Longitudinal MRI Study After Carbon Ion and Photon Irradiation: Shorter Latency Time for Myelopathy is Not Associated With Differential Morphological Changes

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Research

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Abstract

Background

Radiation-induced myelopathy is a severe and irreversible complication that occurs after a long symptom-free latency time if the spinal cord was exposed to a significant irradiation dose during tumor treatment. As carbon ions are increasingly investigated for tumor treatment in clinical trials, their effect on normal tissue needs further investigation to assure safety of patient treatments. Magnetic resonance imaging (MRI)-visible morphological alterations could serve as predictive markers for medicinal interventions to avoid severe side effects. Thus, MRI-visible morphological alterations in the rat spinal cord after high dose photon and carbon ion irradiation and their latency times were investigated.

Methods

Rats whose spinal cords were irradiated with iso-effective high photon (n = 8) or carbon ion (n = 8) doses as well as sham-treated control animals (n=6) underwent frequent MRI measurements until they developed radiation-induced myelopathy (paresis II). MR images were analyzed for morphological alterations and animals were regularly tested for neurological deficits. In addition, histological analysis was performed of animals suffering from paresis II compared to controls.

Results

For both beam modalities, first morphological alterations occurred outside the spinal cord (bone marrow conversion, contrast agent accumulation in the musculature ventral and dorsal to the spinal cord) followed by morphological alterations inside the spinal cord (edema, syrinx, contrast agent accumulation) and eventually neurological alterations (paresis I and II). Latency times were significantly shorter after carbon ions as compared to photon irradiation.

Conclusions

Irradiation of the rat spinal cord with photon or carbon ion doses that lead to 100% myelopathy induced a comparable fixed sequence of MRI-visible morphological alterations and neurological distortions. However, at least in the animal model used in this study, the observed MRI-visible morphological alterations in the spinal cord are not suited as predictive markers to identify animals that will develop myelopathy as the time between MRI-visible alterations and the occurrence of myelopathy is too short to intervene with protective or mitigative drugs.

Introduction

Carbon ions are being increasingly investigated for radiotherapy in clinical trials (1), as they exhibit physical advantages, such as a finite range in tissue leading to a highly conformal dose distributions, as well as biological advantages, associated with their increased linear energy transfer (LET) resulting in a higher relative biological effectiveness (RBE) in the target volume (2). This makes them a promising
candidate for irradiation of radioresistant and critically located tumors. However, despite the increased accuracy of carbon ions, the exposure of tumor surrounding normal tissue to high irradiation doses cannot be avoided. To assure patients’ safety, it is therefore of utmost importance to investigate the biological effect of carbon ions on normal tissue complication in comparison to conventionally used photons.

Late delayed injury like irradiation-induced myelopathy is a severe and irreversible complication that may occur 6 months to 2 years after irradiation (3). Affected patients suffer from numbness, dysesthesias, and incontinence. The neurological deficit develops progressively eventually leading up to quadriplegia in at least 50% of patients (4). On a histological level, irradiation-induced myelopathy is characterized by demyelination, vascular abnormalities, and necrosis in the white matter (5). However, the primary target cell whose death leads to the irradiation-induced necrosis is not yet identified and the hypotheses that the vasculature and endothelial cells are the main cause for the development of late effects is still being investigated (5, 6).

The rat cervical spinal cord is a well-established model for the investigation of irradiation effects on normal tissue (7–11). The biological endpoint irradiation-induced myelopathy can be well detected as paresis of the animals’ forelimbs. The morphological changes accompanying irradiation-induced myelopathy at different pre-designated time points after irradiation have been investigated on a histological level for single exposures of the thoracic and lumbar cord (12–14). However, the results of these studies might not be representative for fractionated treatments used in clinical routine.

Magnetic resonance imaging (MRI) allows repeated and non-invasive examination of morphological tissue alterations that precede neurological disorders. In this way, disease progression may be monitored individually per animal and findings may be transferred to patients. So far, however, only few MRI studies have been performed on the pathogenesis of irradiation-induced myelopathy in the rat spinal cord (15, 16).

