

NCI Contract/Grant No	
IRB Protocol No	

NCI, DIVISION OF CANCER PREVENTION (DCP) SERIOUS ADVERSE EVENT FORM

REQUIRED FIELDS ON ALL REPORTS

Today's Date:		Sponsor: NCI, DCP		Stu	dy (Indication):	
Drug(s) under Investigation	n:	IND No.:				
A. Study Subject Info	rmation					
Study Participant # or PID #	2. Year of Birth:	3. Weight at Time of Event:			4. Height at Time of Event:	
ТЮπ		[] kg [] lbs. [] not available			[] cm [] ft [] not available	
B. Event Information		_				
[] Initial Event Report] Initial Event Report Gender:		circle one) M F		Dose at Event:	
[] Follow-up						
Event Onset Date: (Month/Day/Year)		Primary Event (diagnosis):				
Event Approx. Time: (Indicate A.M./P.M.)						
Event Occurred at:		-				
E diddion of Erag Emposare at Event.			eatment Approx. Time (A.M./P.M.) eatment of Event:):		
Attending Physician (Name Phone/FAX No.: Hospital/Clinic: Address:	e):					
Describe Event (if applicab	ole, include dates of	hospitalizatio	n for event):			
Form Completed by: (Prin	t Name)		Title			
Investigator Signature			Date Phon	ne No		
			(Month/Day/Year)			

SAE Form Revised: 8/09/2006

NCI Contract/Grant No.	
IRB Protocol No.	

ALL FIELDS APPEARING IN THE FOLLOWING PAGES (C-F) MUST BE COMPLETED FOR THE INITIAL REPORT; THEREAFTER, FILL IN ONLY SECTIONS THAT PROVIDE ADDITIONAL/CORRECTIVE INFORMATION.

С.	NIT O	INTOR	mation
·-	SILC	min	mauvn

Investigator Name							
2. Address							
D. Suspect Medication(s)							
Study Design: [] Blind [] Open/Unblind							
Possible Dose (e.g., 300 mg) Free	quency (e.g	g., qd)		Route (e.g., po)		
2. Study Drug	Forr	nulation (e	.g., tablet, s	solution)			
	Lot	No. (If kno	own)				
3. Start Date of Study Drug (Month/Day/Year):							
4. Was blind broken due to event? [] No		[] Yes		[] N	A		
5. Was Study Drug stopped/interrupted/reduced in response to event? [] No [] Yes >> If yes, complete a-e:							
a. If stopped, specify date study drug last taken: (Month/Day	v/Year)	[] NA					
	b. If reduced, specify: New dose Date reduced [] NA (Month/Day/Year)						
c. If interrupted, specify total number of days not given:		[] NA					
d. Did event abate after study drug was stopped or dose reduc	ed?	[] NA	[] Yes	[] No			
e. Did event reappear after study drug was reintroduced? [] NA [] Yes [] No							
6. Was patient taking any other medications concomitantly at the time of the event?[] No [] Yes >> If yes, complete below. (DO NOT LIST DRUGS USED TO TREAT EVENT)							
Drug Name Doses (units, frequency, route, indication for use)	Start Date		Stop Date or mark (X) if continuing			Ş	
	Month	Day	Year	Month	Day	Year	(X)

(continue on a separate sheet if necessary) SAE Form Revised: 8/09/2006

NCI Contract/Grant No	
IRB Protocol No.	

E. Adverse Event 1. Relevant Laboratory/Diagnostic Tests [] No tests performed Results Date Test Actual Value Normal Range Month Day Year (continue on a separate sheet if necessary) 2. Relevant Medical History, including preexisting conditions (e.g., allergies, pregnancy, smoking & alcohol use, hepatic/renal dysfunction, medical/surgical history, etc.) Date (if known) Diseases/Surgeries/Treatment (continue on a separate sheet if necessary) 3. NCI Toxicity GRADE of the Event (use NCI Common Toxicity Criteria): [] 1 [] 2 [] 3 [] 4 [] 5 If not gradable by NCI CTC, check one of the following: [] Mild [] Moderate [] Severe [] Life-threatening [] Fatal 4. Why Serious? [] Results in death Results in persistent or significant disability/incapacity [] Is a congenital anomaly/birth defect [] Other, specify: 5. Outcome of Event (at time of report)

___ [] Improved [] Unchanged [] Worse [] Not available

events and corresponding relationship to study drug in the comments section below.) Check applicable box:

[] Other Investigators participating in this study, if checked, please list names and institutions

Autopsy performed?

6. Investigator's opinion of the relationship between the event and the study drug (If more than one event is being reported, list secondary

[] Possible

Ν

(please attach death certificate and autopsy report, if applicable)

[] Probable

[] Manufacturer/Distributor

[] Definite

(circle one)

SAE Form Revised: 8/09/2006

[] Resolved-date: __

Cause of death:

[] Not related

[] Fatal-date of death: _

(Month/Day/Year)

(Month/Day/Year)

[] Unlikely

7. Was this event reported by the Investigator to (check all that apply): [] IRB

NCI Contract/Grant No	
IRB Protocol No.	

F. Comments/Clarifications:

FOR NCI USE ONLY
1. Date NCI notified of event (Month/Day/Year):
2. Medical Monitor Review:
Medical Assessment of Event (including drug relationship and expectancy):
Is this an FDA reportable (7 calendar days) event? [] Yes [] No
Is this an FDA reportable (15 calendar days) event? [] Yes [] No
>> If No, specify reason:
Is more information expected? [] Yes [] No
>> If Yes, specify:
Is this event to be communicated to other NCI contractors using this investigational drug? [] Yes [] No
>> If Yes, how? By telephone (attach a TC Form): [] Yes, attached TC Form [] No
Other (FAX, mail, e-mail, etc.): [] Yes, attached a copy of the correspondence [] No
Medical Monitor: Print name Signature Date

SAE Form Revised: 8/09/2006