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Cerebromicrovascular dysfunction predicts cognitive decline and gait abnormalities in a mouse model of whole brain irradiation-induced accelerated brain senescence

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Abstract Whole brain irradiation (WBI) is a mainstream therapy for patients with both identifiable brain metastases and prophylaxis for microscopic malignancies. However, it also promotes accelerated senescence in healthy tissues and leads to progressive cognitive dysfunction in up to 50% of tumor patients surviving long term after treatment, due to γ -irradiation-induced cerebromicrovascular injury. Moment-to-moment adjustment of cerebral blood flow (CBF) via neuronal activity-dependent cerebromicrovascular dilation (functional hyperemia) has a critical role in maintenance of healthy cognitive function. To determine whether cognitive decline induced by WBI associates with impaired cerebromicrovascular function,

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Z. Ungvari · P. Hertelendy · E. Farkas · A. Csiszar Department of Medical Physics and Informatics, Faculty of Medicine and Faculty of Science and Informatics, University of Szeged, Szeged, Hungary C56BL/6 mice (3 months) subjected to a clinically relevant protocol of fractionated WBI (5 Gy twice weekly for 4 weeks) and control mice were compared. Mice were tested for spatial memory performance (radial arm water maze), sensorimotor coordination (computerized gait analysis, CatWalk), and cerebromicrovascular function (whisker-stimulation-induced increases in CBF, measured by laser Doppler flowmetry) at 3 to 6 months post-irradiation. We found that mice with WBI exhibited impaired cerebromicrovascular function at 3 months post-irradiation, which was associated with impaired performance in the radial arm water maze. At 6 months, post-irradiation progressive impairment in gait coordination (including changes in the regularity index and phase dispersion) was also evident. Collectively, our findings provide evidence for early and persisting neurovascular impairment after a clinically relevant protocol of fractionated WBI, which predict early manifestations of cognitive impairment.

Keywords Whole brain irradiation · Neurovascular coupling · Functional hyperemia · Dementia · Gait dysfunction · Neurovascular unit · Cellular senescence · DNA damage

Introduction

Whole brain irradiation (WBI) is an important therapeutic treatment in patients with both identifiable brain metastases and prophylaxis for microscopic malignancies (Gaspar et al. 2010; Patil et al. 2012). Over 200,000 tumor patients with brain involvement are treated with

either partial large field or WBI every year in the USA (Lee et al. 2012). Although WBI is clinically effective in eliminating tumor cells, it also results in a range of unwanted side-effects in healthy tissues. In particular, irradiation-induced DNA damage is known to lead to accelerated brain senescence, which promotes functional impairment in the cortex, hippocampus, and white matter months to years after treatment (Khuntia et al. 2006; Welzel et al. 2008a, 2008b). As a result, up to 50% of patients who survive long term after treatment experience progressive dementia, thus severely compromising quality of life (Lee et al. 2012). The available experimental data clearly demonstrate that radiation also leads to a progressive impairment of cognitive function in animal models, mimicking the clinical scenario of cognitive decline in patients after WBI (Lamproglou et al. 2001; Shi et al. 2006; Soussain et al. 2009; Warrington et al. 2012; Welzel et al. 2008a, 2008b). Recent studies show that in addition to its profound harmful effects on cognitive function, WBI can also result in the impairment of other brain functions such as coordination of gait (Hatcher-Martin and Factor 2016). At present, no effective strategies exist to prevent radiation-induced decline in higher brain functions and there are no treatments available to reverse these effects.