Regarding the use of fractionated carbon ion irradiations of the cervical spinal cord, alterations preceding neurological disorders have not yet been investigated. A better understanding of the processes preceding clinical symptoms could help to find an adequate time window for medical interventions to prevent or mitigate radiation-induced myelopathy (17–20).

In this study, the longitudinal development of morphological and neurological alterations preceding radiation-induced myelopathy was observed and analyzed qualitatively by MRI after fractionated iso-effective irradiations with photons or carbon ions, respectively.

**Materials And Methods**

**Animals**
A total of 22 young adult female (242 ± 34 g) Sprague-Dawley rats (Charles River, Sulzfeld, Germany) were used. Animals were randomly distributed into three groups: (I) 8 rats were irradiated with carbon ions, (II) 8 rats were irradiated with photons, and (III) 6 rats served as non-irradiated controls. Animals were kept under the standard laboratory conditions at the German Cancer Research Center (DKFZ, Heidelberg, Germany). All experiments were approved by the governmental review committee on animal care.

**Irradiation setup**

The rat cervical spinal cord was irradiated either with carbon ions or photons. For irradiation, rats were anesthetized with a mixture of 4 Vol% Sevoflurane (Abbott, Wiesbaden, Germany) and oxygen at 2 l/min and animals were irradiated in a hanging position in a customized holding device (7). The rat spinal cord was irradiated from ventral to dorsal with a horizontal beam (field size of 10 × 15 mm²) that covered the cervical segments C1-C6 with C3 being in the center and C1 and C6 being located at the field edge.

Carbon ion irradiations were performed at the Heidelberg Ion-Beam Therapy Center (HIT, Heidelberg, Germany). The cervical spinal cord was positioned in the middle of a 1 cm spread-out Bragg peak (SOBP) with a dose-averaged LET of 91 keV/µm (80–104 keV/µm) and was irradiated with a total dose of 23 Gy given in 6 fractions (3.83 Gy/fraction) on consecutive days. The dose was selected to reach 100% complication probability (8). Photon irradiations (6 MeV, Artiste, Siemens, Erlangen) were performed with an iso-effective total dose of 61 Gy given in 6 fractions (10.167 Gy/fraction) on consecutive days to also reach 100% complication probability (8). The iso-effective dose was determined based on the RBE for the endpoint radiation-induced myelopathy measured in a previous study (8).

**MRI follow-up**

Animals underwent MRI before irradiation and were then imaged on a monthly basis to detect morphological alterations. As soon as an alteration occurred, the respective rat was monitored in shorter time intervals (Fig. 1).

MRI measurements were performed on a 1.5 T whole-body MR-scanner (Symphony, Siemens, Erlangen, Germany) using an in-house built small animal coil. Rats were anesthetized by inhalation anesthesia (Isoflurane, Baxter, 2.5 Vol% in 1.2 l/ min oxygen). During each MRI session, sagittal and axial T2-weighted images (4000/109 ms [repetition time/ echo time], voxel size 0.3 × 0.3 × 1.0 mm) were acquired per animal to assess morphological changes such as edema formation. Additionally, T1-weighted images (600/14 ms [repetition time/ echo time], voxel size 0.2 × 0.2 × 1.0 mm) were recorded before and after contrast agent injection to examine the onset of the blood spinal cord barrier (BSCB) disruption. The contrast agent (0.2 mmol/kg Magnevist®, Bayer, in 0.9% NaCl) was injected intravenously as a bolus via the tail vein and animals were imaged 5 min after injection.

**Follow up and endpoints**
After irradiation, rats were checked weekly for neurological alterations, weight, and general condition. The final biological endpoint was radiation-induced myelopathy (paresis II) within 300 days meaning that both forelimbs showed signs of paralysis (7). As soon as an animal reached paresis II, a final MRI measurement was performed and the animal was sacrificed. Before animals reached the final endpoint, they showed various morphological alterations (see “Image evaluation”) and paresis I (at least one forelimb showed signs of neurological dysfunction (7)). The latency times, defined as the time between the day of the first fraction of irradiation and the occurrence of the respective morphological and neurological alterations after irradiation, were recorded for each animal.

