PRIMETIME – Post-operative avoidance of radiotherapy in minimal risk women: patient selection using biomarkers

Submission date	Recruitment status No longer recruiting	[X] Prospectively registered		
30/01/2017		[X] Protocol		
Registration date	Overall study status	Statistical analysis plan		
07/03/2017	Ongoing Condition category	☐ Results		
Last Edited		Individual participant data		
27/11/2025	Cancer	[X] Record updated in last year		

Plain English summary of protocol

https://www.cancerresearchuk.org/about-cancer/find-a-clinical-trial/study-radiotherapy-women-small-risk-breast-cancer-returning-primetime

Contact information

Type(s)

Public

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Additional identifiers

Clinical Trials Information System (CTIS)

Nil known

Integrated Research Application System (IRAS)

190307

ClinicalTrials.gov (NCT)

Nil known

Central Portfolio Management System (CPMS)

CPMS 33217

Study information

Scientific Title

Post-operative avoidance of radiotherapy: biomarker selection of women categorised to be in a very low risk group by IHC4+C

Acronym

PRIMETIME

Study objectives

The aim of this study is to test whether radiotherapy can be safely avoided in a patient population considered to have such a low risk of local recurrence that the potential absolute gain from radiotherapy is so small as to not outweigh the established risks associated with breast radiotherapy.

Ethics approval required

Old ethics approval format

Ethics approval(s)

East of England – Cambridgeshire and Hertfordshire Research Ethics Committee, 05/09/2016, ref: 16/EE/0305

Study design

Non-randomized; Interventional; Design type: Treatment, Radiotherapy, Management of Care

Primary study design

Interventional

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Breast cancer

Interventions

Patients consent and register for pre-screening Ki67 research testing and 5 slides from the diagnostic block are forwarded for analysis to the central laboratory. Breast conserving surgery will proceed according to standard practice. If a patient is confirmed as eligible (meets inclusion and exclusion criteria) they will then be consented for the main PRIMETIME study. The IHC4+C calculator is then utilised to direct treatment according to the patient's risk category (i.e. if they are deemed 'very low risk' they will be recommended to avoid radiotherapy). All patients will then receive either standard breast radiotherapy or no radiotherapy and standard adjuvant hormone therapy and any additional anti-cancer treatments.

For patients who receive radiotherapy they will have 5 years of annual mammograms, for patients who do not receive radiotherapy they will be required to attend for 10 years of annual mammograms. Patients will be followed up for 10 years in clinic and thereafter via routine data sources.

Added 27/11/2025:

Additional Data Linkage Information:

Participants from this trial will also be included in the INTERACT project which will link to their data held by NHS England. For more information, please see the INTERACT website: https://www.icr.ac.uk/interact.

Intervention Type

Other

Primary outcome(s)

Ipsilateral breast local relapse rate 5 years from study registration is assessed through patient note review.

Key secondary outcome(s))

- 1. Ipsilateral breast local relapse rate is measured by patient note review 10 years from study registration
- 2. Regional relapse rate is measured by patient note reviewat 5 and 10 years
- 3. Distant relapse rate is measured by patient note review at 5 and 10 years
- 4. Overall survival is measured by patient note review at 5 and 10 years
- 5. Breast cancer specific survival is measured by patient note review at 5 and 10 years

Completion date

19/05/2030

Eligibility

Key inclusion criteria

- 1. Provision of written informed consent to participate in the PRIMETIME study
- 2. Provision of slides for research testing and availability of KI67 result (contact ICR-CTSU to confirm)
- 3. Women aged ≥60 years (younger patients are eligible if they are post-menopausal and have comorbidities that imply a high risk of radiotherapy toxicity e.g. significant cardiovascular disease with left sided breast cancer)
- 4. Women having had breast conserving surgery with complete resection of tumour tissue (≥1 mm microscopic, circumferential margins of normal tissue from invasive cancer and DCIS)
- 5. AJCC staging of pT1/pN0/M0 (DCIS is allowed in combination with invasive breast cancer; isolated tumour cells in axillary nodes are allowed)
- 6. Histological confirmation of grade 1 or 2 invasive breast cancer
- 7. Oestrogen receptor (ER) positive according to local practice. The H score must be available.
- 8. Progesterone receptor (PR) positive according to local practice. The percentage positivity must be available.
- 9. Human epidermal growth factor receptor (HER2) negative according to local practice
 10. Patients must be recommended for >5 years adjuvant endocrine therapy according to
- 10. Patients must be recommended for ≥5 years adjuvant endocrine therapy according to local policy and in the investigator's opinion, deemed able to comply with the duration of treatment