Although the mechanisms underlying cognitive deficits after WBI are likely multifaceted, there is increasing pre-clinical and clinical evidence that WBI compromises the structural integrity of the cerebral microcirculation, which significantly contributes to cognitive impairment (Ashpole et al. 2014; Warrington et al. 2011, 2012, 2013). Previous studies demonstrate that clinical series of fractionated WBI induce a 40% decrease in capillary density in the brain regions associated with learning and memory (Warrington et al. 2012), mimicking aging phenotype. Despite these advances, there are no studies available elucidating the functional consequences of WBI-induced accelerated brain senescence on the cerebral microvasculature. Normal brain function is critically dependent on a continuous, tightly controlled supply of oxygen and glucose through cerebral blood flow (CBF). Although energetic demands of neurons are very high, the brain has very little reserve capacity. During periods of intense neuronal activity, there is a requirement for rapid increases in oxygen and glucose delivery. This is ensured by neurovascular coupling (functional hyperemia), a vital mechanism of regulation of CBF that maintains optimal microenvironment of cerebral tissue by adjusting local blood flow to local neuronal activity in a moment-to-moment manner (Attwell et al. 2010; Tarantini et al. 2015, 2016; Toth et al. 2016; Toth et al. 2014a, 2015a, 2015b, 2017; Tucsek et al. 2014). There is increasing evidence that neurovascular dysfunction is causally related to cognitive decline in models of aging (Tarantini et al. 2015, 2016), yet, the effects of WBI on cerebromicrovascular function have never been investigated.

The present study was designed to test the hypothesis that WBI-induced accelerated brain senescence associates with early-onset cerebromicrovascular dysfunction. To achieve this goal, C56BL/6 mice were subjected to a clinically relevant protocol of fractionated WBI (5 Gy twice weekly for 4 weeks). WBI-treated and control mice were tested for spatial memory performance (radial arm water maze), sensorimotor coordination (computerized gait analysis, CatWalk), and cerebromicrovascular function (whisker-stimulation-induced increases in CBF, measured by laser Doppler flowmetry) at 3 and 6 months post-irradiation.

Materials and methods

Experimental animals and experimental design

C57BL/6J (3 months old, n = 51) mice were purchased from the Jackson Laboratories (Bar Harbor, ME) and housed three per cage in the specific pathogen-free animal facility at the University of Oklahoma Health Sciences Center (OUHSC). Animals were kept on a 12-h light/dark cycle and fed standard rodent chow and water ad libitum. One week before radiation treatment, mice were transferred to the conventional facility (OUHSC) and housed under similar conditions. Mice were anesthetized and subjected to clinical series of WBI (n = 31, 5 Gy twice weekly for a total cumulative dose of 40 Gy) or used as a control group (n = 20). Mice were left to recover for 3 or 6 months in the original environment. At the end of the recovery period, mice were experimentally tested for cognitive function, gait coordination, and neurovascular coupling responses. All animal protocols were approved by the Institutional Animal Care and Use Committee of OUHSC.

Fractionated whole brain irradiation protocol

After acclimating to the conventional facility for 1 week, mice were randomly assigned to either anesthetized control or radiated groups. Animals were weighed and anesthetized via intra-peritoneal injection of ketamine and xylazine (100/15 mg per kg). Mice in the radiated group were subjected to clinical series of WBI (5 Gy twice weekly for a total cumulative dose of 40 Gy). Radiation was administered using a ¹³⁷Cesium gamma irradiator (GammaCell 40, Nordion International). A Cerrobend[®] shield was utilized to minimize the dose to the bodies of mice in the radiated group. The radiation dose received by the mice was verified using film dosimetry, as described (Warrington et al. 2011, 2012).