Image evaluation

Two blinded readers (T.W., M.S.) performed qualitative visual inspections of the MR-images with a consensus reading of pathologies to determine the latency times for the morphological alterations. For this, the axial and sagittal T2-weighted images as well as the sagittal T1-weighted images (native and post contrast agent) were used.

Native T1-weighted images were used to identify bone marrow conversion. To examine the onset of loss of BSCB integrity, native and enhanced T1-weighted images were compared. The T2-weighted images were used to identify edema and the dilatation of the canalis centralis (syrinx). All latency times, i.e. the time between the first fraction and the occurrence of MR visible morphological alterations, were recorded.

Histology

When animals reached paresis II they were sacrificed by an overdose of i.p. injected Xylazine/Ketamine (100 µl/kg each). Animals were perfused with a mixture of 4% paraformaldehyde (PFA) in phosphate buffered saline before the cervical spinal cord C1-C6 were dissected and fixed in PFA. After dehydration in ethanol the segments C1-C6 were embedded in paraffin with an axial orientation.

All histological stainings were performed on 9 µm thick paraffin sections that were deparaffinized and rehydrated before staining. For all immunohistochemistry stainings antigen retrieval in sodium citrate buffer (pH 6) was performed for 30 min at 95 °C to unmask antigen sites. At the end, sections were dehydrated and mounted in Eukitt.

For qualitative examination of the extent of demyelination, paraffin sections were stained with Luxol fast blue which binds to the lipoproteins of myelin (21) in combination with hemalaun/eosin (H&E). Sections were incubated in 0.1% Luxol blue at 50 °C over night, and washed in 95% ethanol, and differentiated in 0.05% lithium carbonat and 70% ethanol the next day. Afterwards, sections were counterstained with hematoxylin and eosin.

To study the degree of blood vessel permeability, extravagated endogenous serum albumin was immunohistochemically visualized. Endogenous peroxidase activity was blocked with 3% H$_2$O$_2$. After antigen retrieval, sections were incubated overnight at 4 °C with the primary antibody against albumin
(Acris, 1:6000 diluted in 3% bovine serum albumin in TBS-T) followed by incubation with the secondary antibody (Abcam, 1:500, horse raddish peroxidase) for 30 min at room temperature. 3,3’-diaminobenzidine was used as chromogen (Vector Laboratories, Burlingame, CA, US). In the end, sections were counterstained with 1% Nissl in sodium acetate buffer.

To assess the degradation of the blood spinal cord barrier, sections were stained for the endothelial barrier antigen (EBA), a protein that is exclusively expressed on the luminal surface of non-fenestrated central nervous system blood vessels (22), in combination with the endothelial marker CD34. Autofluorescence was quenched by 30 min incubation with 0.1% Sudan Black B (Sigma-Aldrich®, Missouri, US). After blocking in 10% normal goat serum, sections were incubated with the primary antibodies monoclonal mouse anti-EBA SMI 71 (1:500, Biolegend, San Diego, Ca, US) and recombinant rabbit anti-CD34 antibody (1:1000, abcam, Cambridge, UK) in 3% bovine serum albumin in TBS-T at 4 °C over night. For detection, sections were incubated with the fluorescent-coupled secondary antibodies goat anti-rabbit Alexa Flour 555 (1:1000, Molecular Probes®, Invitrogen AG, Carlsbad, USA) and goat anti-mouse Alexa Fluor 488 (1:1000, Molecular Probes®) at room temperature for 30 min each. Sections were counter stained with DAPI and mounted with Fluoromount-G.

Statistical analyses

The difference in the latency times per morphological and neurological alterations between the irradiation modalities was tested by a two-sided student t-test with a significance level of 0.05. The longitudinal difference of the latency times within one irradiation modality was tested by one way repeated measure ANOVA. All statistical testing was performed in SigmaPlot 14.0 (Systat Software Inc., San Jose, CA, USA).

Results

Skin alterations

Irradiation procedures as well as MRI measurements were well tolerated by all animals. None of the animals died due to acute radiation toxicity or contrast agent intolerance. Acute radiation toxicity developed within the first four weeks after photon treatment and included a slight or complete hair loss as well as moist desquamation restricted to a small focal region at the ventral and dorsal side of the rats’ throat. Skin alterations were less severe at the dorsal than at the ventral side. Due to the narrow SOBP, no skin alterations were observed after carbon ion irradiation.

