Real-world evaluation of robot-assisted surgical services

Submission date	Recruitment status No longer recruiting	Prospectively registered		
07/07/2022		[X] Protocol		
Registration date	Overall study status Ongoing Condition category	Statistical analysis plan		
10/10/2022		Results		
Last Edited		[] Individual participant data		
01/08/2025	Surgery	[X] Record updated in last year		

Plain English summary of protocol

Background and study aims

This study aims to work out if robot-assisted surgery (RAS), a relatively new development, should be routinely available in the NHS and also assess any barriers to its implementation. Over the next 20 years, surgery performed with the help of a robot is expected to increase rapidly around the world, especially for cancer conditions. Previous research shows that when RAS has been introduced in some clinical areas, like urology, it can help surgeons be more precise and can reduce a patient's hospital stay. Using RAS may also speed up training for surgeons to enable them to become experts more quickly. However, RAS has not been tested in all clinical areas and is very expensive with each robot costing over £ 1 million). Also, when RAS is introduced into hospitals it requires special consideration as the set-up can be disruptive. It is not yet clear whether the benefits to patients or the health system of doing surgery this way is worth the cost and the disruption. This study aims to answer that question and provide guidelines for the best way of doing it if robotic surgery is shown to be useful.

We have designed the research to be able to measure the impact of RAS as it is introduced in the UK and scaled up in other hospitals currently performing robotic surgery but planning to expand services. It will study the effects of RAS as it is rolled out at 16 different sites in a planned way. We will measure what happens to patients who get RAS as part of the service and compare their outcomes (e.g., complications, recovery time) to conventional surgery. We will also track how introducing RAS impacts on the staff and the surgeons, and how it affects wider care in hospitals across the country.

This study aims to undertake a real-world, large-scale evaluation of the introduction and scaleup of RAS services evaluating its impact on NHS service delivery, clinical effectiveness, budget and cost-effectiveness.

Who can participate?

Any patient undergoing the specified surgical procedure at a participating hospital site

What does the study involve?

The study involves patients completing a questionnaire before and after surgery to evaluate

their experience of the procedure. We will also interview and survey staff involved with REINFORCE at sites, and calculate the total cost to the NHS (e.g., labour, consumables and other items of surgical equipment).

What are the possible benefits and risks of participating?

There is no direct benefit to participants, but the results of the study are likely to benefit future NHS patients undergoing surgery. Outside of the usual risks associated with surgery and anaesthetic, there are no anticipated risks or disadvantages to participating in the REINFORCE study.

Where is the study run from?

Surgical Intervention Trials Unit (SITU), University of Oxford and Centre for Healthcare Randomised Trials (CHaRT), University of Aberdeen (United Kingdom)

When is the study starting and how long is it expected to run for? January 2022 to August 2025

Who is funding the study?
NIHR Health and Social Care Delivery Research (NIHR HSDR) (United Kingdom)

Who is the main contact? REINFORCE@ndorms.ox.ac.uk

Study website

https://w3.abdn.ac.uk/hsru/REINFORCE/

Contact information

Type(s)

Public

Contact name

Dr Emma Blackmore-Bowes

ORCID ID

https://orcid.org/0000-0002-5848-5333

Contact details

NDORMS
University of Oxford
Nuffield Orthopaedic Centre
Oxford
United Kingdom
OX3 7HE
+44 (0)7721 491938
REINFORCE@ndorms.ox.ac.uk

Type(s)

Principal Investigator

Contact name

Prof David Beard

ORCID ID

https://orcid.org/0000-0001-7884-6389

Contact details

NDORMS
University of Oxford
Nuffield Orthopaedic Centre
Oxford
United Kingdom
OX3 7HE
+44 (0)1865 227695
David.Beard@ndorms.ox.ac.uk

Type(s)