Radial arms water maze testing

To determine how accelerated brain senescence induced by WBI affects cognitive function, spatial memory and long-term memory was tested in mice by assessing performance in the radial arms water maze at 3 or 6 months post-WBI treatment. The maze consisted of eight arms with a submerged escape platform located at the end of one of the arms. Food coloring paint was added into the water to make it opaque. The maze was surrounded by privacy blinds with extramaze visual cues. Intramaze visual cues were placed at the end of the arms. The mice were monitored by a video tracking system directly above the maze as they waded and parameters were measured using Ethovision software (Noldus Information Technology Inc., Leesburg, VA, USA). Experimenters were unaware of the experimental conditions of the mice at the time of testing. During the learning period each day, mice were given the opportunity to learn the location of the submerged platform during two session blocks each consisting of four consecutive acquisition trials. On each trial, the mouse was started in one arm not containing the platform and allowed to wade for up to 1 min to find the escape platform. All mice spent 30 s on the platform following each trial before beginning the next trial. The platform was located in the same arm on each trial. Over 3 days of training, mice gradually improved performance as they learned the procedural aspects of the task. Upon entering an incorrect arm (all four paws within the distal half of the arm), the mouse was charged an error. Learning capability was assessed by comparing performance on days 2 and 3 of the learning period. When eventually both groups learned the procedural aspects of the task, the mice were placed in their home cage for 7 days. Then, the mice were administered the recall trial on day 10. On day 11 (extinction), mice were tested for ability to relearn the task, when the platform has been moved to a different arm which was not adjacent nor diametrically positioned to the previous location. Mice were tested for two session blocks and the second block consisting of four trials has been used for comparison.

Analysis of gait function

To determine how accelerated brain senescence induced by WBI affects gait coordination, we tested the animals using an automated computer assisted method (CatWalk; Noldus Information Technology Inc.) at 3 and 6 month post-WBI. Using the CatWalk system, the detection of paw print size and paw placement patterns during volunteer running on an illuminated glass walkway by a camera placed under the glass surface provides an automated analysis of gait function and the spatial and temporal aspects of inter-limb coordination (Tarantini et al. 2015). Briefly, animals were trained to cross the walkway and then, in a dark and silent room (<20 lx of illumination), animals were tested in three consecutive runs. Data were averaged across ten runs in which the animal maintained a constant speed across the walkway.

The variability of the data has been assessed using quartile dispersion. We adopted a common outlier definition, labeling points more than 1.5 interquartile ranges away from the sample median as extreme values. After variability analyses, the following indices were calculated. The regularity index (%) is a fractional measure of interpaw coordination, which expresses the number of normal step sequence patterns relative to the total number of paw placements. The formula of regularity index is NSSP \times 4/PP \times 100 (%), where NSSP represents the number of normal step sequence patterns and PP the total number of paw placements. In healthy, fully coordinated animals, its value is close to 100%. Phase dispersion provides a quantitative metric of interpaw coordination. Phase dispersion characterizes the placement of two paws ("target" and "anchor") during the cycle of consecutive initial contacts with an "anchor" paw. In a step cycle, base of support gives the distance between the mass midpoints of the fore prints at maximal contact. The results are averaged and expressed in centimeters.

Assessment of neurovascular coupling responses

To determine how accelerated brain senescence induced by WBI affects cerebromicrovascular function, cerebral blood flow responses to whisker stimulation were assessed at 3 months post-WBI treatment. Mice were anesthetized (a-chloralose, 50 mg/kg plus urethane, 750 mg/kg, i.p.), endotracheally intubated and ventilated (MousVent G500; Kent Scientific Co, Torrington, CT). Rectal temperature was maintained at 37 °C using a thermostatic heating pad (Kent Scientific Co, Torrington, CT). End-tidal CO₂ (with dead space) was kept between 3.2 and 3.7% to maintain blood gas values within the physiological range, as reported (Toth et al. 2014a). Animals were immobilized and placed on a stereotaxic frame (Leica Microsystems Inc., Buffalo Grove, IL). The scalp and periosteum were pulled aside and a craniotomy was made with a dental drill over the left somatosensory cortex corresponding to the barrel field as described (Toth et al. 2015b). The dura was gently removed, and the open cranial window was continuously superfused with artificial cerebrospinal fluid (ACSF, composition: NaCl 119 mM, NaHCO₃ 26.2 mM, KCl 2.5 mM, NaH₂PO₄ 1 mM, MgCl₂ 1.3 mM, glucose 10 mM, CaCl₂ 2.5 mM, pH = 7.3, 37 °C). The right femoral artery was cannulated for arterial blood pressure measurement (Living Systems Instrumentations, Burlington, VT) to ensure that the blood pressure was within the physiological range throughout the experiments (90-100 mmHg). To assess neurovascular coupling, a laser Doppler probe (Transonic Systems Inc., Ithaca, NY) was positioned above the barrel cortex (1-1.5 mm posterior and 3-3.5 mm lateral to bregma), and the contralateral whiskers were stimulated for 30 s at 5 Hz from side to side, as previously described (Toth et al. 2014a, 2015b). Changes in CBF (n = 7-8 in each group) were assessed in three trials divided by 5-10 min intervals and are expressed as percent (%) changes from baseline.

