

### **REPORTABLE EVENTS**

A **Reportable Event** is any incident or change that happens throughout the course of research that may impact the participants or the conduct of the study.



### **1. REVIEW REPORTING CRITERIA**

For more information on **Reportable Events**, including detailed descriptions of each category and reporting criteria, visit the Research Ethics Office webpage: <u>https://www.ualberta.ca/research/support/ethics-office/human-research-ethics/research-ethics-boards/reb-4/reporting-requirements</u>

# 2. COMPLETE TEMPLATE / REPORT

For categories linked to corresponding templates, you must complete and upload the template as part of the Reportable Event submission.

Category	Reporting Criteria/Description	Reporting Timeline	Template
Serious Adverse Event (SAE) – Local	Local SAEs are adverse events experienced by research participants at the site(s) under the jurisdiction of the REB. A local SAE is reportable if the PI believes it is an unanticipated problem related to the research, and places research participants or others at a greater risk of harm.	Fatal or life-threatening SAEs should be reported within 7 calendar days of the PI becoming aware of them. All other local SAEs should be reported within 15 calendar days of the PI becoming aware of them.	HREB
Serious Adverse Event (SAE) – Non-Local ( <i>REB4 only</i> )	Non-local SAEs are adverse events experienced by research participants at centres/institutions <b>outside</b> the REB's jurisdiction. A non-local SAE is reportable if the PI believes it is an unanticipated problem (unexpected, related or possibly related to participation in the research and places research participants or others at a greater risk of harm) <b>AND</b> requires a change to the protocol and/or informed consent form or immediate notifications to participants for safety reasons.	Within 15 calendar days of the PI becoming aware of the non-local SAE.	<u>See SOP</u>
Protocol Deviation /Violation	<ul> <li>Protocol Deviations/Violations are departures from the procedures set forth in the REB approved application. These include departures that:</li> <li>Compromise the scientific integrity of the study, and/or</li> <li>Constitute or may constitute a potential safety risk to participants enrolled in the protocol or others affected by the research, and/or</li> <li>Are non-compliant with applicable regulations governing human research, and/or</li> <li>Are non-compliant with the requirements or determinations of the REB, or an allegation of such non-compliance, and/or</li> <li>Consist of any unauthorized collection, use, or disclosure of participant personal information.</li> </ul>	Changes to <b>eliminate immediate safety</b> <b>risks</b> to the study participants should be reported within 7 calendar days. <b>All other violations</b> should be reported within 15 calendar days of the PI becoming aware of the deviation/violation.	<u>HREB</u>
Follow-Up Report	Follow-up report requested by the REB if/when more information becomes available, and/or if the issue remained unresolved in the initial report.		
Report	Written report or memorandum from sponsors, such as summary or periodic safety reports, or data safety monitoring board.	Within 15 calendar days of receiving the report.	
Audit	Report any local audit, inspection, or inquiry by a university, provincial or federal agency to the REB. A copy of the audit findings must also be submitted.	Report that audit will be conducted once it is scheduled. Submit the audit report within 15 calendar days of receipt.	
Suspension	Suspension of active and ongoing research by the sponsor, PI, REB or institution.	Report as information is received.	
Participant Complaint	Complaints made by participants or others affected by the research concerning their well-being (psychological or physical) and/or respectful and fair treatment from the researchers.		



#### 3. INITIATE & COMPLETE REPORTABLE EVENT

Any member of the Study Team and Ethics Administrators can initiate a Reportable Event:

- 1. Login to the on-line system: <u>https://arise.ualberta.ca</u>
- 2. Under the Human tab, navigate to the study.

ALBERTA	ARISE Alberta Research Informa	ation Services				
> Dashbo	ard Home					
Applicant My Roles Applicant Supervisors Were Human Study 2 New Incident Report 2 Request Additional Roles	Page for Hal Hartma • Inbox - Items appearing in this tab requir • General - Link to your profile management Inbox Human Animal Displays all your human related ethic submissi Human Studies I am listed on	ID e your action to move an application form space. Incidents Templates ons.	through the review General	r process.		
	Filter by 🖗 ID 💌 Enter te	ut to coarab for	Add Filter	M Clear All		
	Enter to Enter to			= Data Modified	State	Expiration Data
	Pro00086548 test			<ul> <li>Date Modified</li> <li>5/31/2019 10:26 AM</li> </ul>	Approved - RAA Open	Expiration Date
	Pro00086555 lest - to o monstrate how	to response to REVIEWER NOTES		3/19/2019 8:29 AM	Changes Required By REB Administrator	110ay, April 10, 2020
	Pro00086554 Test 1			1/31/2019 1:33 PM	Pre Submission	
	3 items		€ pa	age 1 of 1 🕨		10 / page

