Combined Local Immunotherapy and Radiotherapy in metastatic Melanoma

Submission date 17/09/2009	Recruitment status No longer recruiting	[X] Prospectively registered [_] Protocol
Registration date 19/11/2009	Overall study status Completed	
Last Edited 12/12/2017	Condition category Cancer	 Individual participant data Record updated in last year

Plain English summary of protocol

Not provided at time of registration

Contact information

Type(s) Scientific

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Contact details

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Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers N/A

Study information

Scientific Title

Combined Local Immunotherapy and Radiotherapy in metastatic Melanoma: an interventional single-arm trial

Acronym

CLIRM-1

Study objectives

Malignant melanoma lesions can be cleared using a combination of local immunotherapy (Tolllike receptor [TLR] agonists and interleukin-2 [IL-2]) and radiotherapy.

Ethics approval required

Old ethics approval format

Ethics approval(s) Wandsworth Research Ethics Committee, approval pending as of 17/09/2009

Study design Interventional single-arm single-centre trial

Primary study design Interventional

Secondary study design Single-centre

Study setting(s) Hospital

Study type(s) Treatment

Participant information sheet Please contact Dr Janet Fernihough at janet@fischertrust.org for patient information sheet

Health condition(s) or problem(s) studied

Metastatic melanoma

Interventions

All participants will receive the following treatments:

1. Aldara® cream applied four times weekly. Each sachet of 250 mg is sufficient to cover 20 square centimeters, so amount of cream used depends on the size of the lesion.

2. Intralesional injections of Proleukin® and Hiltonol®:

2.1. Proleukin® will be diluted to a concentration of 10 micrograms interleukin-2 per millilitre (10 ugml). Each lesion will have 100 microlitre injections, up to three in number depending on the size of the lesion. This will be performed three times weekly.

2.2. Hiltonol® is a 2 mg/ml solution and again, 100 microlitre injections will be made

intralesionally, with a maximum of three injections per lesion, three times weekly. 3. Local radiotherapy

The total duration will be one year if the treatment is effective, but we anticipate an average of 36 weeks, depending on the choice of radiotherapy schedule.

Intervention Type

Drug

Phase Not Applicable

Drug/device/biological/vaccine name(s)

Imiquimod (Aldara®), aldesleukin (Proleukin®), poly-ICLC (Hiltonol®)

Primary outcome measure

Size of lesion

All primary and secondary outcome measures will be measured formally at entry, after 1 month, 6 months and end of trial.

Secondary outcome measures

1. Toxicity associated with this treatment will be measured by the number of adverse and serious adverse events during treatment, and the number of injections delayed in case of excessive local reaction.

2. Frequency and latency of appearance of any new skin lesions local or distal to the treated area will be recorded

3. Levels of markers associated with tumour infiltrating lymphocytes and tumour associated macrophages will be estimated by real time polymerase chain reaction (RT-PCR) of material biopsied from the lesions

4. Systemic immune responses associated with this treatment will be assessed by analysis of serum levels of cytokines and white blood cell properties taken by blood sampling during treatment.

All primary and secondary outcome measures will be measured formally at entry, after 1 month, 6 months and end of trial.

Overall study start date

01/01/2010

Completion date

31/12/2012

Eligibility

Key inclusion criteria

1. Both males and females, aged 16 to 85 years

2. Patients who have one or more malignant skin tumours confirmed histologically at diagnosis

3. Patients with malignant skin tumours for whom other standard therapy options are no longer appropriate or have been refused by the patient

4. Life expectancy of at least 3-6 months

5. Patients with a WHO performance status of 0, 1 or 2

6. Patients who are informed of and are willing and able to comply with the home application of the Aldara® cream

7. Patients who are willing and able to comply with the investigational nature of the study and who have signed a written informed consent form

8. Patients who are willing to receive the number of intra-lesional injections and biopsies required to complete the study

Participant type(s)

Patient

Age group

Adult

Sex

Both

Target number of participants

30

Key exclusion criteria

1. Patients with unstable or severe current medical conditions or active, uncontrolled infection 2. Patients with psychological or sociological conditions, addictive disorders or family problems which would preclude compliance with the protocol

3. Patients undergoing therapy at study entry with other investigational agents that are directly immunosuppressive

4. Patients having procreative potential who are not using adequate contraception

5. Patients with untreated/uncontrolled brain tumours

6. Patients with brain tumours which have been treated but which have not been stable for 3 or more months

7. Patients with known hypersensitivity to Hiltonol®, IL-2, Aldara®, cyclophosphamide (for patients 16 30) or any of the excipients

8. Any condition, which, in the opinion of the investigator might interfere with the safety of the patient or evaluation of the study objectives

Date of first enrolment

01/01/2010

Date of final enrolment 31/12/2012

Locations

Countries of recruitment England

United Kingdom

Study participating centre

St George's University London London United Kingdom SW17 0RE

Sponsor information

Organisation St George's Healthcare NHS Trust (UK)

Sponsor details St George's Hospital London Blackshaw Road Tooting London England United Kingdom SW17 0QT

Sponsor type Hospital/treatment centre

Website http://www.stgeorges.nhs.uk/

ROR https://ror.org/039zedc16

Funder(s)

Funder type Charity

Funder Name Application to Cancer Vaccine Institute (UK) in progress as of 17/09/2009.

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