Latency times

The latency times detected for the different MR-visible morphological alterations and neurological distortions after irradiation with photons or carbon ions are summarized in Table 1 and visualized in Fig. 2.
Table 1
Mean latency times (± standard deviation) of morphological alterations and neurological distortions after photon and carbon ion irradiation.

| Time of onset of morphological and/or neurological alterations [days after irradiation] | BMC  | CA v | CA d | Edema | CA    | Syrinx | P I     | P II    |
|-----------------------------------------------|------|------|------|-------|-------|--------|---------|---------|
| Photons                                      | 108 ±47 | 107 ±29 | 149 ±32 | 194 ±17 | 198 ±18² | 206 ±23¹ | 199 ±22¹ | 212 ±21 |
| ¹²C-ions                                      | 84 ±24 | 88 ±18 | 81 ±13 | 122 ±10 | 125 ±10³ | 124 ±5  | 129 ±7  | 137 ±11 |

BMC = bone marrow conversion, CA v/d = contrast agent enhancement in the ventral/ dorsal musculature outside the spinal cord, respectively, CA = contrast agent enhancement inside the spinal cord, P I (paresis I) = at least one forelimb showed signs of neurological dysfunction, P II (paresis II) = both forelimbs showed signs of paralysis. Bold printed latency times are significantly different (p < 0.001) between photon and carbon ion irradiated animals.

1) 7 out of 8 rats
2) 3 out of 7 animals showed CA without paresis I
3) 5 out of 8 animals showed CA without paresis II

The first irradiation-induced morphological alterations that occurred were located outside the spinal cord (Fig. 3). Bone marrow conversion (BMC) within the cervical vertebral bodies C1-C4 appeared as hyperintense signal in the native T1-weighted MR images. Next, contrast agent started to leak through the vasculature and accumulated in the ventral and dorsal musculature surrounding the cervical spinal cord. While the onset of contrast agent accumulation in the ventral muscle was rather concurrent to the first signs of BMC per modality, the contrast agent accumulation was significantly delayed in the dorsal muscle in photon irradiated animals as compared to carbon ion irradiated animals. The contrast agent accumulated to a greater extend after photon than after carbon ion irradiations.

In the longitudinal course, morphological alterations such as the presence of edema in the spinal cord, a dilatation of the canalis centralis (syrinx), and accumulation of contrast agent alongside the spinal cord were observed concurrently but significantly later than for the spinal cord external alterations in both irradiation modalities (Table 1, Fig. 2, Fig. 4). Out of the seven photon-irradiated animals that showed paresis I, three showed contrast agent accumulation in the spinal cord before signs of paresis. For carbon ion irradiations, the same morphological alterations were observed (Fig. 4), but the latency times were significantly shorter compared to photon irradiations. In five out of eight animals, contrast agent accumulation in the spinal cord was detected before paresis I had developed.

Next, animals developed neurological deficits that rapidly impaired to paresis I and finally to the endpoint paresis II. One animal of the photon treated group went so rapidly to paresis II that paresis I could not be detected separately. Three out of seven animals developed paresis I prior to a syrinx. All carbon ion irradiated animals showed edema, syrinx and contrast agent accumulation along the spinal cord when
they reached the endpoint paresis II. T2-weighted images revealed that at this point edema and syrinx had spread along the irradiated part of C1-C6.

The biological endpoint paresis II occurred at 212 ± 21 days after photon and 137 ± 11 days after carbon ion irradiation. At these time points, the above described morphological changes had intensified. Edema and syrinx had spread further in cranial and caudal directions outside the irradiation field.

Overall, the sequence of morphological and neurological alterations was the same after photon and carbon ion irradiation. However, after the time point of dorsal accumulation of contrast agent, the latency times for the endpoints were significantly shorter for carbon ions than for photons. There was a large spread in latency times between the individual animals per irradiation group.

The mean time between the first morphological alteration (edema) and the onset of paresis I was 4 ± 6 days after photon irradiation and 7 ± 5 days after carbon ion irradiation.

