A phase II uncontrolled study of BAY 73-4506 in previously untreated patients with metastatic or unresectable renal cell cancer (RCC)

Submission date	Recruitment status	Prospectively registered
12/06/2008	No longer recruiting	☐ Protocol
Registration date	Overall study status	Statistical analysis plan
31/07/2008	Completed	[X] Results
Last Edited	Condition category	[] Individual participant data
22/03/2016	Cancer	

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s)

Scientific

Contact name

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Additional identifiers

ClinicalTrials.gov (NCT) NCT00664326

Protocol serial number 11726

Study information

Scientific Title

A phase II uncontrolled study of BAY 73-4506 in previously untreated patients with metastatic or unresectable renal cell cancer (RCC)

Acronym

DAST

Study objectives

Primary hypothesis:

Response rate of patients with advanced renal cell cancer (RCC) to BAY 73-4506.

Secondary hypothesis:

The evaluation of pharmacokinetic and biomarker data.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Brighton East Research Ethics Committee, 06/06/2008, ref: 08/H1107/58

Study design

Interventional single-treatment open-label phase II trial

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Unresectable and/or metastatic renal cell cancer

Interventions

Patients will be treated with BAY 73-4506 160 mg orally (PO) once daily (OD) for three weeks of every four week cycle (i.e. three weeks on, one week off). Patients will continue treatment with BAY 73-4506 until disease progression, intolerable toxicity or patient refusal to continue with the study or at the investigator's decision to remove the patient from the study.

Intervention Type

Drug

Phase

Phase II

Drug/device/biological/vaccine name(s)

BAY 73-4506

Primary outcome(s)

Response rate of patients with advanced RCC to BAY 73-4506.

Key secondary outcome(s))

- 1. Survival
- 2. Progression-free survival
- 3. Time to progression
- 4. Safety (adverse events [AEs], vitals, labs, electrocardiogram [ECG])
- 5. Duration of stable disease
- 6. Pharmacokinetic
- 7. Pharmacodynamic
- 8. Duration of response

Completion date

30/11/2011

Eligibility

Key inclusion criteria

- 1. Greater than or equal to 18 years, either sex
- 2. Unresectable and/or metastatic clear cell renal cell cancer
- 3. Previously untreated disease
- 4. Measurable lesion(s) by computed tomography (CT) scan/magnetic resonance imaging (MRI)
- 5. Intermediate or low Motzer score
- 6. Eastern Cooperative Oncology Group (ECOG) performance status of 0 or 1
- 7. Adequate bone marrow, renal and hepatic function as assessed by specific laboratory tests
- 8. Life expectancy of at least 12 weeks
- 9. Signed informed consent

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Sex

All

Key exclusion criteria

- 1. Previous/concurrent cancer
- 2. Previous systemic treatment of RCC
- 3. Cardiac arrhythmias requiring anti-arrythmics
- 4. History of cardiac disease or congestive heart failure greater than New York Heart Association (NYHA) class 2
- 5. Uncontrolled hypertension despite optimal medical management
- 6. Cardiac ventricular arrhythmias requiring anti-arrhythimics
- 7. Active clinically serious infections

- 8. History of human immunodeficiency virus (HIV) infection or chronic hepatitis B or C
- 9. Know history of symptomatic metastatic brain or meningeal tumours
- 10. Seizure disorders requiring medication
- 11. History of organ allograft
- 12. History or evidence of bleeding diathesis
- 13. Serious non-healing wound, ulcer or bone fracture
- 14. Patients undergoing renal dialysis
- 15. Substance abuse, medical, psychological or social conditions that may interfere with the patient's participation in the study or evaluation of the study results
- 16. Known or suspected allergy to the investigational drug or any agent given in association with the trial
- 17. Any condition which is unstable or which could jeopardise the safety of the patient and his /her compliance in the study
- 18. Pregnant or breast-feeding patients
- 19. Investigational drug therapy outside of the trial within 4 weeks of study entry
- 20. Prior exposure to the drug
- 21. Radiotherapy during study or within 3 weeks of start of study drug
- 22. Major surgery, open biopsy or significant traumatic injury within 4 weeks of start of study
- 23. Autologous bone marrow transplant or stem cell rescue within 4 months of study
- 24. Patients unable to swallow oral medications
- 25. Any malabsorption condition

Date of first enrolment 30/04/2008

Date of final enrolment 30/11/2011

Locations

Countries of recruitment United Kingdom

England

Finland

France

Germany

Poland

United States of America

Study participating centre

Mount Vernon Hospital

Northwood United Kingdom HA6 2RN

Sponsor information

Organisation

Bayer Healthcare Pharmaceuticals Inc. (USA)

ROR

https://ror.org/04hmn8g73

Funder(s)

Funder type

Industry

Funder Name

Bayer Corporation

Alternative Name(s)

Funding Body Type

Private sector organisation

Funding Body Subtype

For-profit companies (industry)

Location

United States of America

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Not provided at time of registration

Study outputs

Output type

Details

Results article	results	01/10/2012	Yes	No
Basic results	Participant information sheet		No	No
Participant information sheet		11/11/2025	11/11/2025 No	Yes