# Randomised phase III study of docetaxel versus active symptom control in patients with relapsed gastric adenocarcinoma

Submission date	Recruitment status No longer recruiting	[X] Prospectively registered		
19/09/2007		<pre>Protocol</pre>		
Registration date	Overall study status	Statistical analysis plan		
14/11/2007	Completed	[X] Results		
<b>Last Edited</b> 19/03/2020	<b>Condition category</b> Cancer	[] Individual participant data		

# Plain English summary of protocol

http://www.cancerhelp.org.uk/trials/a-trial-looking-at-docetaxel-chemotherapy-for-advanced-cancer-of-the-stomach-the-oesophagus-or-the-junction-of-the-stomach-and-oesophagus

# Contact information

# Type(s)

Scientific

#### Contact name

Dr Hugo Ford

#### Contact details

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# Additional identifiers

# EudraCT/CTIS number

2006-005046-37

IRAS number

# ClinicalTrials.gov number

NCT00978549

### Secondary identifying numbers

C21276/A7737

# Study information

#### Scientific Title

Randomised phase III study of docetaxel versus active symptom control in patients with relapsed gastric adenocarcinoma

#### Acronym

COUGAR-02

#### Study objectives

To establish the role of chemotherapy with docetaxel as second line therapy for advanced gastric cancer in terms of overall survival.

On 12/09/2008 the overall trial start and end dates were changed from 01/11/2007 and 30/04/2010 to 10/04/2008 and 10/10/2010, respectively.

On 15/02/2011 the following changes were made to this trial record:

- 1. The overall trial end date was changed from 10/10/2010 to 28/02/2012.
- 2. The target participant number was changed from 320 to 180.

#### Ethics approval required

Old ethics approval format

### Ethics approval(s)

South West Research Ethics Committee, 28/12/2008, ref: 07/H0206/62

# Study design

Open-label two-arm multi-centre phase III randomised controlled trial

# Primary study design

Interventional

# Secondary study design

Randomised controlled trial

# Study setting(s)

Hospital

# Study type(s)

Treatment

#### Participant information sheet

# Health condition(s) or problem(s) studied

Advanced gastric cancer

#### **Interventions**

Arm A: chemotherapy with docetaxel plus active symptom control for 18 weeks. The participants in this arm will have treatment once every 3 weeks and will receive up to 6 cycles of docetaxel intravenously at a dose of 75 mg/m<sup>2</sup> (total of 18 weeks of treatment).

Arm B: active symptom control for 18 weeks. Active symptom control may include administration of analgesics, anti-emetics, steroids, palliative radiotherapy and/or any other supportive measures deemed appropriate by the treating clinician.

#### Intervention Type

Drug

#### **Phase**

Phase III

# Drug/device/biological/vaccine name(s)

Docetaxel

#### Primary outcome measure

Overall survival, measured as the time between the date of randomisation and the date of death from any cause.

#### Secondary outcome measures

- 1. Time to documented progression
- 2. Response rates to docetaxel. This will be assessed using the Response Criteria In Solid Tumours (RECIST) at baseline and after 3 and 6 cycles of treatment. The best overall response achieved over the treatment period will be determined
- 3. Toxicity of docetaxel. Docetaxel related toxicity data will be collected at the end of each of the chemotherapy cycles prior to the start of the next cycle. In addition, all Adverse Events (AEs) will be monitored and recorded from randomisation until 21 days after the last administration of docetaxel
- 4. Quality of life, assessed using the European Organisation for Research and Treatment of Cancer (EORTC) Quality of Life Questionnaire QLQ-C30 vs3 and QLQ-STO22. Quality of life data will be collected at baseline and then after 3, 6, 9, 12, 18 and 24 weeks from start of treatment 5. Health economic evaluation. The EuroQoL (EQ-5D) health outcome instrument is to be used for this assessment. The EQ-5D will be completed at baseline and then at every outpatient visit until death

