

# ARISTOTLE

Trial Number:

Patient Initials:

## FOLLOW UP FORM

☐ 24 months

☐ 36 months

**NB** All dates should be in DD-MM-YYYY format

FU24-36-1/3

### INSTRUCTIONS:

If confirmed residual disease or recurrence has been reported prior to this visit, please complete sections 1 and 3 **only**.

### 1. DATE OF LAST CONTACT

A) Date patient last known to be alive:  /  /

i) How was this information obtained? (*tick one box only*)

- ☐ Patient attended for clinic visit    ☐ Telephone call with patient    ☐ Via GP    ☐ Via local hospital  
☐ Via hospice    ☐ Other (*please specify*) \_\_\_\_\_

*If the patient has died since the last visit please submit a death form*

### 2. PHYSICAL EXAMINATION

☐ N/A (patient did not attend)

A) Weight (kg): .

B) ECOG Performance Status:

### 3. DISEASE STATUS

A) Has the patient's disease recurred since the last visit (*tick one box only*)?

- ☐ Yes—suspected (*complete a Suspected Residual Disease or Recurrence form*)    ☐ Unknown  
☐ Yes—confirmed (*complete a Confirmed Residual Disease or Recurrence form*)    ☐ N/A- patient had residual disease after CRT ± Surgery  
☐ No    ☐ N/A—confirmed recurrence reported previously

### 4. SURGERY

A) Has the patient had surgery for their rectal cancer since the last visit?    ☐ Yes\*    ☐ No

***\*If the patient has had surgery for their rectal cancer, please submit a Surgery Form***

**For UCL CTC use only:** Date Checked: \_\_\_\_\_ Initials: \_\_\_\_\_ Date entered: \_\_\_\_\_ Initials: \_\_\_\_\_

# ARISTOTLE

Trial Number:

A

R

I




Patient Initials:




## FOLLOW UP FORM

☐ 24 months

☐ 36 months

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FU24-36-2/3

### 5. CHEMOTHERAPY

A) Has the patient started any new chemotherapy regimens since the last visit? ☐ Yes ☐ No ☐ Unknown

i) If the patient started a new regimen, give start date of chemotherapy: 
 / 
 /

ii) If yes, what was the function of the regimen

☐ Adjuvant

☐ Palliative

☐ Other (please specify) \_\_\_\_\_

iii) If yes, give details of the regimen

☐ 5FU

☐ Capecitabine

☐ Oxaliplatin + Capecitabine

☐ Oxaliplatin + 5FU

☐ Other (please specify) \_\_\_\_\_

B) Has the patient stopped any ongoing chemotherapy regimens since the last visit? ☐ Yes ☐ No ☐ Unknown

i) If yes, give date of completion of chemotherapy: 
 / 
 /

ii) If yes, give duration of chemotherapy: 
 months

### 6. SURGERY FOR LATE EFFECTS OF RADIOTHERAPY

A) Has the patient required surgery for late effects of radiotherapy since the last visit? ☐ Yes ☐ No ☐ Unknown

i) If yes, date of surgery: 
 / 
 /

ii) If yes, was this for:

☐ Bleeding

☐ Blockage (stricture)

☐ Other (please specify) \_\_\_\_\_

### 7. CT SCAN

Date of scan: 
 / 
 / 



 ☐ No metastatic lesions detected (see point1 below)

Lesion No	Site (please provide a detailed description, e.g. Left lower lobe lung)	Lesion assessment E = equivocal U = unequivocal NV = not visible
1		
2		
3		
4		
5		
6		
7		
8		

#### Instructions:

- Please list sites of metastatic disease only (NB If patient has evidence of recurrence, please complete a Suspected or Confirmed Residual Disease or Recurrence Form). If none detected, tick the box above.
- Once a lesion has been added to the table, it should not be removed from subsequent forms. If it is not visible on subsequent scans, enter NV.
- Please assign the same lesion the same lesion number throughout.
- If there is more than one lesion in the same site, please record only up to a maximum of two lesions per site in the table.

For UCL CTC use only: Date Checked: \_\_\_\_\_ Initials: \_\_\_\_\_ Date entered: \_\_\_\_\_ Initials: \_\_\_\_\_

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Patient Initials:

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### 8. SECOND (UNRELATED) MALIGNANCY

A) Has the patient developed a second unrelated malignancy since the last visit? ☐ Yes ☐ No

i) If yes, specify type: \_\_\_\_\_

ii) Is there histological evidence? ☐ Yes ☐ No

**Please send a copy of the pathology report to the Trials Centre if available.**

### 9. PELVIC FUNCTIONAL QUESTIONNAIRE

A) Has the patient completed a pelvic functional questionnaire? ☐ Yes ☐ No

i) If **yes**, date questionnaire completed: 

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 / 

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 / 

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ii) If **no**, please give reason: ☐ Patient too ill ☐ Patient refused ☐ Patient did not consent ☐ Administrative error  
☐ Other (specify): \_\_\_\_\_

### 10. DYSFUNCTIONING STOMA

A) Does the patient have a dysfunctioning stoma? Yes ☐ No ☐

i) If yes, what type: Ileostomy ☐ Colostomy ☐

ii) If yes, is it intended to be temporary or permanent? Temporary ☐ Permanent ☐

### 11. ASPIRIN

A) Has the patient been entered onto the Add-Aspirin trial? Yes ☐ No ☐

i) If no, is the patient taking aspirin? Yes ☐ No ☐

Completed by (print name):

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Signature:

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Date Completed:

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Site:

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For UCL CTC use only: Date Checked: \_\_\_\_\_ Initials: \_\_\_\_\_ Date entered: \_\_\_\_\_ Initials: \_\_\_\_\_

ARISTOTLE Follow Up Form - 24/36 months, v5, 05/01/2018