# A randomised trial of combination bleomycin, ifosfamide and cisplatinum versus single agent cisplatinum in recurrent cervical cancer

Submission date Recruitment status Prospectively registered 01/07/2001 No longer recruiting [ ] Protocol [ ] Statistical analysis plan Registration date Overall study status 01/07/2001 Completed [X] Results [ ] Individual participant data Last Edited Condition category 04/01/2012 Cancer

**Plain English summary of protocol**Not provided at time of registration

# Contact information

Type(s)

Scientific

Contact name

Dr - -

#### Contact details

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# Additional identifiers

ClinicalTrials.gov (NCT) NCT00003209

Protocol serial number CE3004

# Study information

#### Scientific Title

#### **Study objectives**

Not provided at time of registration

#### 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

Cervix

#### **Interventions**

- 1. Arm A: Chemotherapy with single agent cisplatinum repeated every 21 days for a maximum of six courses.
- 2. Arm B: Multi-drug chemotherapy with bleomycin, ifosfamide and cisplatinum (BIP) repeated every 21 days for a maximum of six courses.

#### Intervention Type

Other

#### Phase

**Not Specified** 

#### Primary outcome(s)

Not provided at time of registration

#### Key secondary outcome(s))

Not provided at time of registration

#### Completion date

28/02/1996

# **Eligibility**

#### Key inclusion criteria

- 1. Histologically proven recurrent invasive squamous cell carcinoma of the cervix uteri
- 2. Symptomatic inoperable pelvic or metastatic disease not amenable to local radiotherapy

- 3. No previous chemotherapy with any of the study agents
- 4. World Health Organisation (WHO) performance status >2
- 5. Adequate renal hepatic and haematological function
- 6. Adequate pulmonary function
- 7. Expected survival of >3 months
- 8. No second primary tumour other than basal cell carcinoma of the skin
- 9. No other serious medical or psychological condition precluding treatment

#### Participant type(s)

**Patient** 

#### Healthy volunteers allowed

No

#### Age group

**Not Specified** 

#### Sex

Female

#### Key exclusion criteria

Not provided at time of registration

#### Date of first enrolment

01/01/1990

#### Date of final enrolment

28/02/1996

## Locations

#### Countries of recruitment

United Kingdom

England

# Study participating centre UKCCCR Register Co-ordinator

London United Kingdom NW1 2DA

# Sponsor information

#### Organisation

Cancer Research UK (CRUK) (UK)

#### **ROR**

https://ror.org/054225q67

# Funder(s)

#### Funder type

Charity

#### **Funder Name**

Cancer Research UK

#### Alternative Name(s)

CR\_UK, Cancer Research UK - London, Cancer Research UK (CRUK), CRUK

#### **Funding Body Type**

Private sector organisation

#### **Funding Body Subtype**

Other non-profit organizations

#### Location

United Kingdom

# **Results and Publications**

Individual participant data (IPD) sharing plan

### IPD sharing plan summary

#### **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article	results	01/09/2000		Yes	No