

Assessment of Collaborative Requesting

Submission date 22/03/2007	Recruitment status No longer recruiting	<input type="checkbox"/> Prospectively registered
Registration date 16/07/2007	Overall study status Completed	<input type="checkbox"/> Protocol
Last Edited 12/10/2009	Condition category Other	<input type="checkbox"/> Statistical analysis plan
		<input checked="" type="checkbox"/> Results
		<input type="checkbox"/> 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

Protocol serial number
Protocol v5.2

Study information

Scientific Title

Acronym
ACRE

Study objectives

Relatives of patients in whom death has been determined using brain stem death criteria are more likely to consent to organ donation when interviewed by the clinical team if a donor transplant co-ordinator is present, compared with relatives interviewed by the clinical team alone.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Approved by Oxfordshire Research Ethics Committee A (Ref: 06/Q1604/119)

Primary study design

Interventional

Study design

Randomised controlled trial

Study type(s)

Other

Health condition(s) or problem(s) studied

Relatives of brain stem dead patients; Organ donation - collaborative requesting

Interventions

The intervention is collaborative requesting which is the addition of a donor transplant co-ordinator to the team requesting organ donation. The routine arm is the relatives interviewed by the clinical team alone.

Intervention Type

Other

Phase

Not Specified

Primary outcome(s)

Proportion of relatives refusing consent for organ donation.

Key secondary outcome(s)

1. Number of patients in whom the family consent to beating heart organ donation
2. Solid organs retrieved by type
3. Solid organs transplanted by type
4. Tissues retrieved by type
5. Consent rates by predefined subgroup (relative relationship to donor; ethnic group and sex of donor and next of kin; clinician seniority and sex; sex of donor transplant co-ordinator)

Completion date

01/08/2009

Eligibility

Key inclusion criteria

Relatives of patients in whom death has been determined using brain stem death criteria

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Not Specified

Sex

Not Specified

Key exclusion criteria

1. Relatives who do not understand written or verbal information for whom an interpreter is not available
2. Relatives who decline discussion on organ donation

Date of first enrolment

01/04/2007

Date of final enrolment

01/08/2009

Locations**Countries of recruitment**

United Kingdom

England

Study participating centre

c/o Miss Sarah Edwards

Oxford

United Kingdom

OX3 9DU

Sponsor information**Organisation**

Oxford Radcliffe Hospitals NHS Trust (UK)

ROR

https://ror.org/03h2bh287

Funder(s)

Funder type

Government

Funder Name

UK Transplant

Results and Publications

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	08/10/2009		Yes	No