# NHS Cancer Vaccine Launch Pad (NHS CVLP)

Submission date 09/07/2024	<b>Recruitment status</b> Recruiting	<ul> <li>Prospectively registered</li> <li>Protocol</li> </ul>
<b>Registration date</b> 09/10/2024	<b>Overall study status</b> Ongoing	<ul> <li>Statistical analysis plan</li> <li>Results</li> </ul>
Last Edited 23/05/2025	<b>Condition category</b> Cancer	<ul> <li>Individual participant data</li> <li>[X] Record updated in last year</li> </ul>

# Plain English summary of protocol

Background and study aims

Personalised cancer vaccines are a type of cancer treatment designed to target an individual's unique cancer cells. They are created by analysing a patient's tumour and identifying specific DNA changes that are unique to that cancer, then using that information to create a vaccine tailored to that patient. The intention is that the cancer vaccine will stimulate the immune system to specifically recognise and destroy the cancer cells.

The NHS Cancer Vaccine Launch Pad (CVLP) is looking to find people who may be able to take part in research trials for personalised vaccines to treat cancer. By supporting these trials, the CVLP aims to speed up the development of personalised treatments for cancer patients.

Who can participate?

NHS cancer patients in England aged over 16 years with capacity to consent.

What does the study involve?

Upon consent, the following will occur:

1. Small sections of the tumor will be cut from the diagnostic block by a histopathologist (a doctor who specializes in studying tissues). These sections will be sent to CVLP partners for nucleic acid extraction and sequencing. Nucleic acids are molecules like DNA and RNA that carry genetic information.

 A blood sample, up to 30 milliliters, will be drawn and sent to CVLP partners. This sample may be tested for germline nucleic acid (genetic information you were born with) and tumor circulating free DNA (genetic material released by the tumor into the bloodstream).
 Baseline demographic (like age and gender) and clinical data will be recorded by the clinical or

research team and sent with the biological material in a pseudonymised form (using codes instead of names to protect your identity). This data will also be centrally held by Southampton Clinical Trials Unit (SCTU) to coordinate with cancer vaccine trial partners and sites. A clinical liaison team will oversee this process.

4. During Phase 1, sequencing data will be held by industry partners. If you are eligible for a cancer vaccine trial, the clinical liaison team will inform your clinical team and local trial sites. Pseudonymised codes will be used to identify patients when communicating with clinical teams or linking patients with NHS cancer vaccine trial sites. Any other information shared outside the NHS CVLP framework will also be pseudonymised.

5. If you are eligible for a cancer vaccine trial, your home clinical or research team, or the team at the nearest cancer vaccine trial site, will approach you for further discussion and a separate consent.

What are the possible benefits and risks of participating?

By taking part in the CVLP, participants might be able to take part in a research trial and receive personalised cancer vaccine treatment before it is widely available. For participants who receive this personalised treatment, it may reduce the chance of their cancer growing or coming back. Not everyone will be suitable to take part in a research trial and the research trial may not benefit all participants directly. This is because we are still finding out how effective the treatment might be.

Participants can also benefit from the knowledge that they are helping medical research to improve cancer treatment. The tests and treatments being used were developed with the help of participants who took part in research years ago. The more participants that take part in research, the faster progress can be made as more human samples can be studied. Where is the study run from?

The study is coordinated by the Southampton Clinical Trials Unit and is being run in NHS hospitals in England. The study is sponsored by NHS England.

When is the study starting and how long is it expected to run for? May 2023 to December 2030

Who is funding the study? NHS England (UK)

Who is the main contact? Southampton Clinical Trials Unit at cvlp@soton.ac.uk

# Study website

https://www.southampton.ac.uk/ctu/cancer-vaccine-launch-pad.page#home

# **Contact information**

**Type(s)** Scientific

**Contact name** Miss Nicole Keyworth

**Contact details** Southampton Clinical Trials Unit, Southampton General Hospital Southampton United Kingdom SO16 6YD +44 2381205154 cvlp@soton.ac.uk

# Additional identifiers

EudraCT/CTIS number

## Nil known

**IRAS number** 325291

**ClinicalTrials.gov number** Nil known

Secondary identifying numbers CPMS 57835. IRAS 325291

# Study information

# Scientific Title

Establishing a UK platform to collect tumour samples and perform DNA/RNA sequencing for the development of personalised cancer vaccines

# Acronym

CVLP

# **Study objectives**

The objectives of the NHS Cancer Vaccine Launch Pad (CVLP) are to:

1. Identify cancer patients undergoing resection or biopsy of their tumours who could be eligible for personalised mRNA therapies such as cancer vaccines.

