

# Analysis of liquid biopsies to improve diagnosis and follow-up in cancer and other conditions

<b>Submission date</b> 08/02/2025	<b>Recruitment status</b> Recruiting	<input type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
<b>Registration date</b> 06/06/2025	<b>Overall study status</b> Ongoing	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
<b>Last Edited</b> 06/06/2025	<b>Condition category</b> Cancer	<input type="checkbox"/> Individual participant data <input checked="" type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

Liquid biopsies (LBs) are minimally invasive samples of body fluids that contain tiny amounts of circulating tumour-derived (ct) DNA, RNA and proteins. Thanks to new, ultrasensitive techniques, LBs have emerged as a potential cancer biomarker. It has been shown that analysis of ctDNA from liquid biopsies using broad screening methods, including whole-genome or targeted gene panel sequencing, and sensitive methods that target specific genetic aberrations is possible. Further techniques are being developed for analysing all the components (including RNA and proteins) for multi-omic analysis of LBs (m-LBs), to increase sensitivity and precision. Together with the doctors involved in cancer care, the research team have identified three core clinical questions which will be used to test LBs: 1. Can LBs provide a molecular diagnosis when tissue biopsies are not available (such as some brain tumours)?; 2. Are LBs as good (or better) as current clinical tests for providing information on prognosis and risk for developing metastatic disease?; and, 3. Can LBs be successfully used to monitor therapy response and detect early relapse?

### Who can participate?

Patients with cancer, risk of cancer and other somatic conditions

### What does the study involve?

LBs (blood, cerebrospinal fluid when available, urine) will be collected from study participants. The analyses are performed retrospectively and will not be used in the treatment or follow-up of the participants.

### What are the possible benefits and risks of participating?

There are no direct benefits to participating in this study. The aim is to improve the diagnosis and follow-up of future patients with the same condition. The risks are minor and include minor complications from taking the samples (done in conjunction with clinical sampling) and the small risk (or benefit depending on how you look at it) that a germline predisposition can be identified that would enable us to offer you more information on possible cancer risks and to refer you to an individualised risk reduction program. The samples are usually collected when other samples are collected clinically which entails minimal risks.

Where is the study run from?  
Karolinska University Hospital, Sweden

When is the study starting and how long is it expected to run for?  
January 2015 to December 2040

Who is funding the study?  
1. The Childhood Cancer Foundation  
2. The Cancer Foundation  
3. Stockholm County Council

Who is the main contact?  
Dr Emma Tham, emma.tham@ki.se

## Contact information

**Type(s)**  
Public, Scientific, Principal investigator

**Contact name**  
Dr Emma Tham

**ORCID ID**  
<https://orcid.org/0000-0001-6079-164X>

**Contact details**  
Clinical Genetics and Genomics  
Karolinska University Hospital  
Stockholm  
Sweden  
171 76  
+46-12370000  
emma.tham@ki.se

## Additional identifiers

**Clinical Trials Information System (CTIS)**  
Nil known

**ClinicalTrials.gov (NCT)**  
Nil known

**Protocol serial number**  
Nil known

## Study information

**Scientific Title**  
Multi-omics analyses of liquid biopsies

**Acronym**

Multi-Liq

**Study objectives**

Liquid biopsies can be used as a complement to standard-of-care methods to provide a molecular diagnosis in patients where no tumour biopsy is available and to refine risk stratification and monitoring during treatment and follow-up of cancer patients

**Ethics approval required**

Ethics approval required

**Ethics approval(s)**

1. approved 12/09/2016, Regional Ethics Review Board (Tomtebodavägen 18, Stockholm, 171 77, Sweden; +46 08-52487000; kansli@stockholm.epn.se), ref: 2016/2-31/1

2. approved 19/01/2025, The Swedish Ethical Review Authority (Box 2110, Uppsala, 75002, Sweden; +46-10-475 08 00; registrator@etikprovning.se), ref: 2024-08384-02

**Study design**

Prospective multi-centre retrospective analysis. Longitudinal with comparison to standard of care performed in parallel.

**Primary study design**

Observational

**Study type(s)**

Diagnostic, Screening

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

Cancer, risk of cancer and other somatic conditions

**Interventions**

The collection and analysis of liquid biopsies using various methods, mostly based on cfDNA, but also with multi-omics approaches. Comparison to diagnosis and follow-up performed within routine clinical care to evaluate the added utility of liquid biopsies.

**Intervention Type**

Not Specified

**Primary outcome(s)**

Additional diagnostic or prognostic information provided by liquid biopsies measured using massive parallel sequencing, methylation analysis or targeted digital PCR at diagnosis

**Key secondary outcome(s)**

Prognostic information and correlation to survival from liquid biopsies measured using massive parallel sequencing, epigenetic analysis, targeted digital PCR and/or proteomics at diagnosis, during or after treatment or during follow-up

**Completion date**

31/12/2040

# Eligibility

## Key inclusion criteria

1. Patients with cancer of a type included in the study
2. Individuals with a predisposition for cancer
3. Individuals with other somatic conditions not classified as cancer
4. Not able to give informed consent

## Participant type(s)

Patient

## Healthy volunteers allowed

No

## Age group

All

## Lower age limit

0 days

## Upper age limit

100 years

## Sex

All

## Key exclusion criteria

Not meeting the participant exclusion criteria

## Date of first enrolment

15/11/2016

## Date of final enrolment

31/12/2035

# Locations

## Countries of recruitment

Sweden

## Study participating centre

Karolinska University Hospital

Eugenivägen 3

Stockholm

Sweden

17176

**Study participating centre**  
Akademiska Sjukhuset  
Akademiska Sjukhuset  
Uppsala  
Sweden  
751 85

## Sponsor information

**Organisation**  
Karolinska University Hospital

**ROR**  
<https://ror.org/00m8d6786>

## Funder(s)

**Funder type**  
Research organisation

**Funder Name**  
Barncancerfonden

**Alternative Name(s)**  
Swedish Childhood Cancer Foundation

**Funding Body Type**  
Private sector organisation

**Funding Body Subtype**  
Trusts, charities, foundations (both public and private)

**Location**  
Sweden

**Funder Name**  
Cancerfonden

**Alternative Name(s)**  
Swedish Cancer Society

**Funding Body Type**

Private sector organisation

**Funding Body Subtype**

Trusts, charities, foundations (both public and private)

**Location**

Sweden

**Funder Name**

Stockholms Läns Landsting

**Alternative Name(s)**

Stockholm County Council

**Funding Body Type**

Government organisation

**Funding Body Subtype**

Local government

**Location**

Sweden

**Funder Name**

Japanese Swedish Foundation

**Funder Name**

Karolinska Institutets Funds

## Results and Publications

**Individual participant data (IPD) sharing plan**

All participants receive written and oral information on the study and provide written informed consent. The results from the liquid biopsy analyses and the clinical data will be stored in a non-publicly available study repository (only available to the researchers involved in the project) and relevant results will be published in peer-reviewed journals

**IPD sharing plan summary**

Stored in non-publicly available repository

**Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Participant information sheet</a>	Participant information sheet	11/11/2025	11/11/2025	No	Yes