# An international, five-arm randomised trial of carboplatin and paclitaxel versus triplet or sequential doublet combination in patients with epithelial ovarian cancer or primary peritoneal carcinoma

Submission date	Recruitment status No longer recruiting	[X] Prospectively registered		
18/05/2001		☐ Protocol		
Registration date 18/05/2001	Overall study status Completed	Statistical analysis plan		
		[X] Results		
Last Edited	Condition category	Individual participant data		
22/10/2018	Cancer			

# Plain English summary of protocol

Not provided at time of registration

# Contact information

# Type(s)

Scientific

#### Contact name

Dr P Harper

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# Additional identifiers

ClinicalTrials.gov (NCT) NCT00011986

#### Protocol serial number

E164/58

# Study information

#### Scientific Title

An international, five-arm randomised trial of carboplatin and paclitaxel versus triplet or sequential doublet combination in patients with epithelial ovarian cancer or primary peritoneal carcinoma

#### Acronym

ICON5/GOG182

#### **Study objectives**

To evaluate a variety of chemotherapy regimes for patients with advanced stage (FIGO III-IV) epithelial ovarian or serious primary peritoneal carcinoma. Efficacy will be determined through analysis of overall survival and progression free survival.

The trial will also compare the toxicities and adverse effects of each treatment regimen, describe the dose density and collative dose delivery for each regime, compare response rates in patients with measurable disease. In the UK, evaluate the impact of regimes on quality of life, evaluate the impact of regimes on resource use and quality-adjusted life-years, collect and store genetic material for future studies of molecular genetics.

The trial includes two stages, at the end of the first stage only those treatment arms that appear promising on the basis of progression free survival will continue in to the second stage which aims to evaluate the impact of the regimes on overall survival.

## Ethics approval required

Old ethics approval format

# Ethics approval(s)

Not provided at time of registration

# Study design

Randomised controlled trial

# Primary study design

Interventional

# Study type(s)

Treatment

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

Ovarian cancer and peritoneal carcinoma

#### **Interventions**

Arm I: Taxol 175 mg/m<sup>2</sup> day one, Carboplatin AUC6 or AUC(EDTA)5 iv day one, eight cycles Arm II: Taxol 175 mg/m<sup>2</sup>, day one, Gemzar 800 mg/m<sup>2</sup> days one and eight, Carboplatin AUC5 or AUC(EDTA)4, day one, eight cycles

Arm III: Taxol 175 mg/m<sup>2</sup> day one, Caelyx 30 mg/m<sup>2</sup> day one every other cycle, Carboplatin AUC5 or AUC(EDTA)4 day one, eight cycles

Arm IV: Hycamtin 1.25 mg/m^2 days one, two and three, Carboplatin AUC5 or AUC(EDTA)4 day three, four cycles, then four cycles of Arm I

Arm V: Gemzar 1000 mg/m $^2$ , days one and eight, Carboplatin AUC6 or AUC(EDTA)5 day eight, four cycles then four cycles of Arm I

In every case cycles are 21 days long. Chemotherapy continues unless disease progression or unacceptable toxicity occurs. Dose reductions or delays for toxicity are defined in the full protocol.

#### Intervention Type

Drug

#### Phase

Not Applicable

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

Carboplatin and paclitaxel

#### Primary outcome(s)

Overall survival.

#### Key secondary outcome(s))

- 1. Progression free survival
- 2. Response rate (in patients with measurable disease)
- 3. Toxicity and symptoms
- 4. Dose and dose intensity
- 5. Patients assessment of quality of life and acceptability of treatment, health economics
- 6. Molecular genetics (future study)

# Completion date

31/12/2007

# **Eligibility**

## Key inclusion criteria

- 1. Stage III or IV ovarian or serious primary peritoneal carcinoma, following appropriate surgery
- 2. Tumor tissue available for histological evaluation
- 3. Adequate bone marrow liver kidney and neurological function
- 4. World Health Organisation (WHO) performance status zero to two
- 5. Fit and able to take part in trial treatments and follow-up
- 6. Informed consent

# Participant type(s)

Patient

# Healthy volunteers allowed

No

#### Age group

Adult

#### Sex

All

#### Key exclusion criteria

- 1. Concomitant or previous malignancies likely to interfere with protocol treatments
- 2. Patient has received radiotherapy or chemotherapy to any abdominal or pelvic tumour
- 3. Acute hepatitis, infection or Gastrointestinal bleeding
- 4. Women of childbearing age who will not use adequate contraception or are breastfeeding

#### Date of first enrolment

01/03/2002

#### Date of final enrolment

31/12/2007

# Locations

#### Countries of recruitment

**United Kingdom** 

England

United States of America

# Study participating centre Medical Oncology

London United Kingdom SE1 9RT

# Sponsor information

#### Organisation

Medical Research Council (MRC) (UK)

# Funder(s)

## Funder type

Research council

#### Funder Name

Medical Research Council (MRC) (UK)

## Alternative Name(s)

Medical Research Council (United Kingdom), UK Medical Research Council, MRC

## **Funding Body Type**

Government organisation

# Funding Body Subtype

National government

#### Location

**United Kingdom** 

# **Results and Publications**

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?
Results article	results:	01/08/2003		Yes	No
Results article	results:	20/03/2009		Yes	No
Plain English results				No	Yes
Study website	Study website	11/11/2025	11/11/2025	No	Yes