Developing genetic tests to diagnose, monitor and guide treatment decisions for children and young people whose cancer has returned

Submission date	Recruitment status	[X] Prospectively registered
14/02/2025	Recruiting	☐ Protocol
Registration date	Overall study status	Statistical analysis plan
28/03/2025	Ongoing	Results
Last Edited	Condition category	Individual participant data
28/03/2025	Cancer	[X] Record updated in last year

Plain English summary of protocol

Background and study aims

This is a UK research study testing tumour (somatic) and normal (germline) DNA and RNA for genetic changes in children and young people with relapsed/refractory cancer. This will be done by comparing the genetic information in the patient's healthy cells with their cancer cells, and following them over time to detect changes earlier than their scans. This information will be used to see if it can help to guide new treatment strategies and personalise cancer treatment for patients based on their genetic information.

Who can participate?

UK children and young adults whose cancer has either come back (relapsed) or not responded to treatment (refractory) and have undergone or will undergo a routine biopsy/surgery to obtain tumour tissue or bone marrow.

What does the study involve?

Participants with a solid tumour provide a blood sample and a piece (or pieces, if available) of tumour collected from their most recent biopsy or surgery. Participants with leukaemia provide a bone marrow sample. The results of the tests are relayed back to the patient's doctor via an expert panel who make recommendations on any available treatments. Patients and/or their parents are asked in advance to consider what information they want to receive in relation to any abnormal genetic results either in the tumour or their normal (germline) genetic code. In addition, the data collected is used and shared for the purposes of clinical research.

What are the possible benefits and risks of participating?

Benefits: It is unlikely there will be an individual benefit for the patient by taking part in the StratMedPaediatrics2 study. The greatest benefits of the work may not be expected for several years and therefore will predominantly help future patients. However, should something be found in the genetic information of the patient's tumour which may help in the understanding or treatment of the patient's cancer then the patient's clinical team will be able to use this information (for example provide a treatment option i.e. clinical trial or early indication that the treatment is not working). For solid tumour patients, as the tumour sample will already have

been or is due to be taken as part of the care at the hospital, the patient will only have blood tests taken whilst on the study. The discomfort of this blood test is just like any other blood test. For leukaemia patients, the bone marrow sample will already have been or is due to be taken as part of the care at the hospital.

Where is the study run from?

- 1. Royal Aberdeen's Children Hospital
- 2. Royal Belfast Hospital for Sick Children
- 3. Birmingham Children's Hospital
- 4. Bristol Royal Hospital for Children
- 5. Addenbrooke's Hospital
- 6. Noah's Ark Children's Hospital for Wales
- 7. Royal Hospital for Sick Children Edinburgh
- 8. Royal Hospital for Children
- 9. Leeds General Infirmary
- 10. Leicester Royal Infirmary
- 11. Alder Hey Children's Hospital
- 12. Great Ormond Street Hospital for Children
- 13. Royal Manchester Children's Hospital
- 14. Royal Victoria Infirmary
- 15. Queen's Medical Centre, Nottingham
- 16. John Radcliffe Hospital
- 17. Sheffield Children's Hospital
- 18. Southampton General Hospital
- 19. University College London Hospital
- 20. Royal Marsden Hospital Sutton

When is the study starting and how long is it expected to run for? March 2023 to October 2032

Who is funding the study? Cancer Research UK

Who is the main contact?
Ms Amina Bukhari, stratmedpaeds2@trials.bham.ac.uk

Plain English summary under review with external organisation

Contact information

Type(s)

Public, Scientific

Contact name

Ms Amina Bukhari

Contact details

Trial Coordinator Children's Cancer Trials Team Cancer Research UK Clinical Trials Unit School of Medical Sciences The University of Birmingham
Edgbaston
Birmingham
United Kingdom
B15 2TT
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StratMedPaeds2@trials.bham.ac.uk

Additional identifiers

Clinical Trials Information System (CTIS)

Nil known

Integrated Research Application System (IRAS)

346034

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

CPMS 64809, Cancer Research UK Grant Code CRCEMA-Jul23/100001

Study information

Scientific Title

Stratified Medicine Paediatrics 2

Acronym

StratMedPaeds2 / SMPaeds2

Study objectives

This is a UK research study testing tumour (somatic) and normal (germline) DNA and RNA for genetic changes in children and young people with relapsed/refractory cancer. This will be done by comparing the genetic information in the patient's healthy cells with their cancer cells. This information will be used to see if it can help to guide new treatment strategies and personalise cancer treatment for patients based on their genetic information.

