

# Cardamon

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	<input type="text"/>	<input type="text"/>	<input type="text"/>				
Trial Number	<input type="text" value="C"/>	<input type="text" value="A"/>	<input type="text" value="R"/>	-	<input type="text"/>	<input type="text"/>	<input type="text"/>

**(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: **[ctc.cardamon@ucl.ac.uk](mailto:ctc.cardamon@ucl.ac.uk)**



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**

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### 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 submitted)  
2 = No

### Pregnancy test (for females of child bearing potential only)

Result:  1 = Negative  
2 = Positive  
3 = Not applicable

Date of pregnancy test

### Bone marrow biopsies (to confirm CR only)

#### Bone marrow aspirate

Date of sample

1= Present, complete % of plasma cells:  
2= Present, not measured  
3= Absent  
4= Not done

%

#### Bone marrow trephine

Date of sample

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/  
Extramedullary lesions?

1= Yes, complete date of test and a separate line for each site involved  
2= No

If yes, date of test

Site involved:

Bidimensional measurements (cm):

Long axis Short axis  
 X

Site involved:

Bidimensional measurements (cm):

X

Site involved:

Bidimensional measurements (cm):

X

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## Efficacy assessments

Date of test

Please complete this section for all myeloma patients:

Paraprotein expression (choose one option only)

- 1= Single paraprotein expressed
- 2= Light chain only
- 3= Biclinal
- 4= Non-secretory

Paraprotein type key: 1 = IgG, 2 = IgA, 3 = IgM, 4 = IgD

Specify paraprotein type:  Serum paraprotein  4= Present, please complete result  (g/L)  
 5= Too faint to quantify  
 6= Absent  
 7= Not Done

Specify paraprotein type:  Serum paraprotein  4= Present, please complete result  (g/L)  
 (If biclinal) 5= Too faint to quantify  
 6= Absent  
 7= Not Done

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:  •  Normal range of Kappa/Lambda FLC ratio:  –

## Urinary light chain measurement

1= Present, quantifiable  
 Please complete 24h BJP result (in g/24h):  •  Light chain type  1= Kappa  
 2= Too faint to quantify (24h BJP only) 2= Lambda  
 3= Absent 3= N/A  
 4= Not done  
 5= Present, not formally quantified (if unable to perform 24h BJP)

## Immunofixation (only to confirm CR)

Immunofixation Serum  1= Positive  
 2= Negative  
 3= Not done

Date of test

Immunofixation Urine  1= Positive  
 2= Negative  
 3= Not done

Date of test

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*Imaging (If clinically indicated or for response assessment if persistent soft tissue plasmacytomas present)*

		Date of test	Lytic or focal lesions? 1= Yes 2= No
MRI	<input type="checkbox"/> 1= Evidence of myeloma <input type="checkbox"/> 2= No evidence of myeloma <input type="checkbox"/> 3= Not done	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/>	<input type="text"/>
CT	<input type="checkbox"/> 1= Evidence of myeloma <input type="checkbox"/> 2= No evidence of myeloma <input type="checkbox"/> 3= Not done	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/>	<input type="text"/>
PET	<input type="checkbox"/> 1= Evidence of myeloma <input type="checkbox"/> 2= No evidence of myeloma <input type="checkbox"/> 3= Not done	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/>	<input type="text"/>
Skeletal survey	<input type="checkbox"/> 1= Evidence of myeloma <input type="checkbox"/> 2= No evidence of myeloma <input type="checkbox"/> 3= Not done	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/>	<input type="text"/>
Other imaging	<input type="checkbox"/> 1= Evidence of myeloma <input type="checkbox"/> 2= No evidence of myeloma <input type="checkbox"/> 3= Not done	<input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/> <input type="text"/>	<input type="text"/>

Specify type of other imaging

Has an increase in number or size of lytic bone lesions been seen on any radiograph?  1 = Yes  2 = No

### Response at the end of maintenance

Date of response assessment

Patient's response to maintenance treatment:  1= sCR  
(choose one option only)  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:

Name of person completing form:  Signature of person completing form:  Date completed:

*The site PI or delegated investigator must sign to confirm that information within the CRF is accurate*

Investigator name:  Investigator signature:  Date completed: