A phase I - II, prospective, double blind, randomised study of the safety and efficacy of sulfasalazine for the treatment of progressing malignant gliomas

Submission date 13/12/2005	Recruitment status No longer recruiting	 Prospectively registered [X] Protocol
Registration date 15/12/2005	Overall study status Completed	 [] Statistical analysis plan [X] Results
Last Edited 15/01/2010	Condition category Cancer	[] Individual participant data

Plain English summary of protocol Not provided at time of registration

Contact information

Type(s) Scientific

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Additional identifiers

EudraCT/CTIS number 2004-004392-11

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers Ulg GBM 04/1

Study information

Scientific Title

Study objectives

Recent evidence suggests that the transcription factor NF-kappaB is constitutively expressed in malignant gliomas and that its inhibition by drugs like sulfasalazine may block the growth of astrocytic tumours in vitro and in experimental models of malignant gliomas. The aim of this study is to evaluate the safety and efficacy of sulfasalazine as a treatment for progressive or recurring malignant gliomas.

Ethics approval required

Old ethics approval format

Ethics approval(s)

The protocol was reviewed and approved by the Ethics Committee of the Faculty of Medicine of the University of Liège (IRB file number 2004/185). It also underwent review and approval by Belgian Federal Authorities (authorisation reference 548/03/05) and was granted the European Trial database (EudraCT) number 2004-004392-11.

Study design A phase I - II, single centre, prospective, randomised, double blind clinical study

Primary study design Interventional

Interventional

Secondary study design Randomised controlled trial

Study setting(s) Not specified

Study type(s) Treatment

Participant information sheet

Health condition(s) or problem(s) studied

Progressive malignant glioma (astrocytic)

Interventions

The randomisation of patients between groups is provided by the Department of Statistics of the Faculty of Medicine of the University of Liège and the drug dosage is communicated only to the hospital pharmacist who prepares and delivers the drug. In order to facilitate the interim

analysis of the data by the review committee, the randomisation algorithm was weighted so that 8 of the first 10 patients receive either the lowest or the highest drug dosage. Neither the investigators nor the patients are aware of the drug dosage that they provide/receive.

Sulfasalazine will be given orally three times a day at total doses of 1.5, 3, 4.5 or 6 g. Four capsules of the drug are to be taken with each meal. Sulfasalazine is to be taken continuously until radiological evidence of tumour progression, complete remission, or the development of serious or intolerable adverse effects. The patient may at any moment decide to discontinue his participation to the study, although every effort will be made to be able to carry on the follow-up. Finally, the independent review committee may decide at any moment to end the study based on safety issues.

Intervention Type

Drug

Phase Phase I/II

Drug/device/biological/vaccine name(s)

Sulfasalazine

Primary outcome measure

 The maximal daily oral dose of sulfaslazine that is tolerated by patients with recurrent or progressive malignant gliomas. Measurements will include the nature, frequency, possible causality and severity of adverse events that occur during treatment
 The assessment of any clinical and/or radiological response of individual tumours to sulfasalazine

Secondary outcome measures

Overall and progression free survival following the initiation of sulfasalazine treatment.

Overall study start date 15/06/2005

Completion date 15/06/2007

Eligibility

Key inclusion criteria

1. Adult patients aged greater than 18 years

2. With recurrent or progressive World Health Organization (WHO) grade 3 or 4 astrocytic gliomas after surgery

- 3. Standard radiation therapy
- 4. A first line of conventional chemotherapy (e.g. temozolomide, CCNU or BCNU)

Recurrence or progression prior to inclusion are based on MacDonalds criteria. Patients are thoroughly informed about the nature of their disease, suspected prognosis, study background and objectives and potential alternative treatments. This information is provided both orally and in written form, prior to obtaining written informed consent from the patient.

Participant type(s)

Patient

Age group

Adult

Lower age limit

18 Years

Sex

Both

Target number of participants

20

Key exclusion criteria

- 1. Present with anaplastic oligodendroglioma (WHO grade 3)
- 2. Allergy to sulfa drugs
- 3. Porphyria
- 4. G-6-PD deficiency
- 5. Psychiatric disorder deemed incompatible with compliance to the study
- 6. Creatinine greater than 15 mg/l
- 7. Aspartate aminotransferase (TGO) greater than 200 UI/l
- 8. Amylase greater than 150 UI/l
- 9. Pregnant or lactating women

10. Patients may not have received any other experimental medication within 30 days (and at least five drug half-lives) prior to inclusion

11. Patients cannot concomitantly take mercaptopurine

Date of first enrolment

15/06/2005

Date of final enrolment 15/06/2007

Locations

Countries of recruitment Belgium

Study participating centre University Hospital of Liege Belgium 4000

Sponsor information

Organisation University of Liege, Department of Neurosurgery (Belgium)

Sponsor details Domaine Universitaire du Sart Tilman, B34 Liege Belgium 4000

Sponsor type University/education

ROR https://ror.org/00afp2z80

Funder(s)

Funder type Research organisation

Funder Name Leon Frederic Fund (Belgium)

Funder Name National Fund for Scientific Research (Belgium)

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?

Protocol article	Protocol	31/01/2006	Yes	No
<u>Results article</u>	results	19/10/2009	Yes	No