Principal Investigator

Contact name

Prof Marion Campbell

ORCID ID

https://orcid.org/0000-0001-5386-4097

Contact details

Health Services Research Unit University of Aberdeen Foresterhill Aberdeen United Kingdom AB25 2ZD +44 (0)1224 273161 m.k.campbell@abdn.ac.uk

Additional identifiers

EudraCT/CTIS number

Nil known

IRAS number

311223

ClinicalTrials.gov number

Nil known

Secondary identifying numbers

CPMS 53026, IRAS 311223

Study information

Scientific Title

A real-world, in-situ, evaluation of the introduction and scale-up of robot-assisted surgical services in the NHS: Evaluating its impact on clinical and service delivery, effectiveness and cost (the REINFORCE study)

Acronym

REINFORCE

Study objectives

This study aims to undertake a real-world, large-scale evaluation of the introduction and scale-up of Robot-Assisted Surgery (RAS) services evaluating its impact on NHS service delivery, clinical effectiveness, budget and cost-effectiveness.

Ethics approval required

Old ethics approval format

Ethics approval(s)

This is an observational study of healthcare professionals. The Sponsor, University of Oxford, reviewed the study as a Service Evaluation project and indicated the project does not need to be submitted for REC review.

Study design

Observational cohort study

Primary study design

Observational

Secondary study design

Cohort 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

Evaluation of robot-assisted surgery

Interventions

- 1. Study design: A stepped-wedge evaluation with integrated process evaluation and economic assessment.
- 2. Study sites: NHS hospitals planning to introduce/expand robot-assisted surgery (RAS) services. Sites will be switched over from non-RAS to RAS-augmented services (or switch up from one level of provision to another) in random order
- 3. Study participants (stepped-wedge evaluation): All patients undergoing the index procedure

(RAS or otherwise) at each site across all time periods of the study

The project is embedded in normal NHS care and it is intended to be non-selective (all patients undergoing surgery for the index procedure are candidates).

4. Participants (process evaluation): Three key personnel including surgeons, theatre staff and service managers will be sampled from 6 REINFORCE sites and invited to interview. Additionally, 3-4 commissioners will also be interviewed and sampled from across the suite of trial sites. The sample size overall will be approximately 40 interviews.

Sample Size: 2,560 procedures

- 5. Objectives:
- 5.1. Impact of RAS system transformation on clinical and service delivery
- 5.2. Budget impact and cost-effectiveness to the NHS of the introduction of RAS at scale
- 5.3. Potential benefits and harms of RAS across and within speciality areas
- 5.4. Mechanisms of change underpinning any change in outcome, including surgeon training

Intervention Type

Other

Primary outcome measure

- 1. Patient level:
- 1.1. Disease-specific quality of life measured using procedure-specific PROM at baseline and 3 months
- 1.2. Overall quality of life measured using the EQ-5D questionnaire at baseline and 3 months
- 1.3. Overall measure of treatment effectiveness/benefit measured using a Patient Questionnaire at baseline and 3 months
- 1.4. Overall measure of complications inc. mortality measured using Clavien-Dindo score at 3 months
- 2. Surgeon/Team level:
- 2.1. Precision/accuracy measured using Surgeon Task Load Index (TLX) on the day of surgery
- 2.2. Visualisation measured using Surgeon Task Load Index (TLX) on the day of surgery
- 3. Organisation level:
- 3.1. Equipment failure measured using a Surgery Form on the day of surgery
- 3.2. Standardisation of operative quality measured using process evaluation interviews pre/peri/post-robot-assisted surgery (RAS) implementation
- 3.3. Overall economic/cost-effectiveness measured using Health Economics review throughout the study
- 4. Population level:
- 4.1 Equity of access measured using Health Economics review throughout the study

Secondary outcome measures

There are no secondary outcome measures

Overall study start date

01/01/2022

Completion date

30/08/2025

Eligibility

Key inclusion criteria

All patients undergoing the index procedure (robot-assisted surgery or otherwise) at each site across all time periods

Participant type(s)

