EudraCT number: 2014-000506-35 Amgen Reference: IST-CAR-567 (20159848) FOR UCL CTC US	USE ONLY SAE ID: CAR — (file SAE Report with SAE Sponsor Review For
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SERIOUS ADVERSE EVENT (SAE) REPORT



Please fax this form within 24 hours of becoming aware of the SAE to the Cardamon Trial Coordinator at the CR UK & UCL Cancer Trials Centre on +44 (0)20 7679 9861

Patient detai	ls									
Patient Trial	Number: CAR- Patient initials: Age	e at onset (years):	Sex: M F	Height:	cm w	eight:	L kg			
Site name: Country: United Kingdo										
Type of Repo	ort: Initial d d m m y y y y	Follow-u	 For all follow up reports please: initial & date all changes throughout the report. fax to the trials centre within 24 hours of becoming aware of significant new information. 							
Date site became aware of SAE: d d m m y y y y y Seconding aware of SAE, provide reason: Patient admitted to another hospital Patient admitted to another properties of the composition of the composi										
Serious Eve	nts (list serious events only)		Continued on a separate sheet?	Total	Total No. of Events					
					Causal relation to event ² : (enter one code for each treatment)					
Event No.	Event Name (refer to CTCAE v4.03)	Severity Grade (CTCAE v4.03)	Dates of Onset & Resolution dd-mm-yyyy	Outcome of Event ¹	Carfilzomib	Cyclophosphamide (N/A in maintenance)	Dexamethasone (N/A in maintenance)			
1			Onset Resolution Resolution							
2			Onset Resolution Resolution							
3			Onset Resolution Resolution							
4			Onset Resolution Resolution							
5			Onset Resolution Resolution							
Codes: (1) Outcome of Event (enter one code per event): 1 = Fatal 2 = Not Resolved 3 = Resolved 4 = Resolved with Sequelae 5 = Resolving (2) Causal Relationship (enter one code): 0 = Not related (no reasonable possibility) 1 = Related (reasonable possibility)										

CARDAMON SAE Report									Patient Trial Number: CAR- Initials Initials	
Why was the SAE serious? (tick all that apply – please refer to the Pharmacovigilance section of the protocol for details)										
		Resulted in	n death							
		Life threa	atening							
Required new or prolonged hospitalisation For new						v hospitalisations only: dd-mm-yyyy admission: Date of discharge:				
Resulted in persistent or significant disability/incapacity										
Resulte	Resulted in congenital anomaly or birth defect									
Other medically significant (e.g. non-serious adverse events of special interest) specify										
If patient has died: Date of death: d d m m y y y y y Cause of death: (specify) (this may be requested by UCL CTC if required)										
Most recent treatment phase: (please select one) (please select on										
	Manufacturer ອີດ ໄດ້ Name ອີດ ໄດ້			Total Daily Dose	ly	_		Overdose¹	Date of First Administration of IMP AND Date of Last Administration of IMP Prior to this Dechallenge/Rechallenge (complete if dose reduced/treatment stopped) Y or N or N/A	
Name	AND Brand Name	Batch Numb	ch Numk ingth (in	Prior to SAE (include units)	Prior to SAE Sinclude S	Formulation (delete as appropriate)	Route	Treatment Ove	Action Taken (s) miproved after stopping or reducing treatment to make a rechallenge performed? Five (stopping or reducing treatment continued) If yes, did the event(s) reappear or reappear or reappear or reducing reducing reappear or reducing reduci	
Carfilzomib	Amgen		60mg			Lyophisilate for solution	IV		First Ongoing Last Ongoing Final Ongoing	
Cyclophosphamide					Days 1, 8 & 15 of 28 day cycle	Tablet OR Powder for solution for injection or infusion			First Ongoing Last Ongoing Final Ongoing	
Dexamethasone					Days 1, 8, 15 & 22 of 28 day cycle	injection			First Ongoing Last Final Final	
Codes: (1) Enter one code: 0 = no overdose 1 = dosing/administration error by site 2 = accidental/intentional overdose by patient 3 = Other (specify)										
ASCT (if patient randomised to this Arm)										
Start date of Melphalan conditioning: Date of Transplant Day 0:										

CARDAMON SAE Report

Any tests/laboratory data applicable to this SAE? If yes, specify below Continued on a separate sheet? Normal range, if applicable Results Pending Date Test Results (check box if result has (specify and include units, if (specify) dd-mm-yyyy (specify units and normal ranges) applicable) not yet been provided) Any non-serious events relevant to this case? If yes, list Adverse Event Term below, with start and stop dates Any relevant medical history/concurrent conditions? (If yes, Please specify details below) Treatment for SAE? If yes, specify below, include: drug name, indication, formulation, dose (including unit), frequency, route, start and end dates Continued on a separate sheet?

Patient Trial Number: CAR-

Initials

CARDAMON SAE Report	Patient Trial Number:	CAR- Initials			
Concomitant medications? If yes please complete the Concomitant Medication Sheet.					
Were any SAEs listed above related to a concomitant medication?	details of adverse event term, drug name, and if there was an interac	ction with the IMP			
Event Name (State term as provided in Serious Events section)	Concomitant Medication (list which concomitant medication is related to adverse ev				
(State term as provided in Section)	(list which concomitant medication is related to adverse ev				
		$\square_{N} \square_{Y}$			
		$\square_{N} \square_{Y}$			
		□ _N □ _Y			
Case Narrative (Give a concise medical description of the events including all relevant signs and symptoms, bo event and trial treatment, include medical judgement considering all relevant factors.)	dy systems, and any additional information deemed relevant to the ca	ase. State the rationale for causal relationship between the			
		Continued on a separate sheet?			
Investigator Assessment: (must be authorised on staff delegation log to review SAEs and perform evaluations of ca	ausal relationship)				
Print Signature:		e of mpletion:			
Form(s) completed by: (must be authorised on staff delegation log to complete CRFs and report SAEs)					
Print Signature:		e of mpletion:			

Date form entered: _

_____ Date form checked: ____

For UCL CTC use only: Date form received: _

Initials: __