2. Create a pathway for collection and transfer of tumour samples, blood samples and pseudonymised clinical data during routine NHS care.

3. Send samples for DNA extraction, molecular analysis and sequencing initially in partnership with industry (Phase 1) and later to be performed by the NHS Genomic Medicine Service (Phase 2).

4. Use the results to accelerate access of eligible cancer patients to cancer vaccine trials.

The programme aims to profile up to 10,000 tumour samples in a phased approach over 7 years. To achieve this we will enrol patients diagnosed with different cancers and seek their consent for:

1. Collection of surplus tumour biopsies or surgical resection samples to be sent to genetic testing laboratories for DNA or RNA extraction and sequencing.

2. Collection of a blood sample to perform relevant tests for cancer vaccines, eg: circulating free tumour DNA or sequencing of healthy cells to compare to the tumour.

3. Collection of routine clinical data on demographics, staging, diagnostic tests, treatment and outcomes.

4. The linkage of this data in a database held by a central team to co-ordinate between clinical sites, industry partners and cancer vaccine trial sites.

5. Onwards referral of potentially eligible patients to CV trials or linked research studies provided relevant research ethics approval has been granted - for which separate consent will be sought.

No specific therapeutic intervention will be carried out under this protocol and any activities for the CVLP will happen in addition to routine genetic testing that is carried out as standard of care. If CV gain regulatory approval we hope the CVLP pathway will become part of routine care.

# Ethics approval required

Ethics approval required

# Ethics approval(s)

Approved 20/08/2023, East of England – Essex Research Ethics Committee (2 Redman Place, Stratford, London, E20 1JQ, United Kingdom; +44 207 104 8177; essex.rec@hra.nhs.uk), ref: 23 /EE/0178

Study design

Observational cohort study

**Primary study design** Observational

**Secondary study design** Cohort study

**Study setting(s)** Hospital, Medical and other records

**Study type(s)** Other

Participant information sheet

See study outputs table

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

Cancer vaccine studies in various cancer types

# Interventions

Eligible patients with a number of different cancers will be approached around the time of diagnostic biopsy or surgical resection. This could be either during their initial diagnosis of cancer or at the time of recurrence. In all cases the patient will be aware that a diagnosis of cancer has already been made, or is a strong possibility. The patient will be approached by the clinical team managing their care to discuss the study and a standard patient information sheet will be provided.

Patients who agree to take part will have their informed consent confirmed and recorded by one of the clinical team or a research associate.

Following consent, the following will occur:

1. Cuttings of sections of tumour from the diagnostic block under the supervision of a histopathologist, for forwarding to CVLP partners (Phase 1) for extraction and sequencing of nucleic acid.

2. Drawing of a blood sample (up to 30mls) also for forwarding to CVLP partners. Potential tests performed on blood samples include germline extraction of nucleic acid for the comparison with the tumour sample or tumour circulating free tumour DNA.

3. Recording of baseline demographic and clinical data will be done by the clinical or research team. This will be sent along with biological material in a pseudonymised form. Data will be held centrally by the Southampton Clinical Trials Unit (SCTU).

4. During Phase 1 sequencing data will be held by industry partners. If a patient is deemed to be eligible for a cancer vaccine trial this will be fed back to the clinical liaison team who will inform

the clinical teams and any local trial sites. The pseudonymised codes will only be used for reidentifying patients when communicating with the clinical team or to link patients with NHS cancer vaccine trial sites. Any other information shared outside the NHS CVLP framework will be pseudonymised.

5. If a patient is deemed eligible for a cancer vaccine trial they will be approached by either their home clinical or research team, or by the team at the nearest cancer vaccine trial site for further discussion and a separate consent.

## Intervention Type

Other

# Phase

Not Specified

# Primary outcome measure

1. Number of participants recruited to the CVLP by September 2024

2. Number of participants recruited to the CVLP measured every 12 months until study closure

3. Proportion of eligible patients approached that consent to participate in the CVLP measured every 12 months until study closure

4. Proportion of tumour samples available for analysis measured at 2 weeks post consent to a cancer vaccine trial

5. Number of participants proceeding to participate in a cancer vaccine trial measured monthly until study closure

# Secondary outcome measures

Number of sequencing outputs from NHS laboratories used for manufacturing of therapeutic personalised cancer vaccines measured every 12 months until study closure

# Overall study start date

01/05/2023

# **Completion date**

31/12/2030

# Eligibility

# Key inclusion criteria

- 1. Be over the age of 16 years
- 2. Have a tumour which has been, or will be resected or biopsied
- 3. Have the capacity to consent to involvement in the CVLP
- 4. Have sufficient tumour available for genomic analyses

