification form

8 AAHRPP ACCREDITATION STANDARDS

II.2.G
III.2.D

9 LINKS INTRANET

10 LINKS INTERNET