WBI could affect functional hyperemia by impairing neural activity evoked by whisker stimulation. To examine this possibility, we recorded spontaneous and evoked neural activity in control and WBI-treated mice. A glass-insulated tungsten microelectrode (impedance, 2–3 M Ω ; Kation Scientific, LLC, Minneapolis, MN, USA) was inserted into the left barrel cortex (1– 1.5 mm posterior and 3–3.5 mm lateral to bregma) through the ACSF-perfused open cranial window for recording local field potentials as described (Toth et al. 2015a). An Ag/AgCl electrode inserted in the neck muscles served as reference. After basal activity was recorded, the right whisker pad was stimulated by a bipolar-stimulating electrode placed to the ramus infraorbitalis of the trigeminal nerve and into the masticatory muscles. The stimulation protocol used to investigate neurovascular coupling and somatosensory evoked field potentials consisted of ten stimulation presentation trials with an intertrial interval of 70 s, each delivering a 15-s train of electrical pulses (2 Hz, 0.2 mA, intensity, and 0.3 ms pulse width) after a 10-s prestimulation baseline period. The signal was amplified with an AC/DC differential amplifier (high pass at 1 Hz, low pass at 1 kHz) (Model 3000; A-M Systems, Inc. Carlsborg, WA, USA) and digitalized by the PowerLab/Labchart data acquisition system (ADInstruments, Colorado Springs, CO, USA) with a sampling rate of 40 kHz. Basal activity was analyzed as distribution of wave amplitude as a function of frequency, and the negative amplitude in the somatosensory evoked field potential response was considered as the excitatory postsynaptic potential (Toth et al. 2014b). Analyses were performed on the average of ten stimulation trials.

Statistical analysis

Statistical analysis was carried out by unpaired t test, one-way ANOVA, or two-way ANOVA for repeated measures followed by Bonferroni multiple comparison test, as appropriate, using Prism 5.0 for Windows (Graphpad Software, La Jolla, CA). A p value less than 0.05 was considered statistically significant. Data are expressed as mean \pm S.E.M.

Results

Cognitive function is impaired in mice by 3 months post-WBI

Previously, we demonstrated that C57BL/6J mice exhibit significant impairment of hippocampal-encoded functions of learning and memory at 5 months post-WBI treatment (Warrington et al. 2013). To gain insight into the time-course of the development of irradiation-induced cognitive decline, in the present study, we assessed learning and memory function in mice 3 months post-WBI and in age-matched controls in the radial arm water maze (Fig. 1a, left and right panel, respectively). We compared the learning performance of mice in each experimental group by analyzing the day-to-day changes in the path length and error rate. The error rate is the sum of all incorrect arm entries, where

each error corresponds to each incorrect entry. The error rate for each mouse is then averaged among each experimental group in every trial block. During acquisition, mice from all groups showed a decrease in the path length (Fig. 1a) and the combined error rate (Fig. 1b) across days, indicating learning of the task. We found that control mice and mice at 3 months post-WBI exhibited similar learning during the first 3 days of testing (Fig. 1b, c). We confirmed that mice exhibited significant impairment of learning function when tested 6 months post-WBI (data not shown). Memory recall 7 days later was also similar in both groups. WBItreated mice showed impaired extinction ability on day 11 (Fig. 1b, c). The extinction blocks describe the ability of the mouse to forget and relearn the task with a different platform location. These findings suggest that

