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).

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

SERIOUS ADVERSE EVENT FORM

PROTOCOL TITLE:	Protocol ID	No.:	Centre:	Centre:			
Subject's Study No.	duct:	Re	port type				
Occupation:			2.1	= Initial = follow up 1 = follow up 2 etc			
Patient Date of birth dd/mm/yy		Age Years	Sex	Height (cm)	Weight (Kg)		
Event onset (dd/mm/yy)	Adverse	Event in MED	ICAL TERMS	:			
Tick ✓ all appropriate to the Event							
Patient Died Date: dd/mm/yy		Prolonged Hospitalization	Disability	Congenital Abnormalit SAE	r		
Description:							
Suspected Product(s):	Dail	y Dose at onset o	of event:	Route of Administratio	n:		
Indication for use:	1						
Therapy dates (from/to), dd/mm	/yy):						
Therapy duration until onset of	SAE						
Was the product stopped? Yes / No If yes, did the event abate after stopping the product? Yes / No/ Not Applicable							
Serious Adverse Event	Protocol ID No			t ID No.:			

	Date Discontinued		tered? Yes / N		comitant Dr	Report Form Were Relevant Conc			
ued For use	I I					TC X7			
ued For use	I I	If Yes, give names and details:							
	(dd/mm/yy)	Continued Y or N	Oate arted nm/yy)	Sta	Dose & Route	Drug Name			
Other Relevant History, laboratory findings and action taken. Medical History (please attach an additional sheet if this space is inadequate):									
				ndings	oratory fin	Relevant test / Labo			
aboratory finding	Comments on laborate		Value	ndings Date	oratory fin Unit	Relevant test / Labo			
aboratory finding	Comments on laborate		Value	_					
aboratory finding	Comments on laborate		Value	_					
laboratory finding	Comments on laborate		Value	_					
laboratory finding	Comments on laborate		Value	_					
laboratory finding	Comments on laborate		Value	_					
	uate):								

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

	Unlikely		Most probably				
			Trost producty				
	D 11		T 00 1 1				
	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		,1 100	ise describe in detail the arrangements	mat will be			
Was	the protocol followed in recruitment of the participant	? Ye	s / 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 vo	our opinion, does this report require that the consent for	rm f	or participants to be revised? Yes / No.				
III yo	ur opinion, does uns report require that the consent rol		participants to be revised. Tes / Tvo				
If Yes, submit <u>two</u> 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	e, address, telephone and e-mail address of the investig	rator					
rain	e, address, terephone and e-man address of the investig	zatoi					
Name	e:Profess	sion	(speciality):				
Department:							
Tel:	e-mail:						
Signature of the Investigator reporting the event:							

Reporting date (dd/mm/yy) PLEASE NOTE THAT THIS DATE MUST BE COM	IPLETED ON THE FORM
Date Received by the Research Office, CMC:	
Signature of the receiver:	