Histology

Histological examination of animals with paresis II exhibited comparable tissue damages after photon and carbon ion irradiation.

Luxol fast blue staining revealed focal demyelination and severe necrosis in the *S. alba*. Focal necroses were located in the right and left *F. lateralis* and the *F. posterior* (Fig. 5A-D). Necrosis was more severe after carbon ion irradiation than after photons. Local bleedings occurred in the *S. grisea* of affected segments. The overall structure remained visually intact.

The extravasation of endogenous albumin, representing the breakdown of the BSCB, was extensive and comparable between photon and carbon ion irradiated spinal cords (Fig. 5E, G). All segments C1-C6 were affected even though the outer segments showed mostly albumin extravasation in the posterior part of the white matter and around the *canalis centralis*. No albumin had extravasated in the age-matched control animals (Fig. 5F, H).

Segments with focal demyelination exhibited hardly any intact vessels as EBA expression was extensively decreased in both, the *S. alba* and *S. grisea*, in those segments of irradiated animals (Fig. 5I-L). CD34 revealed vessel dilatation in the grey matter of irradiated animals.

Discussion

During radiotherapy, tumor surrounding normal tissue is often partially or fully exposed to high doses, which can lead to irreversible side effects. For the spinal cord, radiation-induced myelopathy represents a delayed late effect, which is characterized by a long symptom free latency period of up to 2 years followed by a sudden occurrence of neurological deficits in the patient (3). As carbon ions are increasingly investigated in clinical trials with focus on brain and head and neck tumors, their effect on normal tissue of the central nervous system needs to be explored. For this, we performed a longitudinal
MRI study to characterize the temporal development of irradiation-induced myelopathy in the rat during the symptom free latency period after fractionated iso-effective irradiation with photons or carbon ions and to identify MR-visible morphological alterations that could serve as predictive signs of late radiation injury.

**Acute reactions**

Acute radiation toxicity in the form of slight or complete hair loss and moist desquamation of the ventral and dorsal side of the rats’ throat only occurred after photon but not after carbon ion irradiations. Due to the narrow SOBP, skin was physically spared and no reactions were observed after carbon ions, showing the advantage of the high dose conformity that can be reached with carbon ions. This is similar in patients and may be important for their quality of life (23). Beyond acute reactions, the exposure of smaller volumes with carbon ions may be of importance for tissues exhibiting a distinct volume effect (e.g. brain). As normal tissue tolerance is entangled with the volume effect, conformal irradiations are generally of advantage.

**Fixed sequence of MR-visible morphological alteration and neurological distortions**

Paresis II was preceded by a fixed sequence of morphological alterations in MRI that was identical after iso-effective irradiation with photons and carbon ions. First morphological alterations occurred in the structures surrounding the spinal cord (BMC, CA v/d). Bone marrow conversion is observed due to fatty replacement of the marrow after irradiation (24) and it is non-reversible above a threshold of approximately 30–40 Gy (25). The longer latency time until contrast agent accumulated in the musculature dorsal to the spinal cord after photons but not after carbon ions underlines the independence of carbon ion effectiveness of tissue type (26). Significantly later than the external alterations, morphological changes appeared inside the spinal cord, including edema, syrinx and the breakdown of the BSCB. The latter was indicated by contrast agent accumulation in the spinal cord and by the detection of extravasated endogenous albumin in histological stainings. The observed alterations are in accordance with findings in patients that had developed myelopathy (27, 28).

In our study, the only difference between the two irradiation modalities was found to be their latency times: the sequence of morphological and neurological alterations started significantly earlier after carbon ions than after photons. A shorter latency time for irradiation induced myelopathy was also found in other studies (7, 8, 13, 29, 30). High LET radiation such as carbon ions, induces clustered DNA damage and reduces the cell’s repair capacity significantly (31), which may be the main cause for the shorter latency time. Furthermore, the latency time for irradiation induced myelopathy has also been shown to decrease with increasing LET, fractionation number, and dose (32).

