RISe Clinical application Post approval activity:

Completion



New: Human-Post Approva Activities

Post Approval Activities

- * Select one of the following options to submit to the Research Ethics Board based on the guidelines (Click blue question mark for guidance):
- Annual Renewal
- Amendments to Study
- Completion of Clinical Study
- Request for Acknowledgement

* Nickname

Enter a nickname for this PAA.

(If applicable to your PAA, include descriptive words such as "IB update" or "protocol deviation" or "unanticipated event" or "local SAE" in the nickname)

Completion of Clinical Study Coversheet

Commented [CREB1]: Annual Renewals

For Clinical studies click here for information on annual renewals. If this is an annual renewal of a for-profit (industry or pharmaceutical) sponsored study, a renewal fee is required. For more details about fee payment please consult the applicable REB administration or their website.

Amendments to Study

Amendments are changes to an ongoing study. If you are changing any part of the study (e.g. co-investigators, title, agency, documentation) you must submit an amendment. Click here for more information on amending clinical studies.

Completion of Clinical Study

For Clinical studies click <u>here</u> for criteria on study completion.

Request for Acknowledgement

Protocol deviations, unanticipated problems, new information, safety letters, local serious adverse events, studies on hold, off hold, closed to accrual/enrollment, or miscellaneous information (PI, Sponsor or REB requires acknowledgement). Click here for more information on Request for Acknowledgement criteria. Any other changes to an ongoing study must be submitted through an amendment.

* 1. Date of Completion						
Enter the effective date of completion.						
* 2. Confirmation of Completion of Data Collection						
After reviewing the guidelines on the right, confirm that participant data collection has been completed. Yes No Clear						
163 140 <u>Orear</u>						
3. Number of Participants, Charts or Samples						
3.1. Enter the number of research participants enrolled at the sites/institutions covered by this ethics approval.						
3.2. Enter the number of charts reviewed or samples collected						
* 4. Final Date / Notice Enter the date of the study monitor's final visit or notice, if applicable. If not applicable please select "not applicable" below.						
Not Applicable □						
* 5. Data/ Biospecimen Storage/ Destruction						
5.1. Please describe:						

Commented [u2]: Click <u>here</u> for the definition of study closure for studies involving participant recruitment and studies that do not involve direct human participation (e.g. chart reviews and data registries).

A study is considered complete where there has been either an official "close-out" visit by a Sponsor or there is no further requirement to submit data to the Sponsor. Studies being monitored by some sponsors are not complete until the centre is notified by the sponsor that the study is complete.

Studies that are grant funded may be completed when there is no active grant that requires ethics approval.

Commented [u3]: Question 3.2 should be answered only if you were not required by the REB to consent individuals for the use of their data or tissues. E.g., you received a waiver of consent for secondary use (such as chart reviews) or biological materials (such as tissue from diagnostic tests or surgeries) for part or all of your study. If you consented individuals for the use of their data, please complete 3.1.

Δ١	How long	the study	data/biospe	ecimens	will he	retained	and	where
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- B) Who will have access to the data/ biospecimens in the future and for what
- C) What plans there are for future use of the data/biospecimens (if any)



5.2. If the data/biospecimens will be destroyed, indicate the planned method for erasure/destruction of the data/biospecimens, including when they will be destroyed.



* 6. Reason for Completion

Please provide the reason for the completion of this study (i.e. did the study run its course, or if it ended early, explain why; if the study involved enrollment of participants, comment about enrollment and whether enrollment goals were achieved.) Include any other information required by the study sponsor to be submitted to the Research Ethics Board.



* 7. Submission of study results to ClinicalTrials.gov

Is this a study registered with ClinicalTrials.gov?

○ Yes ○ No Clear

Commented [u4]: Please include the following information:

•Final disposition/storage of all research-related study documents.

For studies reviewed by a UBC REB: According to UBC Policy SC6 (formerly policy 85), study data should be kept for a minimum of 5 years after publication. Clinical trials data must abide by Health Canada's regulations regarding data retention and generally must be kept for 25 years. Click here for more information concerning Health Canada requirements.

- •The procedure that will be followed in response to additional requests for access to the study data/ biospecimens (after the study has been completed and analyzed).
- •Plans for the final disposition of any electronic data or if applicable, the final disposition of any biospecimens.

Commented [u5]: Section 801 of the Food and Drug Administration Amendments Act (FDAAA 801) (PDF) requires Responsible Parties to register and submit summary results of clinical trials with ClinicalTrials.gov.

For more information about this requirement, please refer to Clinical Trials Registration on the Office of Research Services website.

If yes, please confirm that a summary of study results has been submitted to ClinicalTrials.gov. Details should include the name of the individual responsible for submitting results, as well as the date of submission.

8. Reported Results and Sponsor close-out
Add
Title
There are no items to display

Publications/Presentations that have reported results for this study



9. If this study required Health Authority Operational Approval, confirm that the Health Authority has been notified separately of the study's completion.

O Yes O No Clear

Please note: Once the Completion of Study form is reviewed, the REB will issue an Acknowledgement and the study will automatically be listed in RISe as "Terminated" and will show under your "Inactive" tab. The ONLY activity available from that point on is a Request for Acknowledgement if needed. The study cannot be amended or reactivated.

Commented [u6]: List publications that have reported results from this study. If the final report from this study has not yet been published indicate your plans for such publication.

Attach any supporting documents for the Research Ethics Board by selecting "Add".

Please include the official "close-out letter" from the Sponsor, if applicable.

Note: The REB requires at a minimum, an end-of-study report for all studies at study completion.