Ethics approval required

Ethics approval required

Ethics approval(s)

approved 29/11/2024, Yorkshire & The Humber - Leeds East Research Ethics Committee (2 Redman Place, Stratford, London, E20 1JQ, United Kingdom; +44 207 104 8171, 2071048137, 207 104 8357; leedseast.rec@hra.nhs.uk), ref: 24/YH/0218

Study design

Observational clinical laboratory study

Primary study design

Observational

Study type(s)

Diagnostic

Health condition(s) or problem(s) studied

Relapsed/refractory solid tumours (including lymphomas) or leukaemia

Interventions

StratMedPaeds2 is a prospective cohort molecular profiling study that will generate highly annotated clinical and molecular data to highlight novel and potentially druggable dependencies enriched at the time of cancer relapse, in support of a cohort of aligned experimental clinical trials that seek to deploy molecularly targeted drugs and immunotherapies, within molecularly enriched and serially monitored trial designs. Descriptive, highly clinically annotated molecular datasets will be developed on the spectrum and frequency of genetic, epigenetic and immune-microenvironmental changes that characterise relapsed childhood solid tumours. This information will be made available within a shared database, to deliver a harmonised and comprehensive information resource that can be readily used for clinical management and research. This will report in detail on the test performance parameters of the methodologies developed, and within the limitations of a relatively small study in rare disease indications, output statistically powered conclusions where possible on the best use of the tests developed, to support future clinical integration of the technologies. Of particular importance for the study is to investigate the clinical utility of liquid biopsies, and their ability to detect novel and potentially actionable events at relapse compared to existing tissue-biopsy testing.

StratMedPaeds2 will open in the UK across 20 paediatric primary treatment centres. The centres have been chosen based on their clinical expertise in the treatment of children with cancer. The recruitment target for the duration of the study is 400 patients.

For patients who meet the eligibility criteria for the study, the investigator will provide them /their families with information to allow a decision regarding their participation. The patient /parent/guardian will be required to provide consent for the collection and analysis of the patient samples, the collection of relevant clinical information, the return of clinical results back to their investigator, the use of and sharing of data for research, teaching, commercial and scientific publications, and the sharing of samples for other and future ethically approved research projects.

If informed consent is given, the investigator will conduct a screening evaluation to ensure that the patient satisfies all eligibility criteria. Following consent, the site investigators will register the patient onto the study, thus obtaining a patient-specific unique study ID number for that patient. Following registration, the site team must send the baseline ('study entry') following samples to the central sample hub at Great Ormond Street Hospital. All samples are taken outside of the study as part of the patient's standard care, except for one blood sample in solid tumour patients.

The samples are then analysed at the central sample hub at Great Ormond Street Hospital (GOSH), this would also happen as part of the patient's standard care, the Institute of Cancer Research (ICR)/Royal Marsden Hospital, and University of Birmingham (UoB) as part of the StratMedPaeds2 research.

The findings from this analysis will be presented to a Molecular Tumour Board (MTB) based at Great Ormond Street Hospital. At the MTB, selected findings of clinical significance will be presented to (as a minimum) a pathologist, molecular pathologist and an oncologist for a

combined review of the molecular findings in context. The MTB will discuss and finalise the presented findings and will report to the referring clinician, site pathologist and CRCTU.

Information about a patient's participation in a clinical trial and survival outcome will be collected for at least 3 years, this may be extended until all patients have a minimum of 3 years of follow-up.

Intervention Type

Other

Phase

Not Specified

Primary outcome(s)

Genetic changes in the patient's relapsed/refractory cancer cells will be studied using genetic analysis in both the tumour and blood at the time of relapse and throughout their treatment journey to fulfil the following objectives:

- 1. The proportion of patients in whom clinically relevant genomic events are detected at the time of relapse
- 2. The proportion of patients in whom treatment is altered or who have a positive diagnosis as a direct result of either tissue or liquid biopsies.
- 3. The frequency and spectrum of events detected at the time of relapse

Key secondary outcome(s))

Genetic changes in the patient's relapsed/refractory cancer cells will be studied using genetic analysis in both the tumour and blood at the time of relapse and throughout their treatment journey to fulfil the following objectives:

- 1. Proportion of diagnoses that are refined as a result of molecular testing
- 2. Percentage of cases in which long-read sequencing reports a methylation classifier for diagnosis of CNS tumours and sarcoma
- 3. The proportion of patients on therapy who develop actionable or novel treatment resistance
- 4. mutations identified in serial liquid biopsy
- 5. Association of ctDNA levels by serial liquid biopsy during treatment with clinical and/or radiological Indicators of progression
- 6. The turnaround time (TAT) from receipt of sample at GOSH to discussion of results at MTB
- 7. The identification of events which contribute to TAT, beginning from patient consent to final reporting via the MTB

Completion date

01/10/2032

Eligibility

Key inclusion criteria

- 1. Age 0-21 years (patients > 21yo where primary cancer is classified as a "paediatric specific malignancy")
- 2. Patients with relapsed/refractory paediatric solid & CNS tumours, Leukaemia and Lymphoma. Note: refractory leukaemia will only be eligible where no standard 2nd line treatment is available. Contact the leukaemia lead prior to registration
- 3. For solid tumours: Patient has a Formalin fixed paraffin embedded (FFPE) tumour (mandatory) and fresh frozen tumour (if available) from a biopsy, resection or other surgical procedure that

was taken within 8 weeks prior to study entry (as part of NHS SoC). Fresh frozen tumour tissue is highly encouraged*1.