- 3. Click on the study you want to open.
- 4. Click on 🕓 New Reportable Event

ALBERTA	ARISE Alberta Res	earch Infor	mation Services					Hello, Hal Hartman 👻
» Dashboa	rd Home							
Current State Approved - PAA Open	Study:test ( P	ro00086548 ) Test Application						
Printer Version	Principal Investigator:	Hal Supervisor				REB Co	ordinator: Kimberley Kordov	
View Differences	Expiration Date:	Friday, April 10, 202	20			Letter of	f Approval: View	
My Activities	REB:	Health Research Et	thics Board - Health Panel			Legacy	Study #:	
ss Change Personnel ss Change Funding ss Copy Study	Approved Study	Amendment (AM	AE10) > Modified Study					
ss Edit Email List	History R	enewals Am	nendments Documents	Change Log	Reportable Events	Related Studies		
Send Email to REB	Ac	c <b>tivity</b>	ed		Author Kordov, Kimberley		- Activity Date	
Create	View Reportable Ev	/ent: Pro00086548_EN	NT4					
New Reportable Event	(i) Re	portable Event Opene	ed		Lo, Patricia		5/31/2019 10:22 AM	
(Approved)	View Reportable Ev	ent: Pro00086548_EN	NT3					
	Re Re	portable Event Opene	d		Supervisor, Hal		5/30/2019 9:44 AM	
	View Reportable Ev	ent: Pro00086548_EN	NT2					
	Re	portable Event Opene	ed		Lo, Patricia		5/28/2019 9:57 AM	
	View Reportable Ev	/ent: Prouu086548_EN A Opened			Supervisor, Hal		4/25/2019 9:42 AM	



5. Enter a meaningful title that will allow you to quickly identify the **Reportable Event**. The title you enter will be auto-populated onto the acknowledgement letter. The **Reportable Event ID** is generated (top right corner) when saved. We recommend the "type of reportable event" followed by a participant ID if it is related to a particular participant (ie: SAE Participant 001; or DSMB letter Nov 2018).

	ERTA	ARISE Alberta Research Inf	ormation Serv	ices	New: Reportable Event
<b>«</b> Back			🖺 Save	🔒 Print	Continue
	Reportab	* Reportable event title:			]

6. Select applicable category. You are able to select multiple categories for one submission. However, consider submitting categories of events (ie: DSMB Reports 2018-Mar 2019) rather than combining unrelated events (ie: SAE Participant 001, DSMB and Sponsor Memo).

Category	Reporting Criteria/Description	Reporting Timeline	Template
Serious Adverse Event (SAE) – Local	Serious adverse events (as defined in ICH) experienced by a research participants at the local site(s) under the jurisdiction of the REB. A local SAE is reportable if the PI believes it is an unanticipated problem (unexpected, related or possibly related to participation in the research and places research participants or others at a greater risk of harm).	Fatal or life-threatening SAEs should be reported within 7 calendar days of the PI becoming aware of them. All other local SAEs should be reported within 15 calendar days of the PI becoming aware of them	Local SAE Report(0.01)
Serious Adverse Event (SAE) – Non-Local	Non-local SAEs are adverse events experienced by research participants at centres/institutions outside the REB's jurisdiction. A non-local SAE is reportable if the PI believes it is an unanticipated problem (unexpected, related or possibly related to participation in the research and places research participants or others at a greater risk of harm) AND requires a change to the protocol and/or informed consent form or immediate notifications to participants for safety reasons.	Within 15 calendar days of the PI becoming aware of the non-local SAE	
Protocol Deviation/Violations	Protocol Deviations/Violations are departures from the procedures set forth in the REB approved application. These include departures that: Compromise the scientific integrity of the study, and/or Constitute or may constitute a potential safety risk to participants enrolled in the protocol or others affected by the research, and/or Are non-compliant with applicable regulations governing human research, and/or Are non-compliant with the requirements or determinations of the REB, or an allegation of such non-compliance. and/or Consist of any unauthorized collection, use, or disclosure of participant personal information.	Changes to eliminate immediate safety risks to the study participants should be reported within 7 calendar days All other violations should be reported within 15 calendar days of the PI becoming aware of the deviation/violation	Protocol- Violation Form(0.01)
Report	Written report or memorandum from <del>study monitors or</del> sponsors, such as summary or periodic safety reports or data safety monitoring board.	Within 15 calendar days of receiving the report	
Audit	Audit, inspection, or inquiry by a university, provincial or federal agency. Only reports with information relevant to the REB should be submitted.	Within 15 calendar days of receiving the audit report	1
Suspension	Suspension of active and ongoing research by the sponsor, PI, REB or institution.		
Participant	Complaints made by participants or others affected by the research concerning their well-		

 Upload completed template or report, if applicable. Otherwise, upload supporting documentation (ie: site Note to File or Memo). Click +Add to upload your file, or drag and drop the file.