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Sex

Female

Total final enrolment

0

Key exclusion criteria

- 1. Patients known to have lymphovascular space invasion and/or axillary nodal micrometastases or macrometastases.
- 2. Patients with a past history of malignancy excep:
- 2.1. Basal cell skin cancer and CIN cervix uteri
- 2.2.Treated, localised squamous cell carcinoma of the skin
- 2.3. Malignancies treated with curative intent and the patient has been disease free ≥5 years
- 3. Patients who have had an ipsilateral mastectomy
- 4. Patients who have received neoadjuvant therapy (endocrine or cytotoxic chemotherapy with therapeutic intent) or who are deemed by the MDT to require adjuvant cytotoxic chemotherapy 5. Patients with mammographically occult breast cancers, ie. present with lump, but not visible on mammogram

Date of first enrolment

17/03/2017

Date of final enrolment

28/02/2022

Locations

Countries of recruitment

United Kingdom

England

Scotland

Wales

Study participating centre Addenbrookes Hospital

Hills Road

Cambridge England CB2 0QQ

Study participating centre Raigmore Hospital Old Perth Road

Inverness Scotland IV2 3UJ

Study participating centre Wrexham Maelor

Cresnewydd Road Wrexham Wales LL13 7TD

Study participating centre Norfolk and Norwich University Hospital

Colney Lane Norwich England NR4 7UY

Study participating centre Barts Hospital

W Smithfield London England EC1A 7BE

Study participating centre Royal Free Hospital

Pond Street London England NW3 2QG

Study participating centre Mount Vernon Hospital

Rickmansworth Road Northwood England HA6 2RN

Study participating centre Lister Hospital

Coreys Mill Lane, Stevenage England SG1 4AB

Study participating centre Queen Elizabeth II Hospital

Howlands Welwyn Garden City England AL7 4HQ

Study participating centre Charing Cross Hospital

Fulham Palace Road London England W6 8RF

Study participating centre University Hospital of South Manchester

Southmoor Road, Wythenshawe Manchester England M23 9LT

Study participating centre Churchill Hospital

Old Road, Headington Oxford

Study participating centre The Hillingdon Hospital

Pield Heath Road Uxbridge England UB8 3NN

Study participating centre Clatterbridge Cancer Centre

Clatterbridge Health Park, Clatterbridge Road, Birkenhead Wirral England CH63 4JY

Study participating centre The Roval Marsden

203 Fulham Road London England SW3 6JJ

Study participating centre The Royal Marsden

Downs Road Sutton England SM2 5PT

Study participating centre The Royal Bournemouth Hospital

Castle Lane East Bournemouth England BH7 7DW

Study participating centre Royal Glamorgan Hospital

Ynysmaerdy Llantrisant Wales CF72 8XR

Study participating centre Musgrove Park Hospital

Parkfield Drive Taunton England TA1 5DA

Study participating centre West Suffolk Hospital

Hardwick Lane Bury Saint Edmunds England IP33 2QZ

Sponsor information

Organisation

Institute of Cancer Research

ROR

https://ror.org/043jzw605

Funder(s)

Funder type

Charity

Funder Name

Cancer Research UK

Alternative Name(s)

CR_UK, Cancer Research UK - London, Cancer Research UK (CRUK), CRUK

Funding Body Type

Private sector organisation

Funding Body Subtype

Other non-profit organizations

Location

United Kingdom

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary

Data sharing statement to be made available at a later date

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
HRA research summary			28/06/2023	No	No
Interim results article		14/06/2021	13/08/2021	Yes	No
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes
Protocol file	version v3.0	03/05/2019	16/03/2021	No	No
Protocol file	version 4.0	13/01/2021	07/11/2023	No	No
Study website	Study website	11/11/2025	11/11/2025	No	Yes