Mild gait dysfunction becomes evident 6 months post-WBI

We performed gait analyses using the Catwalk system in control and WBI-treated animals. We found that WBItreated mice showed no difference in gait coordination at 3 months post-irradiation (Fig. 2a–f). However, when followed further for another 3 months, significantly lower regularity index and significantly higher phase dispersion was evident in WBI-treated mice as compared to controls (Fig. 3). These data suggest that gait

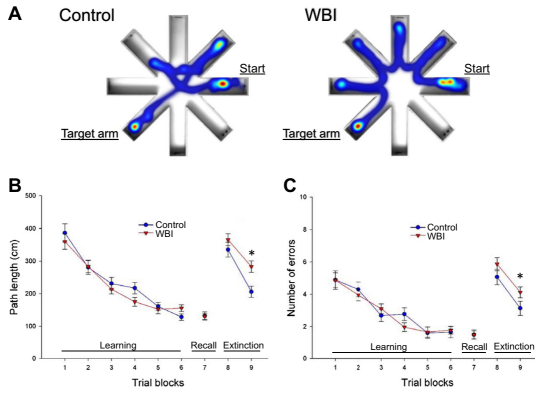


Fig. 1 WBI treatment results in impaired radial arm water maze (RAWM) performance in mice. WBI-treated mice and agedmatched control mice were tested in the RAWM 3 months post-WBI. **a** Heatmaps representing the percentage of time spent in different locations in the maze for a randomly selected control (*left*) and WBI-treated (*right*) animal during experimental day 11 in trial block 9. **b** Average path length required to reach the hidden platform in the RAWM. During acquisition (trial blocks 1 to 6) and memory recall (trial block 7) in WBI-treated mice, the average

path length required to reach the hidden platform did not differ from that in control mice at 3 months post-irradiation. Combined error rates (**c**) were also similar in both groups during acquisition (trial blocks 1 to 6) and memory recall (trial block 7). In contrast, WBI-treated animals traveled longer distance (**b**) and made more errors (**c**) before they were able to relearn the task when the platform was moved to a different location during extinction phase. Data are mean \pm SEM (*p < 0.05)

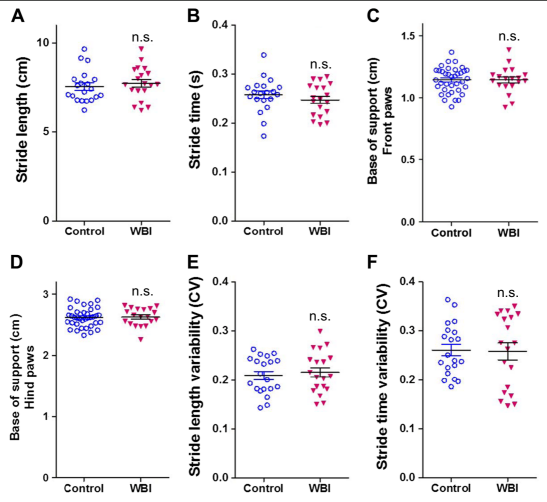


Fig. 2 Effects of WBI on gait parameters at 3 months posttreatment. Stride length (a), stride time (b), base of support (front paws); c), base of support (hind paws); d), stride length variability

(e), and stride time variability (f) in control mice and WBI-treated mice at 3 months post-treatment. Data are mean \pm SEM

abnormalities associated with WBI-induced accelerated brain senescence exhibit later onset than memory impairment.