Interestingly, the time interval between first MRI-visible morphological alterations (edema) and paresis I and II was comparable for both irradiation modalities. This is also in accordance with the histological results, which showed comparable tissue and vascular degradation after irradiation with both modalities. These findings indicate that the progresses that take place on a morphological and functional level
visible in MRI as well as in histology are comparable after low and high LET irradiation. Similar histological results were reported by Okada *et al.* after single fraction irradiation of murine thoracic and lumbar spinal cord with carbon ions and photons (13).

**Predictive markers for medicinal interventions**

From clinical experience, radiation-induced myelopathy develops suddenly after a long symptom-free period and once neurological deficits appear, these are usually irreversible. Therefore, it is of great interest to non-invasively determine preceding morphological changes that could serve as predictive markers to potentially initiate medicinal interventions that might prevent or mitigate clinical symptoms.

In this regard, the vascular hypothesis assumes that radiation-induced vascular damage and subsequent ischemia leads to white matter necrosis (5). The contrast agent accumulation in the spinal cord, as measured by MRI, marks the breakdown of the BSCB of the vasculature, and hence, would be a promising candidate for such a predictive marker. In support of this, we found contrast agent accumulation in the spinal cord before paresis I in half of the animals, suggesting that the breakdown of the BSCB is a prerequisite for the development of neurological symptoms. However, the development of myelopathy is rapid and the mean time between the first detection of an edema and contrast agent accumulation and the onset of paresis I and II was less than one week. Unfortunately, this time window is too narrow for successful medicinal intervention. In contrast, the CA accumulation in the ventral musculature of the spinal cord was observed significantly earlier than paresis I (94 ± 37 days earlier for photon irradiation and 42 ± 18 days earlier for carbon ion irradiation). Whether this can be regarded as an early indication for the development of myelopathy or whether it resembles an independent reaction of the musculature’s vasculature requires further experiments with lower irradiation doses.

Nevertheless, with regard to the low sensitivity of the used MRI scanner, investigation of the molecular pathways prior to the MR visible alterations appear necessary to identify starting points for medicinal interventions. As an example, administration of ACE-inhibitor after irradiation has been reported to mitigate radiation-induced myelopathy in healthy central nervous system tissue (17).

**Conclusion**

The development of myelopathy is free of symptoms over a long time, but then occurs rather rapidly and irreversible. This observational study gives first insights into the time course of morphological alterations in MRI preceding radiation-induced myelopathy. The modality-independent fixed sequence of morphological and neurological alterations has a significantly shorter latency time after carbon ion than after photon irradiation. However, at least in the animal model used in this study, the observed MR-visible morphological alterations in the spinal cord are not suited as predictive markers to identify animals that will develop myelopathy as the time between MR-visible alterations and the occurrence of myelopathy is too short to intervene with protective or mitigative drugs.
Abbreviations

\(^{12}\)C-ions – carbon ions

BSCB – blood spinal cord barrier,

CA – contrast agent,

CA d – contrast agent accumulation in the dorsal musculature of the spinal cord,

CA v – contrast agent accumulation in the ventral musculature of the spinal cord,

EBA – endothelial barrier antigen,

H&E – hemalaun/eosin,

LET – linear energy transfer,

MRI – magnetic resonance imaging,

P I – paresis I,

P II – paresis II,

RBE – relative biological effectiveness

SOBP – spread-out Bragg peak

Declarations

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Consent for publication

Not applicable.

Ethics approval and consent to participate

All experiments were approved by the governmental review committee on animal care (G62/08 and G117/13).
Availability of data and materials

The datasets analyzed during the current study are available from the corresponding author on reasonable request.

Competing interest

JD was financially supported by CRI -The Clinical Research Institute GmbH, View Ray Inc., Accuray International, Accuray Incorporated, RaySearch Laboratories AB, Vision RT limited, Merck Serono GmbH, Astellas Pharma GmbH, Astra Zeneca GmbH, Solution Akademie GmbH, Ergomed PLC Surrey Research Park, Siemens Healthcare GmbH, IQVIA RDS GmbH, Pharmaceutical Research Associates GmbH, Boehringer Ingelheim Pharma GmbH Co, PTW-Freiburg Dr. Pychlau GmbH, and Nanobiotix A.A.

