Clinical investigation to assess the safety and performance of the Organox Metra, for normothermic perfusion of livers, prior to transplantation and to compare with retrospective data from matched controls

Submission date	Recruitment status No longer recruiting	Prospectively registered		
11/10/2012		Protocol		
Registration date	Overall study status	Statistical analysis plan		
27/11/2012	Completed	[X] Results		
Last Edited	Condition category	[] Individual participant data		
25/05/2018	Surgery			

Plain English summary of protocol

Background and study aims

The OrganOx metra is an automated transportable device for preserving donated livers for transplantation. The aim of this study is to assess the safety and performance of the OrganOx Metra as an organ preservation method before liver transplantation.

Who can participate?

Patients aged 18 or older on the waiting list for liver transplantation

What does the study involve?

Participants receive liver transplants that have been preserved using the OrganOx metra device. The functioning of the transplanted liver and the patient survival rate are compared with the results from livers that were preserved using the conventional ice bucket method.

What are the possible benefits and risks of participating?

This study may demonstrate the safety and performance of the OrganOx metra as a medical device for the preservation of livers before transplantation. It is hoped that there will be benefits in terms of both the quality and quantity of livers available for transplantation. We believe that this would reduce the waiting time for liver transplantation and improve outcomes for patients. Liver transplant surgery carries a risk of significant complications, including bile duct complications, bleeding, blood clots, infection, memory and thinking problems, and failure or rejection of the donated liver. Using the OrganOx Metra does not increase the level of risk to the patient.

Where is the study run from? Kings College Hospital (UK) When is the study starting and how long is it expected to run for? October 2012 to December 2013

Who is funding the study? OrganOx Ltd (UK)

Who is the main contact? Dr Toni Day toni.day@organox.com

Contact information

Type(s)

Scientific

Contact name

Prof Nigel Heaton

Contact details

King's College Hospital Institute of Liver Studies London United Kingdom SE5 9RS

Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers ORG/NPL/001

Study information

Scientific Title

Non-randomised, open-label, single-arm, prospective clinical investigation to assess the safety and performance of the Organox Metra, for normothermic perfusion of livers, prior to transplantation and to compare with retrospective data from matched controls

Study objectives

To assess the safety and performance of normothermic perfusion as an organ preservation method prior to liver transplantation using the OrganOx Metra.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Dulwich Research Ethics Committee, 19/07/2012

Study design

Non-randomised open-label single-arm prospective clinical investigation

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

Liver transplant

Interventions

Treatment arm 1: livers transplanted after preservation on the OrganOx metra device, the methodology for current clinical practice for liver transplantation is observed.

Treatment arm 2: Retrospectively collected data from livers transplanted after preservation using the conventional ice bucket, the methodology for current clinical practice for liver transplantation is observed.

Intervention Type

Procedure/Surgery

Phase

Not Applicable

Primary outcome measure

Graft survival at 30 days

Secondary outcome measures

- 1. Histological post-reperfusion biopsies (H&E and immunohistochemistry) 60 minutes after arterial reperfusion (before abdominal closure)
- 2. Markers of endothelial damage and pro-inflammatory cytokines (TNF, IL-6, vWF) (postoperative)
- 3. Markers of liver function (Bilirubin, Alkaline phosphatase, GGT, AST, creatinine (postoperative)
- 4. Six month magnetic resonance cholangiopancreatography (MRCP) appearances
- 5. Six month graft survival
- 6. Six month patient survival

Overall study start date

11/10/2012

Completion date

01/12/2013

Eligibility

Key inclusion criteria

Organ inclusion criteria

- 1. Livers from male or female adult donors (18 years or greater)
- 2. Accepted for transplantation using current (standard and extended) criteria
- 3. Both heart-beating (DBD) and non-heart-beating (DCD) donor organs may be included

Recipient Inclusion criteria

- 1. Male or female, aged 18 years or older
- 2. On the waiting list for liver transplantation at Kings College Hospital, London.
- 3. Provided informed consent for participation in the study
- 4. Able and willing to comply with all study requirements (in the opinion of the investigator or nominated deputy)

Participant type(s)

Patient

Age group

Adult

Lower age limit

18 Years

Sex

Both

Target number of participants

20

Key exclusion criteria

Organ exclusion criteria

- 1. Livers offered from a donor hospital that is located more than 3 hours travelling time (by road) of both Kings College Hospital and Oxford.
- 2. Donors in whom the liver is planned to be split.
- 3. Donor livers that would not be accepted under current (Kings College Hospital) criteria.
- 4. Livers taken from donors aged <18 years.
- 5. Donors with high donor serum Na+ (> 160 mmol/litre) at the time of liver retrieval.

Recipient Exclusion criteria

The participant may not enter the study if ANY of the following apply:

- 1. Not willing to, unable or not capable to provide informed consent
- 2. Recipients aged less 18 years

- 3. Undergoing transplant for fulminant hepatic failure
- 4. Undergoing re-transplantation.
- 5. Undergoing transplantation of other organ(s) in addition to the liver
- 6. Any other significant disease or disorder which, in the opinion of the Investigator, may either put the participant at risk by participating in the study, or may influence the result of the study, or the participants ability to participate in the study.
- 7. Not suitable for any other reason, in the opinion of the Investigator, to take part in the study.
- 8. Enrolled in clinical studies of other unlicensed therapy.
- 9. Participated in another research study involving an investigational product in the previous 12 weeks.
- 10. Pregnant or nursing mothers

Date of first enrolment 11/10/2012

Date of final enrolment 01/12/2013

Locations

Countries of recruitment England

United Kingdom

Study participating centre King's College Hospital London United Kingdom SE5 9RS

Sponsor information

Organisation

OrganOx Ltd (UK)

Sponsor details

c/o Dr Toni Day Magdalen Centre Oxford United Kingdom OX5 1TN

Sponsor type

Industry

Website

http://www.organox.com

ROR

https://ror.org/04fqe7a64

Funder(s)

Funder type

Industry

Funder Name

OrganOx Ltd (UK)

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?
Results article	results	01/06/2016		Yes	No