WT1 TCR gene therapy for leukaemia: a phase I /II safety and toxicity study (WT1 TCR-001)

Submission date	Recruitment status No longer recruiting	Prospectively registered		
31/03/2010		[_] Protocol		
Registration date	Overall study status Completed	[] Statistical analysis plan		
31/03/2010		[X] Results		
Last Edited 21/04/2020	Condition category Cancer	[] Individual participant data		

Plain English summary of protocol

http://www.cancerresearchuk.org/about-cancer/find-a-clinical-trial/a-trial-looking-at-a-type-of-gene-therapy-for-acute-myeloid-leukaemia-and-chronic-myeloid-leukaemia

Contact information

Type(s) Scientific

Contact name Dr Emma Morris

Contact details

Royal Free Hospital Pond Street London United Kingdom NW3 2QG

Additional identifiers

EudraCT/CTIS number 2006-004950-25

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers 5099

Study information

Scientific Title

A phase I/II safety and toxicity study on the use of WT1 TCR gene therapy for adult patients with acute and chronic myeloid leukaemia

Acronym

WT1 TCR-001

Study objectives

One of the main functions of the immune system is to protect the body from infection. It is now clear that the immune system also plays a role in preventing the development or controlling the growth of some cancers. The most important cells of the immune system for this particular task are a group of white blood cells called the T lymphocytes (T cells). It is known from bone marrow transplantation and T cell infusions that leukaemia can be controlled or cured by a strong T cell immune response. Despite this, many other patients develop leukaemia even with a normal immune system. This study wants to test a new way of strengthening the patient's own immune response to their leukaemia by increasing the number of patient T cells which can recognise and kill the leukaemia cells.

The T cells have a receptor (the T cell receptor [TCR]), which enables them to recognise particular abnormalities on the surface of the target cells (virally infected cells or cancer /leukaemia cells). Each TCR recognises a particular fragment of a protein (peptide epitope). Normally, patients have a wide range of different T cells, which recognise many different epitopes on the surface of 'target cells'. It is possible that they have only a very few, or no T cells, which are able to recognise the abnormal leukaemia cells.

The Wilms' tumour antigen 1 (WT1) is a protein, peptides of which are present at abnormally high levels on the surface of leukaemia cells. In the research laboratory we have identified T cells, which specifically kill leukaemia cells by recognising the WT1 on their cell surface. The TCR determines the specificity of the T cell. Not all patients have T cells with the WT1-specific TCR.

We can generate T cells, which recognise WT1 (and can therefore kill leukaemia cells) by transferring the genes for the WT1-specific TCR into T cells, which normally recognise something else.

Ethics approval required Old ethics approval format

Ethics approval(s) Gene Therapy Advisory Committee, 20/12/2007, ref: GTAC 128

Study design Non-randomised interventional multicentre 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: Haematological Oncology; Disease: Leukaemia (chronic), Leukaemia (acute myeloid)

Interventions

Leucapheresis, After recruitment to the study, patients will undergo leucapheresis to harvest peripheral blood lymphocytes. The peripheral blood T-lymphocytes will be cultured for up to 9 days in vitro for transduction with replication defective retroviral vectors containing the WT1-specific TCR. Bulk transduced T-lymphocytes will be intravenously administered using escalating doses of = 2 x 10^7/kg and = 10^8/kg bulk TCR-td T cells (doses based on numbers of allogeneic T cells safely infused post Allo SCT and t).

Follow Up Length: 12 month(s)

Intervention Type

Genetic

Primary outcome measure

- 1. Transduction efficiency at QP release of genetically modified (transduced) T cells
- 2. Toxicity and Side effects every trial visit
- 3. Integration site analysis of transduced T cells batched and to be performed at end of study

Secondary outcome measures

1. Persistence of TCR-Td T cells - blood sample taken at every trial visit after infusion of T cells (analysis to be performed in batches)

2. WT1-specific Immune Responses - blood sample taken at every trial visit after infusion of T cells (analysis to be performed in batches)

3. Disease responses - BM aspirate and trephine at +8 weeks; qPCR at 7 days, 28 days then monthly until 12 months post T cell infusion

Overall study start date

18/01/2010

Completion date 17/01/2012

Eligibility

Key inclusion criteria

General inclusion criteria:

All patients will undergo detailed laboratory based assessment prior to the procedure:

1. Aged greater than or equal to 18 years and less than or equal to 75 years

2. Life expectancy greater than 1626 weeks (46 months)

3. World Health Organisation (WHO) performance status of 0 - 2

4. HLA A*0201 positive

5. Completed previous course of chemotherapy greater than or equal to 4 weeks prior to commencing the initial phase of the trial (leucapheresis for collection of patient peripheral blood mononuclear cells [PBMC])