All authors declare that they have no competing interests.

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Author's contributions

TW was involved in the study design, data interpretation and revising of the manuscript, and performed qualitative visual inspections of the MR-images.

ALB analyzed the data, performed immunohistological stainings, and wrote the manuscript.

CG was involved in the study design, establishment of the MR measurement set-up, data interpretation, as well as in revising the manuscript.

IB was involved in the establishment of the MR measurement setting.

MJ was involved in the establishment of the MR measurement setting.

PP initiated the project and was involved in the study design, data interpretation, as well as in revising the manuscript.

JD initiated the project and was involved in the study design, as well as in the translational assessment of the data.

CPK initiated the project and was involved in the study design. He performed the dosimetry, treatment planning, and was involved in data interpretation and revising the manuscript.

MS was involved in the study design, organized, and performed all irradiations and MR imaging as well as the animal monitoring and tissue dissection during the study. Additionally, she performed qualitative
visual inspections of the MR-images, established protocols for immunhistological stainings, and was involved in writing and revising the manuscript.

All authors read and approved the final manuscript.

References

1. Lazar AA, Schulte R, Faddegon B, Blakely EA, Roach M. Clinical trials involving carbon-ion radiation therapy and the path forward. Cancer. 2018;124(23):4467-76.
2. Allen C, Borak TB, Tsujii H, Nickoloff JA. Heavy charged particle radiobiology: using enhanced biological effectiveness and improved beam focusing to advance cancer therapy. Mutat Res. 2011;711(1-2):150-7.
3. Schultheiss TE, Kun LE, Ang KK, Stephens LC. Radiation response to the central nervous system. Int J Radiation Oncology Biol Phys. 1995;31(5):1093-112.
4. Dropcho EJ. Neurotoxicity of radiation therapy. Neurol Clin. 2010;28(1):217-34.
5. Tofilon PJ, Fike JR. The Radioresponse of the Central Nervous System: A Dynamic Process. Rad Res. 2000;153(4):357-70.
6. Van der Kogel AJ. Radiation-induced damage in the central nervous system: An interpretation of target cell responses. British journal of cancer. 1986;7:207-17.
7. Debus J, Scholz M, Haberer T, Peschke P, Jäkel O, Karger CP, et al. Radiation Tolerance of the Rat Spinal Cord after Single and Split Doses of Photons and Carbon Ions. Rad Res. 2003;160(5):536-42.
8. Karger CP, Peschke P, Sanchez-Brandelik R, Scholz M, Debus J. Radiation tolerance of the rat spinal cord after 6 and 18 fractions of photons and carbon ions: experimental results and clinical implications. International journal of radiation oncology, biology, physics. 2006;66(5):1488-97.
9. Van der Kogel AJ, Barendsen GW. Late effects of spinal cord irradiation with 300 kV X rays and 15 MeV neutrons. Br J Radiol. 1974;47(559):393-8.
10. Saager M, Glowa C, Peschke P, Brons S, Scholz M, Huber PE, et al. Carbon ion irradiation of the rat spinal cord: dependence of the relative biological effectiveness on linear energy transfer. International journal of radiation oncology, biology, physics. 2014;90(1):63-70.
11. Saager M, Peschke P, Welzel T, Huang L, Brons S, Grun R, et al. Late normal tissue response in the rat spinal cord after carbon ion irradiation. Radiation oncology. 2018;13(1):5.
12. Okada S, Okeda R, Matsushita S, Kawano A. Immunohistochemical changes of the blood-brain barrier in rat spinal cord after heavy-ion irradiation. Neuropathology. 1998;18:188-98.
13. Okada S, Okeda R, Matsushita S, Kawano A. Histopathological and Morphometric Study of the Late Effects of Heavy-Ion Irradiation on the Spinal Cord of the Rat. Rad Res. 1998;150:304-15.
14. Tsao MN, Li Y-Q, Lu G, Xu Y, Wong CS. Upregulation of Vascular Endothelial Growth Factor is associated with Radiation-Induced Blood-Spinal Cord Barrier Breakdown. Journal of Neuropathology and Experimental Neurology. 1999;58(10):1051-60.
15. Philippens ME, Gambarota G, Pikkemaat JA, Peeters WJ, van der Kogel AJ, Heerschap A. Characterization of late radiation effects in the rat thoracolumbar spinal cord by MR imaging using USPIO. Magma. 2004;17(3-6):303-12.