Cerebromicrovascular function is significantly impaired in mice by 3 months post-WBI

In order to determine whether cerebromicrovascular dysfunction associates with cognitive decline, we assessed functional hyperemia in the whisker barrel cortex in mice at 3 months post-WBI. We found that CBF responses in the whisker barrel cortex elicited by contralateral whisker stimulation were significantly decreased in WBI-treated mice compared to control animals indicating impaired neurovascular coupling at 3 months post-irradiation (representative CBF tracings are shown in Fig. 4a, summary data are shown in Fig. 4b). WBI could reduce functional hyperemia by impairing neural activity evoked by whisker stimulation. To examine this possibility, we recorded spontaneous and evoked neural activity in control and WBI-treated mice. We found that the amplitude and frequency distribution of the electrocorticogram (data not shown) and the amplitude of the somatosensory field potentials produced by activation of the whisker pad were not different between control and WBI-treated mice (Fig. 4c). Therefore, WBI is unlikely to contribute to impaired functional hyperemia by modulating the neural activity evoked by whisker stimulation.

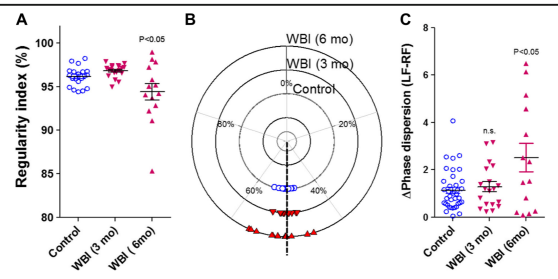


Fig. 3 WBI results in progressive gait abnormalities in mice at 6 months post-treatment. **a** Regularity index in control mice and WBI-treated mice at 3 and 6 months post-treatment. **b** Circular scatter plot showing the distribution of inter-limb coupling values (phase dispersion) in control mice and WBI-treated mice at 3 and 6 months post-treatment (note that the circular plot shows a smaller

Discussion

The key finding of his study is that in mice a clinically relevant protocol of fractionated WBI results in a significant cerebromicrovascular dysfunction, which associates with cognitive impairment, mimicking the aging phenotype.

There is growing evidence that vascular contributions to cognitive impairment and dementia (VCID) are critical in aging (Ungvari and Sonntag 2014). Among them,

phase dispersion scatter in the inner circle (control) as compared to the phase dispersion scatter in the outer circle (assessed 6 month after WBI)). **c** This shows average deviation of phase dispersion (calculated between the right front paw (RF) and left front paw (LF)) from the expected value (50%) under baseline conditions and after WBI. Data are mean \pm SEM.*p < 0.05 vs. control

impaired neurovascular coupling responses (Fabiani et al. 2013; Park et al. 2007; Stefanova et al. 2013; Topcuoglu et al. 2009; Toth et al. 2014a; Zaletel et al. 2005) are thought to importantly contribute to the cognitive decline (Sorond et al. 2013) in the elderly. From the results of this study and from recent findings by other laboratories (Warrington et al. 2013), the picture emerges that WBI-induced DNA damage responses induce an accelerated aging-like phenotype in the neurovascular

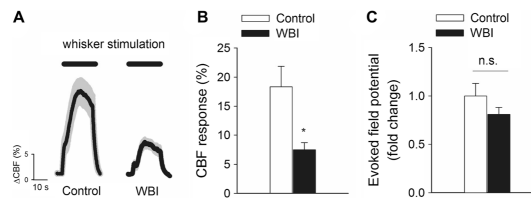


Fig. 4 Whole brain irradiation impairs neurovascular coupling responses 3 months post-WBI. a Representative traces of CBF measured with a laser Doppler probe above the whisker barrel cortex during contralateral whisker stimulation (5 Hz) in control and WBI-treated mice 3 months post-WBI. b Increases in cerebral blood flow (CBF; expressed as % of baseline) measured above the barrel field of the primary somatosensory cortex in response to

whisker stimulation in control mice and in WBI-treated mice. **c** The somatosensory evoked potential responses in the somatosensory cortex evoked by contralateral whisker pad stimulation are comparable in control and WBI-treated mice. The amplitude of the negative wave of the field potentials was not significantly different. Data are mean \pm S.E.M (n = 6, *p < 0.05 vs. control)

