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ORIGINAL ARTICLE

Differences in supratentorial white matter diffusion after radiotherapy – new biomarker of normal brain tissue damage?

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Abstract

Introduction. Therapy-induced injury to normal brain tissue is a concern in the treatment of all types of brain tumours. The purpose of this study was to investigate if magnetic resonance diffusion tensor imaging (DTI) could serve as a potential biomarker for the assessment of radiation-induced long-term white matter injury. Material and methods. DTI- and T1-weighted images of the brain were obtained in 19 former radiotherapy patients [nine men and 10 women diagnosed with astrocytoma (4), pituitary adenoma (6), meningioma (8) and craniopharyngioma (1), average age 57.8 (range 35–71) years]. Average time from radiotherapy to DTI scan was 4.6 (range 2.0-7.1) years. NordicICE software (NIC) was used to calculate apparent diffusion coefficient maps (ADC-maps). The co-registration between T1 images and ADC-maps were done using the auto function in NIC. The co-registration between the T1 images and the patient dose plans were done using the auto function in the treatment planning system Eclipse from Varian. Regions of interest were drawn on the T1-weighted images in NIC based on isocurves from Eclipse. Data was analysed by t-test. Estimates are given with 95% CI. Results. A mean ADC difference of $4.6(0.3;8.9) \times 10^{-5}$ mm²/s, p = 0.03 was found between paired white matter structures with a mean dose difference of 31.4 Gy. Comparing the ADC-values of the areas with highest dose from the paired data (dose > 33 Gy) with normal white matter (dose <5 Gy) resulted in a mean dose difference of 44.1 Gy and a mean ADC difference of $7.87(3.15;12.60) \times 10^{-5}$ mm²/s, p = 0.003. Following results were obtained when looking at differences between white matter mean ADC in average dose levels from 5 to 55 Gy in steps of 10 Gy with normal white matter mean ADC: $5 \text{ Gy}; 1.91(-1.76;5.58) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.29; 15 \text{ Gy}; 5.81(1.53;10.11) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ Gy}; 5.80(2.43;9.18) \times 10^{-5} \text{ mm}^2/\text{s}, p = 0.01; 25 \text{ mm}^2/\text{s}, p = 0.01;$ 10^{-5} mm²/s, p = 0.002; 35 Gy; 5.93(2.89;8.97) × 10^{-5} mm²/s, p = 0.0007; 45 Gy; 4.32(-0.24;8.89) × 10^{-5} mm²/s, p = 0.06; 55 Gy; $-4.04(-14.96;6.89) \times 10^{-5}$ mm²/s, p = 0.39. Conclusion. The results indicate that the structural integrity of white matter, assessed by ADC-values based on DTI, undergoes changes after radiation therapy starting as early as total dose levels between 5 and 15 Gy.

Radiation therapy (RT) is an important treatment modality for both benign and malignant tumours of the central nervous system. The goal of treatment is to deliver a maximum radiation effect to a tumour with minimal effect on normal tissue. Late delayed radiation injury is irreversible but the understanding of its development and the possibility to quantify the amount of injury is still limited [1-3]. Although not fully understood, the mechanisms of the late effects is believed to include vascular damage, white matter injury and coagulation necrosis [3,4]. The use of magnetic resonance imaging (MRI) to characterise late radiation changes is relatively new. Radiation-induced MRI changes, such as T2 signal abnormality, are non-specific and can be seen in a variety of processes including oedema, demyelination and necrosis. Furthermore, central nervous system injury after RT may be present despite a normal appearing MRI [1]. Diffusion-weighted imaging is a promising MRI modality when it comes to describing response to RT [5–8] and damage to normal tissue [9–11]. In diffusion-weighted MRI the MR signal reflect changes in mobility of water molecules in tissue. This mobility, classically called Brownian motion, is due to thermal agitation and is highly influenced by the cellular environment of water [7,12,13]. The structure of the tissue greatly influences whether the diffusion can be characterised as

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isotropic (hindrance or restriction is the same in all directions) or anisotropic (hindrance or restriction is not the same in all directions) [14]. White matter in the brain is organised in large fibre bundles consisting of parallel myelinated fibres which make the tissue anisotropic. Diffusion tensor imaging (DTI) represents a further development of diffusion-weighted imaging measuring both direction and magnitude of water diffusion [15]. DTI is regarded the most sensitive diffusion imaging technique to identify white matter abnormalities and it can detect abnormalities well before structural changes are visible with any other imaging method [9,10].

The purpose of this study was to investigate if DTI could serve as a potential biomarker for the assessment of radiation-induced long-term white matter injury. We hypothesised that in brain tumour patients treated with RT, radiation causes dose dependent progressive loss of structural integrity in white matter.

