# Wearable technology in post breast surgery upper limb rehabilitation

Submission date	Recruitment status	<ul><li>Prospectively registered</li></ul>
27/03/2025	No longer recruiting	☐ Protocol
Registration date	Overall study status	Statistical analysis plan
12/04/2025	Completed	Results
Last Edited	Condition category	Individual participant data
05/09/2025	Musculoskeletal Diseases	[X] Record updated in last year

#### Plain English summary of protocol

Background and study aims

Breast cancer is the most common cancer worldwide, with over 55,000 patients diagnosed in the UK each year. Axillary surgery, which includes procedures like Sentinel Lymph Node Biopsy (SNLB) and Axillary Node Dissection (AND), is crucial for staging breast cancer but can lead to shoulder problems such as pain, reduced movement, lymphoedema, and decreased quality of life. This study aims to explore the benefits of a digital rehabilitation system called OnTrack, which uses a smartwatch and smartphone to help breast cancer patients recover arm function after surgery.

#### Who can participate?

Adults aged 18 and older who are scheduled for sentinel node biopsy or axillary node clearance as part of their breast cancer treatment can participate. Participants must be able to provide informed consent.

#### What does the study involve?

Participants will use the OnTrack system, which provides real-time activity feedback and remote physiotherapy support through a smartwatch and smartphone. This system is designed to help manage arm rehabilitation effectively.

What are the possible benefits and risks of participating?

Benefits include reduced pain and lymphoedema, improved shoulder mobility, and enhanced quality of life. The study also aims to generate cost savings for patients. Risks may include mild discomfort from using the wearable devices, but these are expected to be minimal.

Where is the study run from? Imperial College London (UK)

When is the study starting and how long is it expected to run for? November 2024 to August 2025 Who is funding the study? Imperial MedTechOne (UK) Wellcome Trust (UK)

Who is the main contact? Ahmed Latif, ahmed.latif@imperial.ac.uk Daniel Leff, d.leff@imperial.ac.uk

# Contact information

#### Type(s)

Public, Scientific

#### Contact name

Mr Ahmed Latif

#### **ORCID ID**

https://orcid.org/0009-0001-9066-813X

#### Contact details

10th Floor QEQM Building St Mary's Hospital London United Kingdom W2 1NY +44 7903292540 ahmed.latif@imperial.ac.uk

### Type(s)

Principal investigator

#### Contact name

Mr Daniel Leff

#### Contact details

Breast Surgery Unit, Charing Cross Hospital London United Kingdom W6 8RF +44 7340102203 d.leff@imperial.ac.uk

# Additional identifiers

### Clinical Trials Information System (CTIS)

Nil known

## Integrated Research Application System (IRAS)

327556

#### ClinicalTrials.gov (NCT)

Nil known

#### Protocol serial number

23CX8516

# Study information

#### Scientific Title

A study to investigate the use of wearable technology in upper limb rehabilitation post breast and axillary surgery

#### Study objectives

Wearable technology can facilitate remote shoulder rehabilitation post breast and axillary surgery.

#### Ethics approval required

Ethics approval required

#### Ethics approval(s)

approved 29/02/2024, London - Surrey Research Ethics Committee (2 Redman Place, Stratford, London, E20 1JQ, United Kingdom; +44 207 104 8131; surrey.rec@hra.nhs.uk), ref: 24/PR/0049

#### Study design

Randomized prospective parallel group non-blinded trial

#### Primary study design

Interventional

## Study type(s)

Treatment

## Health condition(s) or problem(s) studied

Shoulder disfunction post breast and axillary surgery

#### **Interventions**

Control Arm: Standard NHS care post breast and axillary surgery. This involves patients receiving an exercise leaflet and asked to perform exercises at home. Some patients will have an inactivated activity monitor on the arm to track their arm movement and some will have no monitor at all. Patients will be followed up for up a minimum of 1 month post surgery. Intervention arm: patients will be asked to wear a wearable rehabilitation platform (OnTrack App on Apple Watch) for a period of 1 month post surgery in addition to the standard NHS care. The App tracks their movement, gives real-time feedback on activity along with push notifications to encourage movement, allowing remote rehabilitation and remote clinician input. They will also be followed up for 1 month.

Randomisation will be using block randomisation via the Sealed Envelope online platform.

#### Intervention Type

Device

#### Phase

Not Applicable

#### Drug/device/biological/vaccine name(s)

OnTrack Rehab

#### Primary outcome(s)

- 1. Patient perceptions, satisfaction and acceptability regarding the use of OnTrack measured using semi-structured interviews and a System Usability Scale at the end of the study period.
- 2. The effect of OnTrack on post-operative upper limb pain measured using validated pain analogue scale at baseline and 1 month after surgery.
- 3. The effect of OnTrack on post-operative upper-limb movement in the affected limb measured using patient reported outcome measures (DASH questionnaire) along with range of movement, using a goniometer and upper limb strength using a dynamometer, at baseline and 1 month after surgery.
- 4. The effect of OnTrack on patient's post operative health related quality of life measured using EQ5D5L QoL questionnaire at baseline and 1 month post surgery

#### Key secondary outcome(s))

- 1. Healthcare professionals' (surgeons, nurse specialists and physiotherapists) perceptions and acceptability regarding the use of OnTrack measured using semi-structured interviews at the end of the study period.
- 2. Impact on the workload of healthcare professionals measured using semi-structured interviews at the end of the study period.

#### Completion date

04/08/2025

# **Eligibility**

#### Key inclusion criteria

- 1. Age ≥ 18 years
- 2. Planned for sentinel node biopsy or axillary node clearance as part of their breast cancer management
- 3. Participants must be able to provide informed consent

## Participant type(s)

Patient

#### Healthy volunteers allowed

No

#### Age group

Adult

#### Lower age limit

18 years

#### Sex

All

#### Total final enrolment

70

#### Key exclusion criteria

- 1. Unstable medical condition
- 2. Pre-existing pre-operative shoulder disability affecting shoulder movement
- 3. Patients unable to provide informed consent

#### Date of first enrolment

30/03/2025

#### Date of final enrolment

30/06/2025

## Locations

#### Countries of recruitment

**United Kingdom** 

England

# Study participating centre

**Charing Cross Hospital** 

Fulham Palace Road London United Kingdom W6 8RF

# Sponsor information

#### Organisation

Imperial College London

#### **ROR**

https://ror.org/041kmwe10

# Funder(s)

#### Funder type

University/education

#### **Funder Name**

#### Funder Name

Wellcome Trust

# **Results and Publications**

### Individual participant data (IPD) sharing plan

The datasets generated and or analysed during current study will be available upon request from Ahmed Latif ahmed.latif@imperial.ac.uk

### IPD sharing plan summary

Available on request

### **Study outputs**

Output type Details Date created Date added Peer reviewed? Patient-facing?

Participant information sheet Participant information sheet 11/11/2025 11/11/2025 No Yes