3.0	Attach completed template and/or relevant supporting documentation: ( <i>if applicable.</i> )						
	+ Add						
	Document Name	Version	Date	Description			
	There are no items to display						



8. Comments: Enter any additional information you would like to communicate to the REB. Then click the **Continue** button to save and exit the form.

4.0	Comments:	

## 4. SUBMIT REPORTABLE EVENT

Submit the prepared **Reportable Event** to the REB by clicking "Submit Reportable Event" on the left. Enter any comment to the REB in the pop-up, and click OK.

	ARISE	_
	🚱 Execute "Submit Reportable Event" on Pro00086548_EVT5 - Google Chrome	Hello, Hal Hartman 🔫
» Dashboa	https://	
	Submit Reportable Event	☆ Follow
Current State		
Pre Submission	Submit Reportable Event	
Edit Reportable Event	Click <b>OK</b> to submit the application for processing after which you will no longer be able to edit the application. You and your study staff can monitor approval progress on-line. You and your study staff will receive email notification if clarifications, additions or deletions are requested by the reviewers/approvers.	nberley Kordov
Print Reportable Event	Comments:	L
View Differences		
Activities		
PT Submit Reportable Event		
ss Withdraw		
	OK Cancel	
		J

Notes:

- a. If you would like to Withdraw your Reportable Event from review, you can click the Withdraw button (shown above).
- b. After submitting the reportable event, the REB Administrator who manages your study will receive the event for review. The event may be forwarded to the REB Chair for review. Should the REB have any questions pertaining to the event, the Study Team will be notified.

# **FREQUENTLY ASKED QUESTIONS**

- Q: When can a **Reportable Event** be created?
- A: Reportable Events can be created any time after ethics approval. This includes studies that are Completed or Closed by the REB Administrator.

- > Q: I have a renewal or amendment open; can I create and submit a Reportable Event at the same time?
- > A: Yes, **Reportable Events** can be created and submitted when an amendment, renewal, or closure is in process.
- Q: Can I create and submit multiple Reportable Events at the same time?
- A: Yes, multiple Reportable Events can be created and submitted at any time. Reportable Events can be acknowledged by REB in any order submitted.
- > Q: An REB Administrator started a **Reportable Event**, is that permitted?
- A: Yes, Reportable Events can be created by an REB Administrator but they can only submitted by the Study Team. After a reportable event is created, the REB Administrator will send the study team an email within the system to notify them to review and submit the reportable event.
- > Q: What happens if a **Reportable Event** is submitted that doesn't meet the REB reporting standard?
- A: The REB Administrator will send communication back to the Study Team (Changes requested) asking them to Withdraw the Reportable Event.
- Q: How do I find the Reportable Events I created?

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A: Look under the Reportable Events tab of the study.

ALBERTA	ARISE Alberta Research Information Servi	ces	Hello, Hal Hartman 🕶
» Dashboa	d Home		
Current State Approved - PAA Open	Study:test ( Pro00086548 ) Description: Test Application		
Printer Version	Principal Investigator: Hal Supervisor		REB Coordinator: Kimberley Kordov
View Differences	Expiration Date: Friday, April 10, 2020		Letter of Approval: View
My Activities	RED. Health Research Ethics Board - Health Panel		Legacy study #:
ss Change Personnel	Approved Study > Amendment (AME10) > Modified S	Study	
ss Change Funding			
ss Copy Study	The terms Describe Among Security Desc	mante Change Lee Dependeble Frante	Peleked Obudies
ss Edit Email List	History Renewais Amendments Doct	Intents Change Log Reportable Events	Related Studies
ss Edit Guest List	Reportable Events		-
Send Email to REB Coordinator	Filter by 🚱 ID 🔹 Enter text to search for	Q + Add Filter X Clear All	
Create	ID Name	▼Date Modified Owner State	REB PI
New Reportable Event	Pro00086548_EVT7 SAE For Participant 007	6/21/2019 9:22 AM Kordov, Kimberley Approved	Health Research Ethics Board - Health Panel Hal Supervisor
(Approved)	Pro00086548_EVT6 SAE Participant 002	6/20/2019 10:36 Kordov, Kimberley Approved	Health Research Ethics Board - Health Panel Hal Supervisor
	Pro00086548_EVT4 Participant Complaint - Lodged by REB	5/31/2019 10:53 Kordov, Kimberley Con REB Administrat	ve Review Health Research Ethics Board - Health Panel Hal Supervisor
	Pro00086548_EVT3 Protocol Violation Report	5/31/2019 10:50 Kordov, Kimberley Correspondence	Review Health Research Ethics Board - Health Panel Hal Supervisor
	Pro00086548_EVT1 Local SAE - Test	5/31/2019 10:15 Kordov, Kimberley Approved	Health Research Ethics Board - Health Panel Hal Supervisor
	Pro00086548_EVT2 SAE Participant 001	5/31/2019 9:39 AM Kordov, Kimberley Approved	Health Research Ethics Board - Health Panel Hal Supervisor
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