

Hitting the Mark: Introducing state-of-the-art MRI for precision radiotherapy of glioblastoma

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Primary objective: • To demonstrate that the probability for reduced coverage of the recurrence volume by a radiotherapy plan based on a CTVbio, compared to the clinical radiotherapy plan (1.5-cm CTV), is lower than 0.20. Secondary objectives: • To...

Ethical review	Approved WMO
Status	Recruiting
Health condition type	Nervous system neoplasms malignant and unspecified NEC
Study type	Observational non invasive

Summary

ID

NL-OMON55947

Source

ToetsingOnline

Brief title

MOSAIC

Condition

- Nervous system neoplasms malignant and unspecified NEC

Synonym

brain tumor, glioblastoma

Research involving

Human

Sponsors and support

Primary sponsor: Erasmus MC, Universitair Medisch Centrum Rotterdam

Source(s) of monetary or material Support: NWO

Intervention

Keyword: glioblastoma, MRI, precision radiotherapy

Outcome measures

Primary outcome

Pattern of failure and dose to organs at risk by the radiotherapy plan based on the 1.5-cm CTV and the theoretical plan created with the CTVbio.

Secondary outcome

- The differences in tumor recurrence coverage and size between the CTVbio (based on all three aMRI-scans) and CTVs based on individual or other combinations of aMRI, i.e. based on two or one aMRI-scan(s).
- The difference in mean and maximum signal intensity on the individual aMRI-scans between the tumor recurrence site and the contralateral normal-appearing white matter.

Study description

Background summary

One of the fundamentals of glioblastoma management is radiotherapy, where ionizing radiation is aimed towards a specific target area in the brain to inhibit further tumor growth. As these brain tumors are notorious for their extensive tumor infiltration, where tumor grows beyond the tumor that is visible on conventional magnetic resonance imaging (MRI), this target area, defined as the clinical target volume (CTV), consists of the visible tumor plus a 1.5-cm isotropic safety margin. In the majority of cases, this unspecific CTV margin adequately covers tumor infiltration, but inevitably also includes considerable amounts of healthy tissue. Radiation-induced side-effects like headaches, nausea, fatigue and cognitive decline can substantially affect the quality of life for these patients.

An opportunity arises to indirectly visualize tumor infiltration with state-of-the-art advanced MRI (aMRI) techniques, providing additional information on physiology rather than only showing anatomical information

through conventional MRI. A workflow has been developed to create a CTV based on these aMRI scans (CTVbio) rather than an isotropic expansion. With the additional information that aMRI provides, it could be possible to more accurately define what needs to be targeted and thus minimize damage to healthy tissue. In this research, the aim is to assess the potential of integrating aMRI into radiotherapy target delineation for patients with a glioblastoma by comparing the pattern of failure (coverage of first tumor recurrence by the radiotherapy plan) and the expected radiation dose to organs at risk between the CTVbio and the 1.5-cm CTV. It is hypothesized that the CTVbio can result in decreased radiation dose to organs at risk, whilst having similar pattern of failure.

Study objective

Primary objective:

- To demonstrate that the probability for reduced coverage of the recurrence volume by a radiotherapy plan based on a CTVbio, compared to the clinical radiotherapy plan (1.5-cm CTV), is lower than 0.20.

Secondary objectives:

- To illustrate a reduction in dose to organs at risk with a radiotherapy plan based on a conceptual CTVbio compared to the clinical radiotherapy plan (1.5-cm CTV).
- To evaluate the synergistic information that each individual aMRI-scan provides for the identification of tumor infiltration.
- To explore the association between pathophysiological changes on aMRI and future tumor recurrence.

Study design

In this prospective cohort study, the clinical standard MRI session used for radiotherapy planning of glioblastoma patients will be extended with aMRI techniques that assess altered oxygenation, angiogenesis and increased protein concentration. Radiation treatment (and patient follow-up) will occur according to the clinical standard, i.e. using the 1.5-cm CTV for radiotherapy planning. The aMRI-scans will be used to create a theoretical CTVbio and corresponding radiotherapy plan. Pattern-of-failure analysis and assessment of dose to organs at risk will be done to compare the radiotherapy plan based on the 1.5-cm CTV with the (theoretical) radiotherapy plan based on the CTVbio. Additionally, various theoretical CTVs based on different combinations of aMRI-scans are generated to explore the added value of the different aMRI techniques. Lastly, the signal intensities on the aMRI-scans at the site of tumor recurrence are compared with contralateral normal-appearing white matter.

Study burden and risks

The patients have the burden of prolonged scan time (+ 20 minutes, total scan time will be approximately 45 minutes instead of 25 minutes) during their standard radiotherapy planning MRI-scan. The remainder of their clinical care will not be altered: Radiotherapy will be given to these patients based on standard 1.5-cm CTVs. Follow-up will follow the clinical protocol. There will be no personal benefit for the patients in this research project.

Contacts

Public

Erasmus MC, Universitair Medisch Centrum Rotterdam

Doctor Molewaterplein 40
Rotterdam 3015GD
NL

Scientific

Erasmus MC, Universitair Medisch Centrum Rotterdam

Doctor Molewaterplein 40
Rotterdam 3015GD
NL

Trial sites

Listed location countries

Netherlands

Eligibility criteria

Age

Adults (18-64 years)

Elderly (65 years and older)

Inclusion criteria

Written informed consent;

Adults (18 years or older);

Diagnosed with IDH-wildtype glioblastoma, as confirmed by pathology including molecular analysis post resection/biopsy;

Referred to the outpatient clinic of the Dept. of Radiotherapy to undergo standard treatment with radiotherapy (30x2 Gy or 15x2.67 Gy) and scheduled for an MRI for radiotherapy planning.

Exclusion criteria

Contraindication for (3 Tesla) MRI;
Contraindication for use of gadolinium-based contrast agent (e.g. subject having renal deficiency or known allergy);
Referred for treatment of recurrent glioblastoma;
Previous radiotherapy to the brain;
Unable to give informed consent.

Study design

Design

Study type: Observational non invasive

Masking: Open (masking not used)

Control: Uncontrolled

Primary purpose: Diagnostic

Recruitment

NL

Recruitment status: Recruiting

Start date (anticipated): 03-01-2024

Enrollment: 53

Type: Actual

Medical products/devices used

Registration: No

Ethics review

Approved WMO

Date: 06-11-2023

Application type:	First submission
Review commission:	METC Erasmus MC, Universitair Medisch Centrum Rotterdam (Rotterdam)
Approved WMO Date:	13-08-2024
Application type:	Amendment
Review commission:	METC Erasmus MC, Universitair Medisch Centrum Rotterdam (Rotterdam)

Study registrations

Followed up by the following (possibly more current) registration

No registrations found.

Other (possibly less up-to-date) registrations in this register

No registrations found.

In other registers

Register	ID
CCMO	NL84994.078.23