

Carfilzomib/Cyclophosphamide/Dexamethasone with maintenance carfilzomib in untreated transplant-eligible patients with symptomatic MM to evaluate the benefit of upfront ASCT

End of Maintenance Form

Patient Initials

Trial Number

C	A	R	_	

(This form has 5 pages including cover sheet)

Please send forms to:

Cardamon Trial Coordinator CR UK & UCL Cancer Trials Centre 90 Tottenham Court Road London W1T 4TJ

General enquires: **020 7679 9860** Randomisations: **020 7679 9860** between 9.00am and 5.00pm Fax: **020 7679 9861** E-mail: <u>ctc.cardamon@ucl.ac.uk</u>



Cancer Research UK and UCL Cancer Trials Centre







Additional instructions for completing forms

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The End of Maintenance form collects details of the patient's response to maintenance treatment. Assessments are to be performed within 14 days of completing the last cycle of maintenance.

Specific Fields

- Please ensure that you are using the correct units (i.e. haemoglobin in g/dL). If your local report uses different units please convert these before entering them on the form
- If any efficacy tests have not been done because they are not clinically indication, please ensure that you complete the boxes with ND to confirm that the tests were not done. A discrepancy will be raised for those fields left completely blank
- Disease response should be confirmed by a local investigator
- Please ensure a progression/relapse form is submitted for patients with progressive disease

Completing forms

- Ensure all entries are clear, legible and written in black ink
- Avoid the use of abbreviations and acronyms
- Do not leave any fields blank. In case of missing data
 - ND (not done) if a test has not been performed or a measure not taken. If applicable state the reason
 - NA (not applicable) if a measure is not applicable
 - NK (not known) if data is unknown. This should only be used once every effort to obtain the data has been exhausted
- The Principal Investigator (PI) is responsible for the accuracy of the data reported on the CRF
- Please ensure that all adverse events are recorded on the adverse event form and the form is attached
- CRFs may only be completed by an appropriately qualified individual delegated as responsible by the PI on the site delegation log
- CRF Footer section
 - The "completed by" Name should be legible
 - Each CRF should be signed and dated by the person completing the form
 - Do not complete the UCL CTC Use only section
- The CRF should be sent/faxed to the Cancer Trials Centre (CTC) with a copy retained at the Site (ensure when photocopying the page that the copy is added to the CRF booklet in the same place where the original was stored)

If you have any questions about how to complete this form please contact the Cardamon Trial Coordinator on: 020 7679 9860





Cardamon Trial Number C A R - Patient Initials	
End of Maintenance Form	Page 3 of 5
Adverse events	
Did the patient experience any adverse events between their last cycle of maintenance and their end of maintenance assessment? 1 = Yes (please ensure adverse event form is submit 2 = No	ted)
Pregnancy test (for females of child bearing potential only)	
Result:1 = Negative 2 = Positive 3 = Not applicableDate of pregnancy testDDMMYYYY	
Bone marrow biopsies (to confirm CR only)	
Bone marrow aspirate Date of sample D D M M Y Y Y Y	
1= Present, complete % of plasma cells: 2= Present , not measured 3= Absent 4= Not done 4= Not done	
Bone marrow trephine Date of sample D D M M Y Y Y Y	
1= Present, complete % of plasma cells: 2= Present , not measured 3= Absent 4= Not done	
If No to the above, specify a reason:	
Soft tissue plasmacytoma/Extramedullary lesions	
Does the patient have any soft tissue plasmacytomas/ 1= Yes, complete date of test and a separate line for ea Extramedullary lesions? 2= No	ch site involved
If yes, date of test D D M M Y Y Y Y L Long ax	is Short axis
Site involved: Bidimensional measurements (cm):	x
Site involved: Bidimensional measurements (cm):	x
Site involved: Bidimensional measurements (cm):	x

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Cancer Research UK and UCL Cancer Trials Centre



Cardamon	Trial Number CAF		Patient Initials
End of Maintenance	Form		Page 4 of 5
Efficacy assessments			
Date of test D D M	M Y Y Y Y		
	e <u>one</u> option only)	Single paraprotein expressed Light chain only Biclonal Non-secretory	
Paraprotein type key: 1 = IgG, 2 = IgA	, 3 = IgM, 4 = IgD		
Specify paraprotein type:	Serum paraprotein	4= Present, please complete r 5= Too faint to quantify 6= Absent 7= Not Done	esult (g/L)
Specify paraprotein type:	Serum paraprotein	4= Present, please complete r 5= Too faint to quantify 6= Absent 7= Not Done	esult (g/L)
Serum free light chain: Kappa (mg/L	.)	OR	Tick if not done
Serum free light chain: Lambda (mg	/L)	OR	Tick if not done
Serum free light chain Kappa/Lambda ratio:		nal range of a/Lambda FLC ratio:	_
Urinary light chain measurement			
1= Present, quantifiable <i>Please complete 24h</i> 2= Too faint to quantify (24h BJ 3= Absent 4= Not done 5= Present, not formally quanti (<i>if unable to perform 24h BJP</i>)	P only)	• (plea	chain type se choose one_only): 1= Kappa 2= Lambda 3 = N/A
Immunofixation (only to confi	rm CR)		
Immunofixation Serum	1= Positive 2= Negative Date of tes 3= Not done	t D D M M Y	Y Y Y
Immunofixation Urine	1= Positive 2= Negative Date of tes 3= Not done	t D D M M Y	Y Y Y

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CANCER RESEARCH UK	Cancer Research L	JK and UCL Cancer Trials	Centre	CL
Cardamon	Trial Number	AR –	Patient Initials	
	ntenance Form ally indicated or for response		- 6 + +	-
MRI	1= Evidence of myeloma 2= No evidence of myeloma 3= Not done	D D M M		
ст	1= Evidence of myeloma 2= No evidence of myeloma 3= Not done	D D M M	Y Y Y Y	
PET	1= Evidence of myeloma 2= No evidence of myeloma 3= Not done	D D M M	Y Y Y Y	
Skeletal survey	1= Evidence of myeloma 2= No evidence of myeloma 3= Not done	D D M M	Y Y Y Y	
Other imaging	1= Evidence of myeloma 2= No evidence of myeloma 3= Not done	D D M M	Y Y Y Y	
Spec	ify type of other imaging			
	number or size of lytic bone lesion	is been seen on any radiograp	1 = Yes 2 = No	
Response at the er Date of response asse	D	D M M Y Y	Y Y	
Patient's response	to maintenance treatment: (choose <u>one</u> option only)	1= sCR 2= CR 3= VGPR 4= PR 5= MR 6= SD 7= PD 8= Unable to assess—Specify	reason:	
Investigator name (print):		Investigator signature:		
		Date signed:	D M M Y Y Y	Y
Name of person completing	g form: Signature of pe	erson completing form:	Date completed:	
				Y
The site PI or delegated in Investigator name:	nvestigator must sign to confirm that inforn Investigator sig		Date completed:	
				Y

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