



## FORMAT FOR REPORTING UNANTICIPATED OR SERIOUS ADVERSE EVENTS IN HUMAN RESEARCH PARTICIPANTS AT IIT DELHI

(Please do not delete the headings and subheadings in the attached form)

Institute Ethics Committee

Indian Institute of Technology, Delhi

Ph: 011- 26591221

### Terminology:

**Serious Adverse Event (SAE):** A serious adverse event (SAE) in human drug trials are defined as:

1. Any untoward medical occurrence that at any dose results in
2. Death
3. Is life-threatening
4. Requires inpatient hospitalization or prolongation of existing hospitalization
5. Results in persistent or significant disability/incapacity, or
6. Is a congenital anomaly/birth defect.

### Procedure for reporting:

All the research proposals approved by the institute Ethics Committee of IITD will come under the purview of this policy (drugs, devices, and behavioral or educational interventions; single or multiple armed trials, randomized or non-randomized).

**For all SAE reports:** Within **24 hours** of learning about an unanticipated or serious adverse event, the principal investigator is responsible for notifying the **DCGI, the Study Sponsor (if external), the Institute Ethics Committee ([iitd\\_iec@iitd.ac.in](mailto:iitd_iec@iitd.ac.in))** with a cc to -----  
----- --. A hard copy of this document must also be sent to IEC Secretariat, Institute Ethics Committee Office, Block IV, 3rd Floor, Room 4B9, IIT Delhi.

Within **10 days** the principal investigator is to submit a follow up report to the same list of people as above. **IF IT IS A DEATH REPORT THEN THIS MUST ALSO BE SENT TO THE EXPERT COMMITTEE AND THE HEAD OF THE INSTITUTION (both should have a copy of the original report to the DCGI).**

**Expert Committee address:**

**Chairman,Expert Committee,  
The Drug Controller General of India,  
FDA Bhavan, ITO,Kotla Road,  
New Delhi -110002**

**Within 10 days the completed access database should be sent to -----  
---**

## SERIOUS ADVERSE EVENT FORM

<b>PROTOCOL TITLE:</b>										<b>Protocol ID No.:</b>					<b>Centre:</b>						
<b>Subject's Study No.</b>					<b>Investigation Product:</b>										<b>Report type</b> <div style="display: flex; justify-content: space-around; border: 1px solid black; width: 100px; height: 25px; margin-bottom: 5px;"></div> <b>1.0 = Initial</b> <b>2.1 = follow up 1</b> <b>2.2 = follow up 2 etc</b>						
<b>Occupation:</b>																					
<b>Patient Initials:</b>			<b>Date of birth dd/mm/yy</b>							<b>Age Years</b>		<b>Sex</b>			<b>Height (cm)</b>			<b>Weight (Kg)</b>			
<b>Event onset (dd/mm/yy)</b> <div style="display: flex; justify-content: space-around; border: 1px solid black; width: 150px; height: 25px; margin-top: 5px;"></div>										<b>Adverse Event in MEDICAL TERMS:</b>											
<b>Tick ✓ all appropriate to the Event</b>																					
<b>Patient Died Date: dd/mm/yy</b> <div style="display: flex; justify-content: space-around; border: 1px solid black; width: 150px; height: 25px; margin-top: 5px;"></div>								<div style="border: 1px solid black; width: 40px; height: 25px; margin: 5px auto;"></div>		<b>Life Threatening</b> <div style="border: 1px solid black; width: 40px; height: 25px; margin: 5px auto;"></div>		<b>Prolonged Hospitalization</b> <div style="border: 1px solid black; width: 40px; height: 25px; margin: 5px auto;"></div>		<b>Significant Disability</b> <div style="border: 1px solid black; width: 40px; height: 25px; margin: 5px auto;"></div>		<b>Congenital Abnormality</b> <div style="display: flex; justify-content: space-around; border: 1px solid black; width: 60px; height: 25px; margin-top: 5px;"></div>			<b>Other SAE</b> <div style="border: 1px solid black; width: 40px; height: 25px; margin: 5px auto;"></div>		
<b>Description:</b>																					
<b>Suspected Product(s):</b>										<b>Daily Dose at onset of event:</b>					<b>Route of Administration:</b>						
<b>Indication for use:</b>																					
<b>Therapy dates (from/to), dd/mm/yy:</b>																					
<b>Therapy duration until onset of SAE</b>																					
<b>Was the product stopped? Yes / No</b> <b>If yes, did the event abate after stopping the product? Yes / No/ Not Applicable</b>																					
<b>Serious Adverse Event</b>					<b>Protocol ID No.:</b>					<b>Patient ID No.:</b>											

[illegible]

<b>Serious Adverse Event Report Form Contn.</b>		<b>Protocol ID No.:</b>	<b>Patient ID No.:</b>
Action taken by the Investigator:		Please tick appropriate box	
<input type="checkbox"/>	None	<input type="checkbox"/>	Concomitant drug discontinued
<input type="checkbox"/>	Trial dosage changed	<input type="checkbox"/>	New drug therapy added
<input type="checkbox"/>	Trial drug discontinued	<input type="checkbox"/>	Prolonged hospitalization
<input type="checkbox"/>	Non-drug Therapy		
Outcome:		Please tick appropriate box	
<input type="checkbox"/>	Completely recovered on (dd/mm/yy)	<input type="checkbox"/>	Condition deteriorated
<input type="checkbox"/>	Recovered with sequel	<input type="checkbox"/>	Death, autopsy done (attach summary)
<input type="checkbox"/>	Condition improving	<input type="checkbox"/>	Death, autopsy not done
<input type="checkbox"/>	Condition still unchanged		
Casualty Assessment by investigator (is there any relationship with the test product?):			
<input type="checkbox"/>	Not related	<input type="checkbox"/>	Probable
<input type="checkbox"/>	Unlikely	<input type="checkbox"/>	Most probably
<input type="checkbox"/>	Possible	<input type="checkbox"/>	Insufficient data to assess
Could the SAE be related to the study procedure?:			
<input type="checkbox"/>	Not related	<input type="checkbox"/>	Probable

	Unlikely		Most probably
--	----------	--	---------------

	Possible		Insufficient data to assess
--	----------	--	-----------------------------

What is the long-term prognosis for the patient and will the patient continue to receive treatment? Will the costs of treatment be covered by insurance or other arrangements? (Please describe in detail the arrangements that will be made)

Was the protocol followed in recruitment of the participant? Yes / No

Did the participant meet the exclusion / inclusion criteria of the protocol? Yes / No

Was informed consent obtained as outlined in the protocol? Yes / No ..... If no please explain:

In your opinion, does this report require that the consent form for participants to be revised? Yes / No

If Yes, submit **two** revised consent forms (one soft copy of each and one hard copy). One with the proposed changes emphasized in some fashion (highlighter, bolded, etc.) and another clean copy.

Name, address, telephone and e-mail address of the investigator

Name: \_\_\_\_\_ Profession (speciality): \_\_\_\_\_

Department: \_\_\_\_\_

Tel: \_\_\_\_\_ e-mail: \_\_\_\_\_

Signature of the Investigator reporting the event: \_\_\_\_\_

Reporting date (dd/mm/yy) PLEASE NOTE THAT THIS DATE MUST BE COMPLETED ON THE FORM

\_\_\_\_\_

Date Received by the Research Office, CMC:

\_\_\_\_\_

Signature of the receiver:

\_\_\_\_\_