# Overall study start date

10/04/2008

# Completion date

10/10/2010

# **Eligibility**

# Key inclusion criteria

- 1. Patients with histologically confirmed adenocarcinoma of the stomach (including Siewert-Stein type II and III adenocarcinoma of the oesophago-gastric junction)
- 2. Age 18 years or older
- 3. Advanced disease not amenable to curative treatment

- 4. Documented progressive disease while receiving or within 6 months of completion of first line chemotherapy with a platinum and fluoropyrimidine based therapy either for advanced disease or as neoadjuvant/perioperative therapy
- 5. Estimated life expectancy greater than or equal to 12 weeks
- 6. Eastern Cooperative Oncology Group (ECOG) performance status 0, 1 or 2
- 7. Satisfactory haematologic (haemoglobin [Hb] greater than or equal to 10 g/dL, White Blood Cells [WBC] greater than or equal to 3.0 x  $10^9$ /L, Absolute Neutrophil Count (ANC) greater than or equal to  $1.5 \times 10^9$ /L, Platelets (Plt) greater than or equal to  $100 \times 10^9$ /L), renal (creatinine less than or equal to Upper Limit of Normal [ULN]) and hepatic (total Bilirubin less than or equal to ULN, Alanine aminotransferase (ALT) less than or equal to  $1.5 \times ULN$ , Alkaline Phosphatase (ALP) less than or equal to  $5 \times ULN$ ) function
- 8. Ability to give informed consent
- 9. Completion of baseline questionnaires (European Organisation for Research and Treatment of Cancer Quality of Life Questionnaire [EORTC QLQ-C30] and EORTC-QLQ-STO22 [gastric cancerspecific QLQ] and EuroQoL (EQ-5D) questionnaire)
- 10. Patients of both sexes with reproductive potential must be willing to employ barrier contraceptives whilst on treatment and for 3 months after completion of treatment

#### Participant type(s)

**Patient** 

#### Age group

Adult

#### Lower age limit

18 Years

#### Sex

Both

# Target number of participants

180

#### Key exclusion criteria

- 1. Cerebral or leptomeningeal metastasis
- 2. Prior chemotherapy with taxanes
- 3. Peripheral neuropathy
- 4. More than one prior chemotherapy regimen in advanced setting
- 5. Previous malignancy except for curatively treated basal cell carcinoma of the skin or cervical intraepithelial neoplasia
- 6. Pregnant or breastfeeding female patient
- 7. Any medical or psychiatric condition which would influence the ability to provide informed consent
- 8. Any other serious or uncontrolled illness which, in the opinion of the investigator, makes it undesirable for the patient to enter the trial

#### Date of first enrolment

10/04/2008

# Date of final enrolment

10/10/2010

# Locations

## Countries of recruitment

England

**United Kingdom** 

Study participating centre Addenbrooke's Hospital Cambridge

Cambridge United Kingdom CB2 0QQ

# Sponsor information

# Organisation

Cambridge University Hospitals NHS Foundation Trust (UK)

# Sponsor details

Box 277
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England
United Kingdom
CB2 0QQ
+44 (0)1223 245151
webmaster@addenbrookes.nhs.uk

#### Sponsor type

Hospital/treatment centre

#### Website

http://www.addenbrookes.org.uk/

#### **ROR**

https://ror.org/04v54gj93

# Funder(s)

# Funder type

Charity

#### Funder Name

Cancer Research UK (UK) (ref: C21276/A7737)

#### Alternative Name(s)

CR\_UK, Cancer Research UK - London, CRUK

# **Funding Body Type**

Private sector organisation

## **Funding Body Subtype**

Other non-profit organizations

#### Location

United Kingdom

# **Results and Publications**

# Publication and dissemination plan

Not provided at time of registration

# Intention to publish date

Individual participant data (IPD) sharing plan

#### IPD sharing plan summary

Not provided at time of registration

# **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Basic results				No	No
Plain English results				No	Yes
Results article	results	01/01/2014		Yes	No