Material and methods

Patients

Forty-six former patients who had undergone radiotherapy towards the brain in the period 2004–2010 were invited to participate. The study was approved by the local Ethics Committee and the Danish Data Protection Agency. Each patient provided informed written consent prior to inclusion.

MR imaging

MRI data acquisition was performed using a 3T MR scanner (Signa HDx R14M5, GE Healthcare, Milwaukee, WI, USA). The MRI protocol included a T1-weighted axial 3D BRAVO fast spoiled gradient echo (FSPGR) sequence (TR = 10.864 ms, TE = 4.58 ms, FOV = $240 \times 240 \text{ mm}^2$, matrix 352×224 , slice thickness = 1.2 mm, scantime 4:30 minutes) and a diffusion-weighted spin echo echo-planar imaging (SE-EPI) sequence axial 2D DTI (TR = 9000 ms, TE = 105 ms, isotropic voxel size of $2.4 \times 2.4 \times 2.4 \text{ mm}^3$, scantime 5:42 minutes) with 32 gradients and two b-values (0 and 1300 s/mm^2).

This MRI protocol is comparable to the protocol used by Catani et al. when creating their atlas of tractography [16], and was already implemented at our Department of Neurosurgery.

Post processing

Apparent diffusion coefficient maps (ADC-maps) were generated based on the eigen values of the tensor in each voxel using NordicICE (version 2.3.11, NordicImagingLab, Bergen Norway).

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The ADC-maps were co-registered to the morphological 3D BRAVO images using the automatic co-registration in NordicICE. The co-registration was visually inspected before approval. The BRAVO sequences were then co-registered to the patients' dose plans using the auto function in the dose-planning system (Eclipse Registration module, Varian). Again all co-registrations were visually inspected before approval. Three sets of regions of interest (ROIs) were defined on the BRAVO-images in NordicICE based on the isocurves projected on the BRAVO-images in the dose-planning system: 1) Normal appearing white matter in an area with less than 5 Gy; 2) High (dose > 33 Gy) and low (dose < 33 Gy) dose ROIs in paired or parallel white matter structures (Figure 1); 3) ROIs in normal appearing white matter at dose levels 0-10(5), 10-20(15), 20-30(25), 30-40(35), 40-50(45) and 50-60(55) Gy. The ROI size is based on a standard ROI of 66 mm² circle which equals 11 voxels. The standard ROI is then reduced in each case to the largest circular/oblong ROI that fit the white matter area between the relevant dose lines. The standard ROI is placed with centre on the relevant dose level, e.g. 15 Gy and then reduced to fit between the lower (10 Gy) and the upper (20 Gy) dose line limits. In order to get as close to the standard ROI area of 66 mm² as possible the ROIs were positioned in white matter areas on the BRAVO-images where the distance between the dose lines were as big as possible. ROIs were positioned with consideration of avoiding areas with susceptibility artefacts and distortions (e.g. areas near air cavities and compact bone) within the ADC-maps.

Finally the ADC-maps were placed as an overlay on the BRAVO images including the applied ROIs and the ADC data within each ROI were obtained.

Data analysis

Although a quantitative parameter the ADC value should be used with consideration. The ADC value varies among other parameters with the choice of b-values [13,17], scanner vendor [18] and patient age [19–21]. By using the same scanner throughout the study and looking at ADC-differences among the patients instead of absolute values these confounding factors are minimised. ADC also varies in the different white matter areas of the brain [22]. This was accounted for by an analysis of paired/parallel white matter areas exposed to high and low dose of radiation.

Data was analysed by t-test. Estimates are given with 95% confidence intervals and 2-sided p-values.

Results

Nineteen patients agreed to participate [nine men and 10 women, average age 57.8 (range 35-71)



Figure 1. Top row: patient with pituitary adenoma, example of ROI placement in parallel white matter areas. The ROI size is based on a 66 mm² circle, reduced in each case to the largest circular/oblong ROI that fit the white matter area between the relevant dose lines. Bottom row: patient with astrocytoma grade III, example of ROI placement in paired white matter areas. Left column: ADC-map placed as an unsmoothed overlay on T1 weighted images. Right column: Dose plan with absolute dose in steps of 5 Gy placed as an overlay on the T1 weighted images.

years]. Five different diagnoses were represented (Table I). Average time from radiation to DTI MRI scan was 4.6 (range 2.0–7.1) years.

A mean ADC difference of $4.6(0.3;8.9) \times 10^{-5}$ mm²/s, p = 0.03 was found between paired white matter structures with a mean dose difference of 31.4 Gy (Table II). Although not significant the same trend where seen when looking at the maximum and the minimum ADC values (p = 0.06 and p = 0.17, respectively). A mean ADC difference of 7.5(2.6;12.4) × 10⁻⁵ mm²/s, p = 0.005 was found when comparing the mean ADC values of the areas with highest dose from the paired data with mean ADC values of the areas with normal white matter. The mean dose difference was 44.4 Gy. The same trend, although not significant, was observed when comparing the mean ADC values of the areas of

lowest dose from the paired data with the mean ADC values of the areas with normal white matter p = 0.21.