7. Peripheral blood total lymphocyte count greater than 0.5 x 10^9/L

8. Informed consent in writing and ability to co-operate with treatment and follow up

9. Willing, able and available for collection of PBMC/T-cells by leucapheresis

10. Hepatitis B and C, HTLV-1, human immunodeficiency virus (HIV) negative

11. Free from serious concurrent illness

12. Female patients of child-bearing age must have a negative pregnancy test and agree to use reliable contraceptive methods for the duration of the therapy and for 6 months afterwards13. Male patients must agree to use appropriate medically approved contraception during the trial and for six months afterwards

14. Haematological and Biochemical Indices:

14.1. Haemoglobin (Hb) = 7.0 g/dl; neutrophils = 0.2 x 10^9/L; total lymphocytes >0.5 x 10^9/L; platelets (Plts) = 40 x 10^9/L

14.2. Serum bilirubin, Alanine amino-transferase (ALT) and/or aspartate amino-transferase (AST) less than 3 x upper normal limit

14.3. Calculated creatinine clearance = 30 ml/min (uncorrected value) or isotope clearance measurement = 30 ml/min

Disease-specific inclusion criteria:

AML or CML proven by morphology, histology, immunophenotyping and cytogenetics (where available):

1. Acute myeloid leukaemia (AML):

Patients not eligible for BMT procedure with:

1.1. AML in 2nd CR or greater

1.2. Good and Standard Risk* AML in 1st CR or PR in patients greater than 60 years; and

1.3. Poor Risk* AML in 1st CR CR/PR or later (slow remitters and/or adverse cytogenetics)

1.4. AML at 1st relapse post BMT in CR or PR after re-induction and consolidation.

[NB, *risk category as defined by MRC criteria: Good Risk: t(15;17), t(8;21), inv 16; Poor Risk: -5; -7; del (5q); abn (3q) or complex (=/>4 abn)].

2. Chronic myeloid leukaemia (CML):

2.1. Patients in chronic phase resistant to Glivec/Imatinib and 2nd generation tyrosine kinase inhibitors (e.g., Dasatinib), AND NOT eligible for allogeneic BMT

2.2. Patients in chronic phase resistant to Glivec/Imatinib and with an identified mutation known to be resistant to 2nd generation tyrosine kinase inhibitors, AND NOT eligible for allogeneic BMT 2.3. Patients = 50 years (and ineligible for myeloablative allo-BMT), with a suboptimal response to Glivec/Imatinib and an identified mutation known to be resistant to 2nd generation tyrosine kinase inhibitors. These patients are at high risk of disease progression. Patients in this group would stop Glivec/Imatinib prior to leucapheresis and receiving TCR-transduced T cells. They will have monthly quantitative RT-PCR for Bcr-Abl and restart Glivec/Imatinib in the event of a log increase in transcript numbers.

2.4. Patients in chronic phase, resistant to Glive/Imatininb and NOT eligible for allo BMT without access to 2nd generation tyrosine kinase inhibitors may be considered after discussion with the Chief Investigator and Sponsor

Resistance to Glivec is defined as (European Leukemianet Criteria, 2006): No Haematological Response (HR) at 3 months Incomplete HR or No Cytogenetic Response (CgR) at 6 months Less than partial CgR (Ph+ >35%) at 12 months Less than complete CgR at 18 months Loss of HR or CgR Development of highly resistant mutations

Suboptimal response to Glivec is defined as (European Leukemianet Criteria, 2006): Less than complete Haematological response at 3 months Less than partial CgR (Ph+ >35%) at 6 months Less than complete CgR at 12 months Less than major MoR at 18 months Loss of major MoR Development of a mutation

Participant type(s)

Patient

Age group

Adult

Lower age limit 18 Years

Sex

Both

Target number of participants

Planned Sample Size: 18; UK Sample Size: 18

Total final enrolment

7

Key exclusion criteria

1. Aged less than 18 years or greater than 75 years

2. Patients should not receive concurrent systemic corticosteroids whilst on the study

3. Major thoracic and/or abdominal surgery in the preceding three to four weeks from which the patient has not yet recovered

4. Patients who are high medical risks because of non-malignant systemic disease, as well as those with active uncontrolled infection

5. Patients with any other condition, which in the Investigator's opinion would not make the patient a good candidate for the clinical trial

6. Patients known to be serologically positive for Hepatitis B, C, HTLV-1 or HIV

7. Concurrent congestive heart failure or prior history of New York Heart Association (NYHA) class III/IV cardiac disease

8. Positive pregnancy test or reluctance to use contraception

Pregnant and lactating women are excluded
History of severe allergy

Date of first enrolment 18/01/2010

Date of final enrolment 17/01/2012

Locations

Countries of recruitment England

United Kingdom

Study participating centre Royal Free Hospital London United Kingdom NW3 2QG

Sponsor information

Organisation University College London (UCL) (UK)

Sponsor details Gower Street London England United Kingdom WC1E 6BT

Sponsor type University/education

Website http://www.ucl.ac.uk/

ROR https://ror.org/02jx3x895

Funder(s)

Funder type Government

Funder Name Department of Health (UK) (ref: 66807)

Funder Name Leukaemia Research Fund (UK) (ref: 08001)

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?
Basic results			21/04/2020	No	No