16. Philippens ME, Gambarota G, Van der Kogel AJ, Heerschap A. Radiation Effects in the Rat Spinal Cord: Evaluation with Apparent Diffusion Coefficient versus T2 at Serial MR Imaging. Radiology. 2009;250(2):387-97.

17. Saager M, Hahn EW, Peschke P, Brons S, Huber PE, Debus J, et al. Ramipril reduces incidence and prolongates latency time of radiation-induced rat myelopathy after photon and carbon ion irradiation. J Radiat Res. 2020.

18. Singh VK, Fatanmi OO, Wise SY, Newman VL, Romaine PL, Seed TM. The Potentiation of the Radioprotective Efficacy of Two Medical Countermeasures, Gamma-Tocotrienol and Amifostine, by a Combination Prophylactic Modality. Radiat Prot Dosimetry. 2016;172(1-3):302-10.

19. Sridharan V, Tripathi P, Sharma S, Corry PM, Moros EG, Singh A, et al. Effects of late administration of pentoxifylline and tocotrienols in an image-guided rat model of localized heart irradiation. PloS one. 2013;8(7):e68762.

20. van der Veen SJ, Ghobadi G, de Boer RA, Faber H, Cannon MV, Nagle PW, et al. ACE inhibition attenuates radiation-induced cardiopulmonary damage. Radiotherapy and oncology : journal of the European Society for Therapeutic Radiology and Oncology. 2015;114(1):96-103.

21. Atkinson S, Li Y-Q, Wong CS. Changes in oligodendrocytes and myelin gene expression after radiation in the rodent spinal cord. International Journal of Radiation Oncology*Biology*Physics. 2003;57(4):1093-100.

22. Sternberger N, Sternberger LA, Kies MW, Shear CR. Cell surface endothelial proteins altered in experimental allergic encephalomyelitis. Journal of Neuroimmunology. 1989;21:241-8.

23. Dormand E-L, Banwell PE, Goodacre TE. Radiotherapy and wound healing. Int Wound J. 2005;2(2):112-27.

24. Vogler JB, Murphy WA. Bone marrow imaging. Radiotherapy and oncology : journal of the European Society for Therapeutic Radiology and Oncology. 1988;168(3):679-93.

25. Daldrup-Link HE, Henning T, Link TM. MR imaging of therapy-induced changes of bone marrow. European radiology. 2007;17(3):743-61.

26. Glowa C, Karger CP, Brons S, Zhao D, Mason RP, Huber PE, et al. Carbon ion radiotherapy decreases the impact of tumor heterogeneity on radiation response in experimental prostate tumors. Cancer Lett. 2016;378(2):97-103.

27. Alfonso ER, De Gregorio MA, Mateo P, Escó R, Bascón N, Morales F, et al. Radiation myelopathy in over-irradiated patients: MR imaging findings. European radiology. 1997;7(3):400-4.

28. Mahta A, Borys E, Kanakamedala MR, Kesari S. Radiation induced myelopathy in a patient with tongue cancer: a case report. Acta oncoligica. 2012;51(3):409-11.

29. Okada S, Okeda R. Pathology of radiation myelopathy. Neuropathology. 2001;21:247-65.
30. Leith JT, McDonald M, Powers-Risius P, Bliven SF, Howard J. Response of Rat Spinal Cord to Single and Fractionated Doses of Accelerated Heavy Ions. Rad Res. 1982;89:176-93.

31. Georgakilas AG, O'Neill P, Stewart RD. Induction and repair of clustered DNA lesions: what do we know so far? Radiation research. 2013;180(1):100-9.

32. Saager M, Glowa C, Peschke P, Brons S, Grun R, Scholz M, et al. Fractionated carbon ion irradiations of the rat spinal cord: comparison of the relative biological effectiveness with predictions of the local effect model. Radiation oncology. 2020;15(1):6.