

# A pilot study to investigate how best to integrate a PET-CT scan into the radiotherapy planning pathway for lymphoma

|                                        |                                                   |                                                                                                   |
|----------------------------------------|---------------------------------------------------|---------------------------------------------------------------------------------------------------|
| <b>Submission date</b><br>22/06/2016   | <b>Recruitment status</b><br>No longer recruiting | <input type="checkbox"/> Prospectively registered<br><input type="checkbox"/> Protocol            |
| <b>Registration date</b><br>28/06/2016 | <b>Overall study status</b><br>Completed          | <input type="checkbox"/> Statistical analysis plan<br><input checked="" type="checkbox"/> Results |
| <b>Last Edited</b><br>10/07/2023       | <b>Condition category</b><br>Cancer               | <input type="checkbox"/> Individual participant data                                              |

## Plain English summary of protocol

Not provided at time of registration

## Contact information

### Type(s)

Scientific

### Contact name

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### Contact details

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## Additional identifiers

### Protocol serial number

RD12/10519

## Study information

### Scientific Title

A pilot study to optimise the use of FDG PET-CT and deformable image co-registration for lymphoma radiotherapy planning

**Study objectives**

Study aim is to evaluate the impact of pre-chemotherapy PET-CT performed in the radiotherapy treatment position upon the accuracy of subsequent radiotherapy target volume definition

**Ethics approval required**

Old ethics approval format

**Ethics approval(s)**

NRES Committee Yorkshire & The Humber - Leeds East, 24/05/2013, ref: 13/YH/0139

**Primary study design**

Interventional

**Study design**

Single centre non-randomised interventional pilot study

**Study type(s)**

Other

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

High grade lymphoma

**Interventions**

The study aims to recruit up to 20 patients with early stage Hodgkin lymphoma or high grade non-Hodgkin lymphoma. A routine staging PET-CT will be followed at the same session by a PET-CT in the radiotherapy treatment position appropriate radiotherapy immobilisation devices.

Participation in the study will not affect treatment decisions or the radiotherapy planning process. The process of planning radiotherapy will not be systematically altered by the study.

**Intervention Type**

Device

**Phase**

Not Applicable

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

-

**Primary outcome(s)**

The impact of pre-chemotherapy PET-CT performed in the radiotherapy treatment position upon the subsequent radiotherapy target volume on retrospective analysis. (This will not alter routine treatment for participants). This endpoint will be evaluated using methods of positional analysis to compare radiotherapy target volumes that are constructed with and without the benefit of the PET-CT acquired in the radiotherapy position.

**Key secondary outcome(s)**

To assess the accuracy of image coregistration between pre-chemotherapy PET-CT and the post-chemotherapy planning CT scan.

**Completion date**

31/12/2017

## **Eligibility**

**Key inclusion criteria**

1. Age  $\geq 18$  years
2. WHO Performance status 0-2 (Appendix A)
3. Histologically proven Hodgkin lymphoma or high grade non-Hodgkin lymphoma
4. Ann Arbor Stage I/II disease based upon clinical examination and any radiology investigations performed
5. Clinical decision to proceed with sequential chemotherapy and radiotherapy if stage I/II disease is confirmed on subsequent PET-CT staging
6. Residual disease in situ after biopsy (either palpable or on any imaging acquired pre-PET-CT)
7. Able to provide fully informed written consent
8. Able to lie flat for 1 hour
9. Not be pregnant or breast feeding. Female patients of childbearing potential must agree to use effective contraception, be surgically sterile, or be post-menopausal

**Participant type(s)**

Patient

**Healthy volunteers allowed**

No

**Age group**

Adult

**Lower age limit**

18 Years

**Sex**

All

**Total final enrolment**

19

**Key exclusion criteria**

1. Hypersensitivity to  $^{18}\text{F}$ Fluorine-FDG
2. Hypersensitivity to iodinated contrast media
3. Poorly controlled diabetes
4. Acute renal failure or moderate renal impairment (estimated glomerular filtration rate  $< 30 \text{ mL/min}$ )
5. Uncontrolled pain
6. Urinary incontinence
7. Female patients must not be pregnant and if of child bearing age using adequate contraception
8. Breast feeding
9. Serious psychiatric comorbidity

**Date of first enrolment**

12/08/2013

**Date of final enrolment**

31/12/2016

## **Locations**

**Countries of recruitment**

United Kingdom

England

**Study participating centre**

**Leeds Cancer Centre**

Beckett Street

Leeds

United Kingdom

LS9 7TF

## **Sponsor information**

**Organisation**

Leeds Teaching Hospitals NHS Trust

**ROR**

<https://ror.org/00v4dac24>

## **Funder(s)**

**Funder type**

Hospital/treatment centre

**Funder Name**

The Leeds Teaching Hospitals Charitable Foundation

## **Results and Publications**

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

## IPD sharing plan summary

Available on request

### Study outputs

| Output type                          | Details | Date created | Date added | Peer reviewed? | Patient-facing? |
|--------------------------------------|---------|--------------|------------|----------------|-----------------|
| <a href="#">Results article</a>      |         | 06/09/2018   | 10/07/2023 | Yes            | No              |
| <a href="#">HRA research summary</a> |         |              | 28/06/2023 | No             | No              |