**Participant type(s)** Patient

**Age group** Adult

**Lower age limit** 16 Years **Sex** Both

**Target number of participants** 10,000

**Key exclusion criteria** 1. Under the age of 16 years 2. Incapable of giving informed consent

Date of first enrolment 31/08/2024

Date of final enrolment 31/03/2026

# Locations

## **Countries of recruitment** England

United Kingdom

**ME16 9QQ** 

#### Study participating centre Maidstone Maidstone Hospital Hermitage Lane Maidstone United Kingdom

# Study participating centre Tunbridge Wells Hospital

The Tunbridge Wells Hospital Tonbridge Road Pembury Tunbridge Wells United Kingdom TN2 4QJ

#### **Study participating centre Southampton** Southampton General Hospital Tremona Road Southampton

United Kingdom SO16 6YD

#### **Study participating centre University Hospital Lewisham** Lewisham High Street London United Kingdom SE13 6LH

# Study participating centre Queen Elizabeth Hospital

Woolwich Stadium Road Woolwich London United Kingdom SE18 4QH

#### **Study participating centre Royal Surrey County Hospital** Egerton Road Guildford United Kingdom GU2 7XX

**Study participating centre Royal Sussex County Hospital** Eastern Road Brighton United Kingdom BN2 5BE

### **Study participating centre Conquest Hospital** The Ridge St. Leonards-on-sea

United Kingdom TN37 7RD

#### Study participating centre Eastbourne District General Hospital

Kings Drive Eastbourne United Kingdom BN21 2UD

# Study participating centre Addenbrookes

Addenbrookes Hospital Hills Road Cambridge United Kingdom CB2 0QQ

#### Study participating centre Queens Hospital

Rom Valley Way Romford United Kingdom RM7 0AG

# Study participating centre

**Colchester General Hospital** Colchester District General Hosp. Charter Way Turner Road Colchester United Kingdom CO4 5JL

# Study participating centre

**Ipswich Hospital** Heath Road Ipswich United Kingdom IP4 5PD

**Study participating centre Stoke Mandeville Hospital** Mandeville Road Aylesbury United Kingdom HP21 8AL

#### **Study participating centre Wycombe Hospital** Queen Alexandra Road High Wycombe United Kingdom HP11 2TT

#### **Study participating centre Hillingdon Hospital** Hillingdon Hospital Pield Heath Road Uxbridge United Kingdom

UB8 3NN

OX3 7LE

# Study participating centre Churchill Hospital Old Road Headington Oxford United Kingdom

#### **Study participating centre Queens Medical Centre, Nottingham University Hospital** Derby Road Nottingham United Kingdom NG7 2UH

# Study participating centre

**Nottingham City Hospital** Hucknall Road Nottingham United Kingdom NG5 1PB Study participating centre Warwick Hospital Lakin Road Warwick United Kingdom CV34 5BW

Study participating centre New Cross Hospital Royal Wolverhampton Wolverhampton Road Heath Town Wolverhampton United Kingdom WV10 0QP

**Study participating centre Walsall Manor Hospital** Moat Road Walsall United Kingdom WS2 9PS

**Study participating centre Royal Stoke University Hospital** Newcastle Road Stoke-on-trent United Kingdom ST4 6QG

Study participating centre Salford Royal Hospital Stott Lane Eccles Salford United Kingdom M6 8HD

#### The Royal Oldham Hospital

Rochdale Road Oldham United Kingdom OL1 2JH

### Study participating centre Fairfield General Hospital

Fairfield General Hospital Rochdale Old Road Bury United Kingdom BL9 7TD

# Study participating centre

**Rochdale Infirmary** Whitehall Street Rochdale United Kingdom OL12 0NB

#### **Study participating centre St. James's University Hospital** Beckett Street Leeds United Kingdom LS9 7TF

#### Study participating centre Leeds General Infirmary

Great George Street Leeds United Kingdom LS1 3EX

### Study participating centre

Weston Park Hospital Whitham Road Sheffield United Kingdom S10 2SJ Study participating centre Derriford Hospital Derriford Road Derriford Plymouth United Kingdom PL6 8DH

**Study participating centre Musgrove Park Hospital** Parkfield Drive Taunton United Kingdom TA1 5DA

**Study participating centre Yeovil District Hospital** Higher Kingston Yeovil United Kingdom BA21 4AT

**Study participating centre Royal Devon and Exeter Hospital** Royal Devon & Exeter Hospital Barrack Road Exeter United Kingdom EX2 5DW