Table III shows the differences between mean ADC values in normal white matter compared to

Table I. Diagnosis and treatment overview of the included patients.

Diagnose	Number of patients	Pre RT surgery/biopsy [Yes/No]	Radiation therapy [Gy/fractions]
Craniopharyngioma	1	Yes	50.4/28
Astrocytoma grade III	3	Yes	59.4/33
Pilocytic astrocytoma	1	Yes	45/25
Pituitary adenoma	6	Yes	45/25
Meningioma	8	Yes	54/30

RT, radiation therapy.

Table II. ADC and dose differences between paired white matter structures at unequal total radiation dose levels.

	High dose level	Low dose level	Mean diff	CI	Р	N
Mean ADC	$79.72 \\ 44.44$	75.13	4.6	0.3;8.9	0.03	18
Mean dose		13.06	31.4	25.9;36.9	0.00	18

Mean diff = High dose level – Low dose level. CI, 95% confidence interval; n, number of patients; p, 2-sided p-value. ADC-units $[10^{-5} \text{ mm}^2/\text{s}]$. Dose-units [Gy].

white matter at different isodose levels. The number of patients is not the same at each isodose level, because of the heterogeneity of the patient group At dose level 35 Gy the patient group is reduced with one meningioma patient and at dose level 45 Gy the patient group is reduced with additional; one astrocytoma, two pituitary adenoma and one craniopharyngiom patient. At dose level 55 Gy the patient group only consists of three astrocytoma and three meningioma patients. According to this the mean ADC in normal white matter differs between the patient groups present at the different dose levels (71.92– 73.30×10^{-5} mm²/s).

The boxplots in Figure 2 shows the relationship between the mean ADC values. There is a significant difference (Table III) between normal white matter mean ADC and normal appearing white matter mean ADC at dose level 15, 25 and 35 Gy of about 8%. Although not statistically significant the trend is the same at dose level 45 Gy. The same pattern is seen when looking at minimum ADC and maximum ADC within the ROIs (data not illustrated).

At dose level 55 Gy the mean ADC seems to drop below mean ADC of normal white tissue (not statistically significant). There is only six patients left in this group; three meningioma patients with mean ADC of 72.78×10^{-5} mm²/s (range 66.68–80. 10×10^{-5} mm²/s) with an increase of 2.83 (-4.69; 10.35)× 10^{-5} mm²/s in mean ADC when comparing with normal ADC for these three patients and three astrocytoma patients with mean ADC of 65.74×10^{-5} mm²/s (range $62.43-69.3 \times 10^{-5}$ mm²/s) with a decrease of $-11.0(-38.1;16.3) \times 10^{-5}$ mm²/s in mean ADC when comparing with normal ADC for these three patients. Again the same pattern is seen when looking at minimum ADC and maximum ADC within the ROIs (data not illustrated).

It was not possible to find comparable paired regions in the white matter structures when looking at the wide span of dose levels from 5 to 55 Gy. However the differences between mean ADC in the paired analysis (Table II) is within the range of the differences in mean ADC found in the dataset of continuous dose levels (Table III).

Discussion

This study shows that ADC values calculated on the basis of DTI are able to detect diffusion differences in normal appearing white tissue as a late effect to radiation therapy.

Although not directly comparable as described in the data analysis section the absolute mean ADC values in this study is within range of values reported by others [20–22]. A mean ADC increase of 8% compared to normal white matter tissue was detectable from total radiation dose levels of 15–45 Gy. This increase in mean ADC corresponds well with previously reported histopathologic findings of postirradiation diffuse white matter injury such as reactive astrocytosis, oedema or demyelination [4,23]. The mean increase in ADC-value seemed rather constant in the interval of 15–45 Gy which could indicate a threshold dose for the observed changes.

At 55 Gy the mean ADC seemed to decrease indicating restricted diffusion. Only data from six patients were obtained at this dose level – three patients with astrocytoma and three patients with meningioma. It seemed that it was the astrocytoma patients who solely contributed to the decrease in ADC. At the dose level of 55 Gy, data was obtained

Table III. Comparison of mean ADC-values in normal white matter and normal appearing white matter at different total radiation dose levels to the brain.