unit, which results in impaired CBF responses upon neuronal activation. We posit that impaired functional hyperemia per se may play a significant causal role in irradiation-induced cognitive impairment. Experimental studies support this concept, demonstrating that pharmacologically induced selective neurovascular uncoupling in mice mimics important aspects of both age-related and WBI-induced cognitive impairment (Tarantini et al. 2015). There are also studies extant linking impaired neurovascular coupling to gait dysfunction in the elderly (Sorond et al. 2011). Thus, we speculate that WBIinduced neurovascular coupling may also contribute to the gait/motor coordination abnormalities observed in our present study as well as in previous reports (Brown et al. 2016; Hatcher-Martin and Factor 2016). Further studies are warranted to determine whether changes in gait after WBI correlate with the severity of post-radiation cerebromicrovascular deficits. Previous studies by us and others have shown that WBI also results in a significant microvascular rarefaction (Ljubimova et al. 1991; Warrington et al. 2012), particularly in the hippocampus, the region responsible for spatial learning and memory (Warrington et al. 2011). It is likely that neurovascular uncoupling and microvascular rarefaction act synergistically to impair regional blood supply in the brain (Fuss et al. 2000; Taki et al. 2002), exacerbating cognitive decline. It is important that both neurovascular dysfunction and microvascular rarefaction are manifested relatively early (within 3 months) after WBI and their onset coincides with the appearance of cognitive symptoms.

The mechanisms underlying WBI-induced neurovascular dysfunction are presently unknown and should be elucidated in future studies. Neurovascular coupling depends on an intact functional network of neurons, astrocytes, and vascular cells (Attwell et al. 2010; Chen et al. 2014). Unlike neurons, cells of the neurovascular unit (including astrocytes and endothelial cells) are radiosensitive (Ungvari et al. 2013). There is evidence that WBI significantly impacts astrocytes, the functional impairment of which has also been linked to radiation-induced blood brain barrier disruption (Gaber et al. 2004; Yuan et al. 2006; Yuan et al. 2003; Yuan et al. 2005). The astrocytic mechanisms underlying neurovascular coupling include production and release of vasodilator metabolites of arachidonic acid, including epoxygenase-derived epoxyeicosatrienoic acids (EETs) and cyclooxygenase-derived prostaglandins (Peng et al. 2002; Takano et al. 2006; Zonta et al. 2003). Future studies should determine how DNA damage responses in astrocytes alter the synthesis/release of these mediators. There is also strong experimental evidence that endothelial NO production contributes to glio-vascular coupling (Chen et al. 2014; Girouard et al. 2007; Longden and Nelson 2011; Ma et al. 1996; Stobart et al. 2013; Toth et al. 2015b). Because the phenotype of cerebromicrovascular endothelial cells can also be altered by irradiation-induced DNA damage (Ungvari et al. 2013; Warrington et al. 2013); future studies should also determine the role of endothelial dysfunction in WBI-induced impairment of functional hyperemia.

In conclusion, our results add to the growing evidence that stressors that cause DNA damage promote the development of complex aging-like phenotypic changes in the brain, which are characterized by functional and structural impairment of the cerebral microcirculation. It is significant that cognitive decline due to microvascular causes is potentially reversible (Warrington et al. 2011, 2012). Our findings, taken together with the results of earlier studies (reviewed in Warrington et al. (2013)), point to potential benefits of interventions that rescue glio-vascular coupling mechanisms and promote microvascular health for prevention of cognitive decline both in WBI-treated patients and the elderly. There is increasing realization that DNA damage-induced cellular senescence programs play a critical role in brain aging process (reviewed in Chinta et al. (2014)), thus we propose that the present clinically relevant model of fractionated WBI should be considered to study fundamental mechanisms of brain and cognitive aging as well.

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Compliance with ethical standards

Disclosure None.

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