**Study participating centre North Devon District Hospital** Raleigh Park Barnstaple United Kingdom EX31 4JB

## Bristol Haematology & Oncology Centre

Horfield Road Bristol United Kingdom BS2 8ED

#### Study participating centre Southmead Hospital

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

# Study participating centre

**Royal United Hospital Bath** Combe Park Bath United Kingdom BA1 3NG

#### **Study participating centre Cheltenham General Hospital** Sandford Road Cheltenham United Kingdom GL53 7AN

#### **Study participating centre Gloucestershire Royal Hospital** Great Western Road Gloucester United Kingdom GL1 3NN

# Study participating centre

**Royal Lancaster Infirmary** Ashton Road Lancaster United Kingdom LA1 4RP **Study participating centre Royal Blackburn Hospital** Haslingden Road Blackburn United Kingdom BB2 3HH

**Study participating centre Cumberland Infirmary** Newtown Road Carlisle United Kingdom CA2 7HY

**Study participating centre West Cumberland Hospital** Homewood Hensingham Whitehaven United Kingdom CA28 8JG

**Study participating centre Sunderland Royal Hospital** Kayll Road Sunderland United Kingdom SR4 7TP

**Study participating centre South Tyneside District General Hospital** Harton Lane South Shields United Kingdom NE34 0PL

**The James Cook University Hospital** Marton Road Middlesbrough

United Kingdom TS4 3BW

**Study participating centre North Tyneside General Hospital** Rake Lane North Shields United Kingdom NE29 8NH

Study participating centre Broomfield University Hospital Broomfield Hospital Court Road Chelmsford United Kingdom CM1 7ET

**Study participating centre Basildon University Hospital** Nethermayne Basildon United Kingdom SS16 5NL

**Study participating centre Southend University Hospital** Prittlewell Chase Westcliff-on-sea United Kingdom SS0 0RY

**Study participating centre Royal Bournemouth General Hospital** Castle Lane East Bournemouth United Kingdom BH7 7DW **Study participating centre Poole Hospital** Longfleet Road Poole United Kingdom BH15 2JB

**Study participating centre St Marks Hospital** Watford Road Harrow United Kingdom HA1 3UJ

Study participating centre East Surrey Hospital Canada Avenue Redhill United Kingdom RH1 5RH

**Study participating centre Basingstoke and North Hampshire Hospital** Aldermaston Road Basingstoke United Kingdom RG24 9NA

**Study participating centre University Hospital of North Tees** Hardwick Road Stockton-on-tees United Kingdom TS19 8PE

**Study participating centre Great Western Hospital** Marlborough Road Swindon United Kingdom SN3 6BB

#### Study participating centre Milton Keynes University Hospital Standing Way Eaglestone Milton Keynes United Kingdom MK6 5LD

**Study participating centre Royal Hampshire County Hospital** Romsey Road Winchester United Kingdom SO22 5DG

### **Study participating centre Castle Hill Hospital** Castle Road Cottingham United Kingdom HU16 5JX

#### **Study participating centre Bedford Hospital South** Kempston Road Bedford United Kingdom MK42 9DJ

**Study participating centre Luton and Dunstable University Hospital** Lewsey Road Luton United Kingdom LU4 0DZ

#### **Study participating centre West Middlesex University Hospital** Twickenham Road Isleworth United Kingdom TW7 6AF

#### Study participating centre

**University Hospital Coventry & Warwickshire** Clifford Bridge Road Walsgrave Coventry United Kingdom CV2 2DX

#### Study participating centre Russells Hall Hospital

Pensnett Road Dudley United Kingdom DY1 2HQ

#### **Study participating centre Manchester Royal Royal Infirmary** Cobbett House Oxford Road Manchester United Kingdom M13 9WL

### **Study participating centre Wythenshawe Hospital** Southmoor Road Wythenshawe Manchester United Kingdom

M23 9LT

## **Chesterfield Royal Hospital**

Chesterfield Road Calow Chesterfield United Kingdom S44 5BL

#### **Study participating centre Harrogate District Hospital** Lancaster Park Road Harrogate United Kingdom HG2 7SX

#### **Study participating centre Rotherham General Hospital** Moorgate Road Rotherham United Kingdom S60 2UD

# Sponsor information

# Organisation

NHS England

### Sponsor details

Wellington House, 133-155 Waterloo Road London England United Kingdom SE1 8UG

england.cancervaccinelaunchpad@nhs.net

**Sponsor type** Government

# Funder(s)

Funder type

Industry

Funder Name

NHS England as sponsor will be receiving funding from multiple industry companies

# **Results and Publications**

**Publication and dissemination plan** Planned publication in a peer-reviewed journal

# Intention to publish date

31/12/2031

# Individual participant data (IPD) sharing plan

The current data sharing plans for this study are unknown and will be available at a later date.

## 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?
Participant information sheet	version 8	22/03/2024	23/07/2024	No	Yes