

# Study to see if GSK2110183 with chemotherapy can treat ovarian cancer

<b>Submission date</b> 16/01/2014	<b>Recruitment status</b> No longer recruiting	<input type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
<b>Registration date</b> 16/01/2014	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan <input checked="" type="checkbox"/> Results
<b>Last Edited</b> 14/04/2021	<b>Condition category</b> Cancer	<input type="checkbox"/> Individual participant data

## Plain English summary of protocol

<http://www.cancerresearchuk.org/cancer-help/trials/a-study-looking-cell-changes-ovarian-cancer-aktres>

## Contact information

### Type(s)

Scientific

### Contact name

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## Additional identifiers

### EudraCT/CTIS number

2012-002483-27

### IRAS number

### ClinicalTrials.gov number

NCT01653912

### Secondary identifying numbers

13614, PKB116611

## Study information

### Scientific Title

AKTRES study: A Biologic Study of the early effects and determinants of AKT inhibition using GSK2110183 alongside chemotherapy in patients with platinum RESistant adenocarcinoma of the ovary

### Acronym

AKTRES

### Study objectives

Activation of the AKT signalling pathway in ovarian cancer cells causes chemotherapy resistance. An AKT inhibitor, given alongside chemotherapy, could potentially reverse this resistance and enable chemotherapy to be effective once again. This study is designed for women whose ovarian cancer has relapsed within 6 months of receiving carboplatin or cisplatin-containing chemotherapy and who are receiving an AKT inhibitor given alongside 3 weekly carboplatin and paclitaxel in the PKB116611 trial. Participants in the AKTRES study undergo a maximum of two biopsies from their tumour and some additional blood tests. The aims of the study are to better understand the how chemotherapy-resistant cancer cells are affected by AKT inhibitors.

### Ethics approval required

Old ethics approval format

### Ethics approval(s)

London Hampstead, 07/11/2012, ref. 12/LO/1174

### Study design

Non-randomised; Interventional; Design type: Process of Care, Treatment

### Primary study design

Interventional

### Secondary study design

Non randomised study

### Study setting(s)

Hospital

### Study type(s)

Treatment

### Participant information sheet

Not available in web format, please use the contact details below to request a patient information sheet

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

Topic: National Cancer Research Network; Subtopic: Gynaecological Cancer; Disease: Ovary /Fallopian tube

## Interventions

Blood sampling, [3 x 10 ml whole blood collected in greentopped (heparinised) tube] will be collected at baseline (within 14 days prior to first dose), on day 1 (immediately preceding chemotherapy), day 8 and day 15 of cycle 1 only. Blood sampling will subsequently be performed at day 1 only of subsequent chemotherapy treatment cycles and thereafter 3 weekly for those on maintenance GSK2110183 alone. A final 3 x 10 ml blood sample will be taken at end of study visit.

Coagulation profile, will be collected immediately prior to the first biopsy during the screening period and postcycle 1 biopsy on day 22. This is in addition to the screening coagulation blood test required for PKB116611 study.

Collection of ascites: If patients are undergoing ascitic or pleural drainage for symptom relief prior to, or during, this study a sample of ascites (or pleural effusion) can be taken and stored. The preferred times for collection are during screening, day 22 and at study completion, but additional/alternative collections can be obtained at the investigators discretion.

Tumour Biopsy: A tumour biopsy performed under image guidance within 14 days prior to first dose. Three tumour cores will be taken, two will be fresh frozen and one will be formalin fixed. Following one complete treatment cycle (day 22), =3 days prior to starting cycle 2 day 1, a second tumour biopsy will be performed under image guidance. Again, three tumour cores will be taken, two will be fresh frozen and one will be formalin fixed. Patients can decline the second biopsy if they wish to.

## Intervention Type

Drug

## Phase

Phase I/II

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

GSK2110183 (afurasertib)

## Primary outcome measure

Overall Response, CA125 response and Progression free survival measured by ELISA detected decrease in phosphorylated PRAS40 (as ratio of total PRAS40) and increase in pAKT S473

## Secondary outcome measures

Not provided at time of registration

## Overall study start date

10/12/2012

## Completion date

04/12/2014

## Eligibility

### Key inclusion criteria

1. Female at least 18 years of age at the time of signing the informed consent form
2. Capable of giving written informed consent
3. Having measurable tumour tissue and Platinum-resistant ovarian cancer as defined as radiological evidence of disease progression within 6 months of completion of platinum-

containing chemotherapy

4. Tumour tissue measuring >2cm long axis amenable to direct biopsy or biopsy via image-guidance

**Participant type(s)**

Patient

**Age group**

Adult

**Lower age limit**

18 Years

**Sex**

Female

**Target number of participants**

Planned Sample Size: 24; UK Sample Size: 12

**Total final enrolment**

59

**Key exclusion criteria**

Concurrent medication with warfarin or low molecular weight heparin (heparin use is acceptable if it has been discontinued for 2 days preceding biopsy)

**Date of first enrolment**

10/12/2012

**Date of final enrolment**

04/12/2014

## **Locations**

**Countries of recruitment**

England

United Kingdom

**Study participating centre**

Hammersmith Hospital

London

United Kingdom

W12 0HS

## **Sponsor information**

**Organisation**

Imperial College of Science, Technology and Medicine (UK)

**Sponsor details**

Department of Cancer Medicine  
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**Sponsor type**

University/education

**ROR**

<https://ror.org/041kmwe10>

**Funder(s)****Funder type**

Charity

**Funder Name**

Ovarian Cancer Action (UK)

**Alternative Name(s)****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?
<a href="#">Basic results</a>				No	No
<a href="#">Basic results</a>				No	No
<a href="#">Results article</a>		01/03/2019	14/04/2021	Yes	No