Patient

Age group

Adult

Lower age limit

18 Years

Sex

Both

Target number of participants

Planned Sample Size: 2560; UK Sample Size: 2560

Total final enrolment

2836

Key exclusion criteria

Does not meet the inclusion criteria

Date of first enrolment

01/07/2022

Date of final enrolment

31/05/2025

Locations

Countries of recruitment

England

United Kingdom

Wales

Study participating centre University Hospital of Wales

Cardiff Urology
Dept of Urology
Cardiff & Vale University Health Board
Heath Park
Cardiff
United Kingdom
CF144XW

Study participating centre University Hospital of Wales

Cardiff Colorectal
Dept of Gastroenterology
Cardiff & Vale University Health Board, University Hospital of Wales
Heath Park,
Cardiff
United Kingdom
CF144XW

Study participating centre Ysbyty Gwynedd

Betsi Cadwaladr Gynaecology Dept of Gynaecology Penrhosgarnedd Bangor Gwynedd United Kingdom LL57 2PW

Study participating centre University Hospital of Wales

Cardiff Gynaecology
Dept of Gynaecological Oncology
Cardiff & Vale University Health Board
Heath Park
Cardiff
United Kingdom
CF144XW

Study participating centre Norfolk & Norwich University Hospital

Colney Lane Colney Norwich United Kingdom NR4 7UY

Study participating centre

North Bristol NHS Trust

Southmead Hospital Southmead Road Westbury-on-trym Bristol United Kingdom BS10 5NB

Study participating centre South Tees Hospitals NHS Foundation Trust

James Cook University Hospital Marton Road Middlesbrough United Kingdom TS4 3BW

Study participating centre Swansea Bay University Local Health Board

Tonna Hospital
Tonna Uchaf
Tonna
Neath
United Kingdom
SA11 3LX

Study participating centre Stockport NHS Foundation Trust

Stepping Hill Hospital Poplar Grove Stockport United Kingdom SK2 7JE

Study participating centre Salford Royal Hospital

Stott Lane Eccles Salford United Kingdom M6 8HD

Study participating centre Salisbury District Hospital

Salisbury District Hospital Odstock Road Salisbury United Kingdom SP2 8BJ

Study participating centre Wrightington Hospital NHS Trust

Hall Lane Wrightington Wigan United Kingdom WN6 9EP

Study participating centre

Cwm Taf Morgannwg University Local Health Board

Dewi Sant Hospital Albert Road Pontypridd United Kingdom CF37 1LB

Study participating centre

Oxford University Hospitals NHS Foundation Trust

John Radcliffe Hospital Headley Way Headington Oxford United Kingdom OX3 9DU

Study participating centre Barts Health NHS Trust

The Royal London Hospital 80 Newark Street London United Kingdom E1 2ES

Sponsor information

Organisation

University of Oxford

Sponsor details

University Offices Oxford England United Kingdom OX1 2JD +44 (0)1865 270000 ctrg@admin.ox.ac.uk

Sponsor type

University/education

Website

https://www.ox.ac.uk/

ROR

https://ror.org/052gg0110

Funder(s)

Funder type

Government

Funder Name

National Institute for Health and Care Research Evaluation, Trials and Studies Co-ordinating Centre (NETSCC); Grant Codes: NIHR131537

Alternative Name(s)

National Institute for Health Research, NIHR Research, NIHRresearch, NIHR - National Institute for Health Research, NIHR (The National Institute for Health and Care Research), NIHR

Funding Body Type

Government organisation

Funding Body Subtype

National government

Location

United Kingdom

Results and Publications

Publication and dissemination plan

Planned publication in a high-impact peer-reviewed journal

Intention to publish date

31/03/2026

Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study will be available upon request from the Surgical Intervention Trials Unit (SITU) at situ@ndorms.ox.ac.uk. Further details will be made available at a later date.

IPD sharing plan summary

Available on request

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol file	version 3.0	20/03/2025	30/05/2025	No	No