	No	Irradiated white matter					
Number of patients	matter Mean ADC [10 ⁻⁵ mm ² /s]	Dose [Gy]	Mean ADC $[10^{-5} \text{ mm}^2/\text{s}]$	Mean diff [10 ⁻⁵ mm ² /s]	CI [10 ⁻⁵ mm ² /s]	Р	
19	71.92	5	73.83	1.9	-1.8;5.6	0.29	
19	71.92	15	77.74	5.8	1.5;10.1	0.01	
19	71.92	25	77.72	5.8	2.4;9.2	0.002	
18	72.14	35	78.07	5.9	2.9;9.0	0.0007	
14	72.89	45	77.21	4.3	-0.2;8.9	0.06	
6	73.30	55	69.26	-4.0	-15.0;6.9	0.39	

Mean diff=mean ADC at dose level (\times) – mean ADC normal white matter. CI, 95% confidence interval; p, 2-sided p-value.



Figure 2. Boxplot of mean ADC ($\times 10^{-5}$ mm²/s) for normal white matter (radiation total dose 0 Gy) and the irradiated normal appearing white matter (radiation total dose 5–55 Gy).

within the volume defined as target in the radiation dose plan. Although the ROIs were placed in normal appearing white matter, there is relatively high risk of infiltrative tumour growth in astrocytoma patients. Such hypercellularity would explain the observed decrease in mean ADC [5,7,24]. One of the astrocytoma patients had a mean ADC in normal appearing white matter in an area with less than 5 Gy (normal tissue) of 88.5×10^{-5} mm²/s and a mean ADC of 65.5×10^{-5} mm²/s in normal appearing white matter at dose 55 Gy (a ADC reduction of 26%). The patient experienced clinical relapse three months after the DTI scan. The relapse was considered to be a further progression of the tumour from astrocytoma grade III to glioblastoma multiforme and was diagnosed 7.5 years after RT. Another of the patients had a mean ADC in normal tissue of 70.5×10^{-5} mm²/s and a mean ADC of 62.4×10^{-5} mm²/s in normal appearing white matter at dose 55 Gy (a ADC reduction of 11%). The patient experienced clinical relapse 1.5 years after the DTI scan. This was believed to be a further development of the astrocytoma grade III and was diagnosed six years after RT. The last astrocytoma patient was diagnosed and received RT two years prior to the DTI scan. This patient had a mean ADC in normal tissue of 71.0×10^{-5} mm²/s and a mean ADC of 69.3×10^{-5} mm²/s in normal appearing white matter at dose 55 Gy (a ADC reduction of 2). This indicates that DTI could have the potential as an early predictor for tumour recurrence. This should be explored further in future studies.

In the study by Nagesh et al. [9] a nine gradient DTI with b-values (0 and 1000 s/mm²) at 1.5T was

used to determine acute and sub-acute radiationinduced changes in normal appearing white matter fibres (genu and splenium of the corpus callosum) in patients with cerebral tumours. This study demonstrated that the mean ADC increased linearly and significantly with time to a point of 18% 45 weeks after start of RT which was the end of follow-up. In the present study the late effects to white matter mean ADC were in the range of 8–10% with an average time from radiation to the DTI scan of 4.6 years. This indicates that the radioresponse of white matter tissue in the brain is a dynamic process which is in good accordance with other studies [25].

The changes of 8–10% in ADC found in this study are relatively small. This makes the results rather sensitive to noise. Although the SS-EPI technique has a high signal to noise ratio (SNR) per unit of scanning time, the use of high b-values in diffusion-weighted imaging makes the technique more SNR-limited than conventional MR imaging such T2-weighted imaging. To boost the SNR it is common practice to repeat acquisition of the same DWIs (increasing NEX) or as in the current study apply more gradients to the DTI. It has been shown that applying 32 gradient directions at 1.5 T makes a robust estimation of mean ADC which is reproducible within 2.6% [26].

Another thing to consider is the precision at which the different image sequences were co-registered. For each patient the DTI and the BRAVO sequences were obtained in the same scan session. This reduces the risk of co-registration inaccuracy considerably. No mismatch was registered at the visual inspection prior to co-registration approval. The BRAVO sequence was also co-registered to the CT-based dose plan. The mismatch was visually assessed to be in the range of 1-2 mm. To reduce the influence of such mismatch the ROIs were attempted placed in areas with low dose gradients.

The use of DTI is a promising technique to assess radiation-induced white matter injury. Besides ADCmaps at a higher resolution than common DWI, DTI provides a wide range of diffusion-related parameters such as fractional anisotropy (FA-maps) which provides information about the main diffusion direction of each voxel [7,11,15] and tensor eigen values (λ -maps) which can help differentiate between white matter axonal injury or demyelination [9]. In order to interpret these parameters a Pre-RT DTI scan is needed, as the interstructural white matter variation in the parameters is relatively high [22].

In conclusion ADC-values based on DTI seem to have the potential to serve as a biomarker for the assessment of radiation-induced long-term white matter injury. Further studies are needed to evaluate the clinical value of the method.

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