- 4. For leukaemia –Viable fresh or frozen Bone Marrow aspirate sample taken at a prior assessment within 8 weeks prior to study entry*2 (taken as part of NHS SoC) For BM and combined relapses where bone marrow is unavailable, a peripheral blood sample can be provided if circulating blasts are confirmed on morphology or flow cytometry.
- 5. For isolated CNS/Combined relapses where bone marrow is unavailable or BM is uninvolved with leukaemia, a CSF sample should be provided (taken as part of NHS SoC).
- 6. Written informed consent of patient/parent/guardian
- *1. To allow full multi-omic analysis both fresh frozen and Formalin fixed paraffin embedded (FFPE) tumour plus a blood sample for constitutional (germline) and circulating tumour (ct) DNA will need to be available. Original diagnostic slides should be submitted at the same time as block from current relapse/refractory episode either in the same shipment (see laboratory manual for further details). For CNS tumours only: blood and CSF samples paired with initial SoC tumour tissue samples
- *2. Where available, a cerebrospinal fluid (CSF) sample in the event of an isolated or combined CNS relapse should also be provided in addition to the bone marrow aspirate.

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Mixed

Lower age limit

0 years

Upper age limit

21 years

Sex

All

Key exclusion criteria

Not meeting the participant inclusion criteria

Date of first enrolment

14/04/2025

Date of final enrolment

14/09/2029

Locations

Countries of recruitment

United Kingdom

England

Northern Ireland

Scotland

Wales

Study participating centre NHS Grampian Summerfield House 2 Eday Road Aberdeen United Kingdom AB15 6RE

Study participating centre
Cambridge University Hospitals NHS Foundation Trust
Cambridge Biomedical Campus
Hills Road
Cambridge
United Kingdom
CB2 0QQ

Study participating centre
Alder Hey Childrens NHS Foundation Trust
Alder Hey Hospital
Eaton Road
West Derby
Liverpool
United Kingdom
L12 2AP

Study participating centre Belfast City Hospital 51 Lisburn Rd Belfast United Kingdom BT9 7AB

Study participating centre

Birmingham Women's and Children's NHS Foundation Trust

Steelhouse Lane Birmingham United Kingdom B4 6NH

Study participating centre University Hospitals Bristol and Weston NHS Foundation Trust

Trust Headquarters Marlborough Street Bristol United Kingdom BS1 3NU

Study participating centre Cardiff & Vale University Lhb

Woodland House Maes-y-coed Road Cardiff United Kingdom CF14 4HH

Study participating centre Lothian

Waverleygate 2-4 Waterloo PLACE Edinburgh City of Edinburgh United Kingdom EH1 3EG

Study participating centre Gartnavel Royal Hospital

1055 Great Western Road Glasgow United Kingdom G12 0XH

Study participating centre Great Ormond Street Hospital for Children NHS Foundation Trust Great Ormond Street

London United Kingdom WC1N 3JH

Study participating centre St James University Hospital NHS Trust

St James's University Hospital Beckett Street Leeds United Kingdom LS9 7TF

Study participating centre Leicester Royal Infirmary

Infirmary Square Leicester United Kingdom LE1 5WW

Study participating centre Manchester University NHS Foundation Trust

Cobbett House Oxford Road Manchester United Kingdom M13 9WL

Study participating centre

The Newcastle upon Tyne Hospitals NHS Foundation Trust

Freeman Hospital
Freeman Road
High Heaton
Newcastle upon Tyne
United Kingdom
NE7 7DN

Study participating centre Nottingham University Hospitals NHS Trust

Trust Headquarters Queens Medical Centre Derby Road Nottingham United Kingdom NG7 2UH

Study participating centre John Radcliffe Hospital

Headley Way Headington Oxford United Kingdom OX3 9DU

Study participating centre The Royal Marsden NHS Foundation Trust

Fulham Road London United Kingdom SW3 6JJ

Study participating centre Sheffield Childrens Hospital

Western Bank Sheffield United Kingdom S10 2TH

Study participating centre Southampton General Hospital

Tremona Road Southampton United Kingdom SO16 6YD

Study participating centre

University College London Hospitals NHS Foundation Trust

250 Euston Road London United Kingdom NW1 2PG

Sponsor information

Organisation

Institute of Cancer Research

ROR

https://ror.org/043jzw605

Funder(s)

Funder type

Government

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

The datasets generated during and/or analysed during the current study will be stored in a non-publicly available repository, Palantir's Foundry SaaS platform, hosted within the United Kingdom (UK).

- The type of data that will be shared: Pseudonymised Health data, genetic data
- When the data will become available and for how long: The data will be available throughout the study and into the follow-up period (a total of 7.5 years) Analysed full and partial data will be undertaken again throughout the study with the first potential availability being 1 year after the study opening
- By what access criteria the data will be shared including with whom: Trial Data Management Group decide access on a case-by-case basis, based on their legal need for access. Other researchers and groups.

- For what types of analyses, and by what mechanism: Sequencing, genome, spatial
- Whether consent from participants was obtained: Consent will be obtained for all study participants
- Comments on data anonymisation: Data is pseudonymised at the point of study enrolment (study ID is allocated upon registration and used throughout the study)
- Any ethical or legal restrictions, any other comments: Bound by clinical REC approval conditions and sponsor conditions

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